Effect of n-3 PUFA supplementation at different EPA:DHA ratios on the spontaneously hypertensive obese rat model of the metabolic syndrome

Eunice Molinar-Toribio¹, Jara Pérez-Jiménez¹*,†, Sara Ramos-Romero¹,², Marta Romeu³, Montserrat Giralt³, Núria Taltavull³, Mónica Muñoz-Cortes³, Olga Jáuregui⁴, Lucía Méndez⁵, Isabel Medina⁵ and Josep Lluís Torres¹

¹Institute of Advanced Chemistry of Catalonia (IQAC-CSIC), Barcelona, Spain
²Biomedical Research Networking Center in Bioengineering, Biomaterials, and Nanomedicine (CIBER-BBN), Zaragoza, Spain
³Unidad de Farmacología, Facultad de Medicina y Ciencias de la Salud, Universidad Rovira i Virgili, Reus, Spain
⁴Scientific and Technological Centers of the University of Barcelona (CCIT-UB), Barcelona, Spain
⁵Instituto de Investigaciones Marinas (IIM-CSIC), Vigo, Spain

(Submitted 17 June 2014 – Final revision received 3 December 2014 – Accepted 17 December 2014 – First published online 27 February 2015)

Abstract
The increasing incidence of the metabolic syndrome (MetS), a combination of risk factors before the onset of CVD and type 2 diabetes, encourages studies on the role of functional food components such as long-chain n-3 PUFA as preventive agents. In the present study, we explore the effect of EPA and DHA supplementation in different proportions on spontaneously hypertensive obese (SHROB) rats, a model for the MetS in a prediabetic state with mild oxidative stress. SHROB rats were randomised into four groups (n 7), each supplemented with EPA/DHA at ratios of 1:1, 2:1 and 1:2, or soyabean oil as the control for 13 weeks. The results showed that in all the proportions tested, EPA/DHA supplementation significantly lowered total and LDL-cholesterol concentrations, compared with those of the control group. EPA/DHA supplementation at the ratios of 1:1 and 2:1 significantly decreased inflammation (C-reactive protein levels) and lowered oxidative stress (decreased excretion of urinary isoprostanes), mainly at the ratio of 1:2. The activity of antioxidant enzymes increased in erythrocytes, abdominal fat and kidneys, with magnitudes depending on the EPA:DHA ratio. PUFA mixtures from fish affected different MetS markers of CVD risk factors in SHROB rats, depending on the ratios of EPA/DHA supplementation. The activation of endogenous defence systems may be related to the reduction of inflammation and oxidative stress.

Key words: Metabolic syndrome; n-3 PUFA; Spontaneously hypertensive obese rats; EPA/DHA

The metabolic syndrome (MetS) is a combination of features such as insulin resistance hypercholesterolaemia, hypertriacylglycerolaemia, hypertension or obesity(1) that makes subjects with the MetS highly prone to developing type 2 diabetes mellitus and CVD. The MetS is also accompanied by low-grade chronic inflammation(2) and increased oxidative stress (OS)(3).

The increasing incidence of the MetS has encouraged the search for preventive strategies, based on either drugs or dietary approaches, including diet supplementation with bioactive compounds from food sources. Long-chain n-3 PUFA of marine origin are key components of cell membranes that may be used as a functional food and dietary supplement. The two most common n-3 PUFA in fish oils are EPA (20 : 5) and DHA (22 : 6). Studies using different animal models, such as rats with sucrose-induced or fructose-induced MetS, have shown that regular supplementation with n-3 PUFA decreases blood pressure and levels of insulin, TAG, cholesterol, NEFA or total lipids (4,5). Because n-3 PUFA play the role of mediators in nuclear events, they modulate these parameters via different mechanisms, including interactions with other nutrients and effects on the expression of genes involved in lipid and glucose metabolism(6).

In human subjects, n-3 PUFA have been shown to modulate certain characteristic MetS features such as insulin resistance,
hypertension and dyslipidaemia, although their effects on other aspects of CVD, such as the incidence of myocardial infarction or arrhythmia, remain controversial\(^7\). However, it should be noted that, to date, most trials have not considered that the relative proportion of EPA and DHA may influence the results, which might explain some of these controversies. Indeed, it has been observed that different proportions of \(n\)-3 and \(n\)-6 PUFA have different effects on health\(^8\), so different proportions within the \(n\)-3 series may also differentially affect MetS risk factors. Recently, we compared the effects of supplementation with EPA/DHA at different ratios on the markers of CVD and OS in healthy Wistar Kyoto (WKY) rats, and found that the proportion 1:1 was especially favourable for the improvement of some risk factors of type 2 diabetes mellitus, such as glycated Hb levels, as well as for a reduction in OS, as shown by increased plasma antioxidant capacity and decreased protein oxidation\(^5,10\).

Spontaneously hypertensive obese (SHROB or Koletsky) rats are an animal model of the MetS. The strain exhibits a mutation that introduces a premature stop codon in the extracellular leptin receptor domain, which results in truncated receptors that are insensitive to leptin\(^11,12\). The rats are obese, hypertensive and hyperinsulinaemic, although they are not hyperglycaemic. They have been suggested as an appropriate model for the MetS in a prediabetic state\(^13,14\). The aim of the present study was to evaluate the effect of supplementation with EPA/DHA mixtures at different proportions found in fish and fish oil-based supplements, namely 1:1, 2:1 and 1:2, on the markers of CVD and OS in SHROB rats, as an animal model for the MetS. To the best of our knowledge, the present study was the first to investigate the effect of \(n\)-3 PUFA in SHROB rats.

**Materials and methods**

**Materials and reagents**

The standard A04 diet was obtained from Harlan Iberica. Oils with different EPA:DHA ratios were obtained by mixing appropriate amounts of the commercial fish oils AFAMPES 121 EPA (AFAMSA), EnzerZona Omega 3 RX and Ogilen liquid DHA 80% (IFIGEN-Equip 98, S.L.). Soyabean oil, obtained from unrefined organic soyabeans (first cold pressing), was from Clearspring Limited. Other reagents and chemicals were as follows: ketamine chloride from Merial Laboratorios; xylazine from Quimica Farmaceutica; \(\beta\)-glucuronidase (90 U/\(\mu\)l) from Sigma-Aldrich. Also, 15-F\(_2\)-isoprostanes (IsoP), \([\text{H}^4]\text{15-F}\(_2\)-IsoP, 5-F\(_2\)-IsoP, 5-F\(_3\)-IsoP, 8-F\(_3\)-IsoP, 15-F\(_3\)-IsoP and 2,3-dinor-15-F\(_2\)-IsoP were obtained from Cayman Chemical, which were used without further purification.

Ethanol (analytical grade) was purchased from Carlo Erba. Methanol, heptane, acetonitrile (all HPLC grade) and formic acid (analytical grade), hydrochloric acid fuming 37%, ethanol (absolute), ethyl acetate, chloroform, \(\text{Na}_2\text{CO}_3\), \(\text{NaHCO}_3\), EDTA, \(\alpha\)-phthalaldehyde, \(N\)-ethylmaleimide and albumin were obtained from Merck. Iodoacetamide, EDTA, Tris—HCl, acetone, phosphomolybdic acid, Bradford reagent, NADPH, GR (glutathione reductase from baker’s yeast), GSH (1-glutathione reduced, minimum 99.9\%), tert-butyl hydroperoxide solution, GSSG (1-glutathione oxidised) and fluorescein-5-thiosemicarbazide were all obtained from Sigma-Aldrich. \(\text{N},\text{O}\)-Bis(trimethylsilyl)trifluoro-acetamide was from Supelco. Drabkin’s reagent was purchased from Quimica Clinica Aplicada. \(\text{Na}_2\text{HPO}_4\), \(\text{NaH}_2\text{PO}_4\), \(\text{NaCl}\), epinephrine, \(\text{KH}_2\text{PO}_4\), \(\text{K}_2\text{HPO}_4\), TCA and \(\text{NaOH}\) were purchased from Panreac Quimica. Water for the assay solutions was obtained using a Milli-Q water purification system from Millipore Corporation.

**Animals**

A total of twenty-eight female SHROB rats (Charles River Laboratories), aged 11–14 weeks, were used for the present study. They were kept in a room containing no male animals to minimise the influence of hormonal alterations. All the procedures strictly adhered to the European Union guidelines for the care and management of laboratory animals, and approved by the CSIC Subcommittee of Bioethical Issues (reference no. AGL2009-12 374-C03-03).

**Experimental design**

Rats were kept in Makrolon cages (\(n\) 2–3 per cage) under controlled conditions of stable humidity (50 ± 10%) and temperature (22 ± 2°C) with a 12 h light–12 h dark cycle. Rats were randomly divided into four groups, each (\(n\) 7) supplemented with a different oil mixture: EPA/DHA 1:1, EPA/DHA 2:1, EPA/DHA 1:2 or soybean oil as the control (isoenergetic without \(n\)-3 long-chain PUFA). After 1 week of acclimatisation, all the groups were fed a standard pelleted A04 diet, given water (Ribes) ad libitum, and orally administered with the corresponding oil mixture or soybean oil at a weekly dose of 0.8 ml/kg body weight for 13 weeks. All the diets had similar fat and energy content (see online supplementary Table S1). Moreover, the weekly dose of oil orally administered did not alter the overall intake of the animals in terms of macronutrients, compared with a standard diet, with 20% of energy derived from proteins, 67% from carbohydrates and 13% from fat (see online supplementary Table S1). The fatty acid composition of the supplements is also provided in Table S2 (available online). Because PUFA are extremely susceptible to oxidation and the by-products are potentially toxic, lipid oxidation levels were checked throughout the experiment (peroxide values < 5 mEq oxygen/kg oil). Intragastric administration was selected because PUFA have a strong tendency to oxidise when incorporated into the animal feed, as demonstrated by our previous study (data not shown). Oxidised PUFA may seriously alter the results, as described for commercial pills\(^15\). After week 16, rats were fasted overnight, anaesthetised intraperitoneally with ketamine and xylazine (80 and 10 mg/kg body weight, respectively), and then killed by exsanguination.

**Sample collection**

For urine collection, at week 14/15 of the experiment, the rats were placed in metabolic cages and deprived of food for 18h,
after which the samples were collected. After 1 week, blood was collected by cardiac puncture under anaesthesia, and plasma, serum and erythrocytes were also collected. Abdominal fat was weighed and immediately frozen. Tissue samples collected from the heart, brain, liver and kidneys were washed with 0·9% NaCl solution, dried, weighed and immediately frozen in liquid N₂. All the samples were stored at −80°C until analysis. Before analysis, the tissue samples were homogenised with sodium phosphate buffer and ultra-centrifuged.

**CVD risk factors**

**Physical measurements.** Body weight was monitored weekly throughout the experiment. Abdominal fat was collected post-mortem, weighed and expressed as a percentage of body weight.

**Determination of plasma lipid profile.** Plasma total cholesterol, LDL-cholesterol (LDL-C), HDL-cholesterol and TAG, as well as ApoA1 and ApoB100 concentrations were determined by spectrophotometric methods using the corresponding kits from Cusabio Biotech Co. Plasma oxidised LDL (LDL-ox) was determined using an ELISA kit (Cusabio Biotech Co.) by measuring absorbance at 450 nm with a PowerWave XS2 spectrophotometer (Biotek Instruments, Inc.).

**Determination of glycaemia.** Blood glucose levels were determined by the enzyme electrode method using an Ascensia ELITE XL glucose meter (Bayer Consumer Care). Plasma insulin level was determined using an ELISA kit (Millipore). Blood glyated Hb level was measured using a spectrophotometric kit (Spinreact).

**Markers of endothelial function, inflammation and thrombotic activity.** The corresponding ELISA kits from Cusabio Biotech Co. were used to measure the following parameters in plasma: vascular cell adhesion molecule-1 and intercellular adhesion molecule-1, as markers of endothelial function; C-reactive protein (CRP, detection range 0·16–10 ng/ml), as an inflammation marker; plasminogen activator inhibitor-1, as a marker of thrombotic activity.

**Endogenous antioxidant systems**

The activities of superoxide dismutase (SOD) and catalase (CAT) were measured in erythrocytes, kidneys, abdominal fat, heart and brain using standard spectrophotometric methods[16,17] .

GR and glutathione peroxidase (GPx) activities were determined in erythrocytes and kidneys by spectrophotometric methods[18,19] . GSH and GSSG activities were determined in kidneys using fluorometric methods at wavelengths of 350 nm (excitation) and 420 nm (emission)[20] . Measurements in erythrocytes were normalised to the amount of Hb by the Drabkin method[21] , while those in tissue were normalised to the amount of protein estimated by the Bradford method[22] .

**Isoprostanes**

IsoP of the F₂ and F₃ series were determined in urine samples by liquid chromatography coupled to tandem MS after electrospray ionisation (LC–ESI–QqQ–MS/MS), according to a procedure described previously[23,24] , with some modifications. Briefly, stock standard solutions of all IsoP were prepared in ethanol (1 ng/µl) and stored at −20°C under N₂. The mixture of standards was prepared by mixing and diluting the appropriate stock solutions to final concentrations between 0·5 and 400 pg/µl. The internal standard [²H₄]15-F₂t-isoP was prepared in ethanol (1 ng/µl), and added to the mixture of standards and samples at a final concentration of 0·5 ng/µl. Urine samples (500 µl) were acidified with formic acid to pH 4·5–5·0. Then, 10 µl of β-glucuronidase (90 U/µl) were added to each sample and incubated at 37°C for 2 h. Following incubation, the samples were cooled to room temperature, and then the pH was adjusted again to approximately 2·5 with formic acid. After the addition of the isotope-labelled internal standard [²H₄]15-F₂t-isoP, the samples were subjected to solid-phase extraction on a C18 Sep-Pak cartridge (Waters Associates). The cartridges were preconditioned with 5 ml methanol followed by 7 ml of acid water (water acidified to pH 3 with formic acid) before loading the samples. To remove interfering components, the cartridges were washed with 10 ml acid water, 10 ml acid water–methanol (9:1, v/v), and 10 ml heptane in succession. Then, the IsoP were eluted with 1 ml methanol, the eluate was evaporated under N₂, and the residues reconstituted with 100 µl of the liquid chromatography starting mobile phase (A, see below). The IsoP were analysed by LC–ESI–QqQ–MS/MS using an Agilent 1100 HPLC series system; equipped with an autosampler with the temperature set to 8°C; fitted with a Luna C18 column (50 X 2·0 mm inner diameter; 3·0 µm packing; Phenomenex) and coupled to an AB Sciex API 3000 triple quadrupole mass spectrometer. The instrument was operated in the negative-ion mode with a Turbolon spray source to obtain MS and MS/MS data, and the source temperature at 450°C. In the separation step, elution was accomplished with a binary system consisting of 0·1% aqueous formic acid (A) and 0·1% formic acid in CH₃CN (B), under an increasing linear gradient (v/v) of B (0·1 % aqueous formic acid) at room temperature, and then the pH was adjusted again to 3·0 with formic acid. The IsoP were detected in multiple reaction monitoring experiments, and their identity was confirmed using the corresponding standard. Concentrations were interpolated in the calibration curves of each standard (R² > 0·99 in all cases). Sample concentrations were calculated from the equation y = mx + b, as determined by weighted (1/x²) linear regression of the standard curve.

The limit of detection was established at a signal:noise ratio of 3, and the values were between 0·5 and 3·0 pg/µl depending on the standard. The limit of quantification was established at a signal:noise ratio of 10, and the values were between 1·0 and 10·0 pg/µl, depending on the standard. Peak integrations and signal:noise calculations were performed using 1.4.2 Analyst software (Applied Biosystems). The dwell time for multiple reaction monitoring experiments was 100 ms, and the cycle time was 2 s. The declustering potential and collision energy in multiple reaction monitoring mode were optimised for each standard in infusion experiments: 55 V and 28 eV for [²H₄]15-F₂t-isoP (357 → 197); 55 V and 38 eV for 15-F₂t-isoP (355 → 193); 50 V and 22 eV for 2,3-dinor-15-F₂t-isoP (325 → 237); 55 V and 34 eV for...
The results are expressed as pg IsoP/mg creatinine to correct for urine concentration. Creatinine levels from the urine samples were determined by a colorimetric method using a commercial kit (Cromatest Linear Chemicals) by measuring absorbance at 510 nm on a SpectraMax M5 spectrophotometer (Molecular Devices). Examples of the detection of IsoP in the standard mixture and in a sample are shown in Figs. S1 and S2 (available online).

Statistical analyses

Results from the fatty acid composition (see online supplementary Table S1) are expressed as means and standard deviations, and those from the measurements in rat samples are expressed as means with their standard errors. Levene’s test and the Kolmogorov–Smirnov test were applied to assess variance equality and normal distribution, respectively. One-way ANOVA followed by Tukey’s post hoc significance test was used when the assumptions of normality and equal variance were met. Otherwise, non-parametric tests (Kruskal–Wallis and Mann–Whitney U rank-sum) were used to assess significance. Differences were considered significant at \( P < 0.05 \), and considered to indicate a tendency at \( P < 0.1 \). All statistical analyses were performed using the statistical package SPSS IBM version 19 for Windows.

Results

Physical measurements

Supplementation with EPA/DHA in different proportions did not modify either the animal body-weight time course, or the percentage of abdominal fat compared with the control group (data not shown).

Endogenous antioxidant systems

SOD and CAT activities. The activities of these antioxidant enzymes were evaluated in erythrocytes, kidneys, abdominal

Table 1. CVD risk factors in spontaneously hypertensive obese rats supplemented with EPA/DHA at different proportions (1:1, 2:1 and 1:2) and soyabean oil as the control diet

<table>
<thead>
<tr>
<th>Category</th>
<th>Parameter</th>
<th>Control</th>
<th>1:1</th>
<th>2:1</th>
<th>1:2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Mean</td>
<td>SEM</td>
<td>Mean</td>
<td>SEM</td>
</tr>
<tr>
<td>Plasma lipid profile</td>
<td>Total cholesterol (mg/ml)§</td>
<td>3·2</td>
<td>0·5</td>
<td>2·1†</td>
<td>0·2</td>
</tr>
<tr>
<td></td>
<td>LDL-C (mg/ml)§</td>
<td>0·9</td>
<td>0·3</td>
<td>0·3*</td>
<td>0·1</td>
</tr>
<tr>
<td></td>
<td>LDL-ox (mg/ml)†</td>
<td>154·6</td>
<td>23·4</td>
<td>279·3***</td>
<td>14·6</td>
</tr>
<tr>
<td></td>
<td>HDL-C (mg/ml)</td>
<td>0·8</td>
<td>0·1</td>
<td>0·7</td>
<td>0·1</td>
</tr>
<tr>
<td></td>
<td>Total TAG (mg/ml)</td>
<td>14·6</td>
<td>3·4</td>
<td>5·7*</td>
<td>0·8</td>
</tr>
<tr>
<td>Glycaemia</td>
<td>Glucose (mg/ml)</td>
<td>0·6</td>
<td>0·02</td>
<td>0·6</td>
<td>0·01</td>
</tr>
<tr>
<td></td>
<td>Insulin (ng/ml)</td>
<td>5·8</td>
<td>1·0</td>
<td>12·2**</td>
<td>2·0</td>
</tr>
<tr>
<td></td>
<td>Glycated Hb (%)</td>
<td>6·8</td>
<td>0·4</td>
<td>4·2***</td>
<td>0·1</td>
</tr>
<tr>
<td>Inflammation</td>
<td>CRP (µg/ml)</td>
<td>285·9</td>
<td>23·8</td>
<td>206·0*</td>
<td>15·9</td>
</tr>
<tr>
<td>Endothelial dysfunction</td>
<td>VCAM-1 (µg/ml)§</td>
<td>2·7</td>
<td>0·3</td>
<td>3·8</td>
<td>0·1</td>
</tr>
<tr>
<td></td>
<td>ICAM-1 (ng/ml)†</td>
<td>1·0</td>
<td>0·1</td>
<td>0·9</td>
<td>0·1</td>
</tr>
<tr>
<td>Thrombotic activity</td>
<td>PAI-1 (µg/ml)</td>
<td>11·6</td>
<td>0·6</td>
<td>10·5</td>
<td>0·8</td>
</tr>
</tbody>
</table>

LDL-C, LDL-cholesterol; LDL-ox, oxidised LDL; HDL-C, HDL-cholesterol; CRP, C-reactive protein; VCAM-1, vascular cell adhesion molecule-1; ICAM-1, intercellular adhesion molecule-1; PAI-1, plasminogen activator inhibitor-1.

Mean value was significantly different from that of the control group: * \( P < 0.05 \); ** \( P < 0.01 \); *** \( P < 0.001 \).

† Mean value tended to be different from that of the control group (\( P < 0.1 \)).

‡ Mean value was significantly different from that of the 1:1 group (\( P < 0.05 \)).

§ Comparisons were made using one-way ANOVA and Tukey’s post hoc significance tests.

∥ Comparisons were performed using Kruskal–Wallis and Mann–Whitney U tests.

Downloaded from https://www.cambridge.org/core. IP address: 54.70.40.11, on 13 Oct 2018 at 21:09:39, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms. https://doi.org/10.1017/S0007114514004437
Compared with the soya bean oil-supplemented control group, SOD activity was significantly higher in the erythrocytes, kidneys, abdominal fat, heart and brain of the 1:1 EPA/DHA group, in the kidneys and heart of the EPA/DHA

---

**Fig. 1.** (a), (c), (e), (g), (i) Superoxide dismutase and (b), (d), (f), (h), (j) catalase activities in different organs and tissues ((a), (b) erythrocytes; (c), (d) kidneys; (e), (f) abdominal fat; (g), (h) heart; (i), (j) brain) of spontaneously hypertensive obese rats supplemented with EPA/DHA at different proportions (1:1, 2:1 and 1:2) and soya bean oil as the control diet. Values are means, with their standard errors represented by vertical bars. Mean value was significantly different from that of the control group: *P < 0.05, **P < 0.01, ***P < 0.001 (Kruskal–Wallis and Mann–Whitney U tests). † Mean value was significantly different from that of the 1:1 EPA/DHA group (P<0.05; Kruskal–Wallis and Mann–Whitney U tests).
2.1 group \((P<0.05)\), and in the heart of the 1:2 EPA/DHA group \((P<0.05; \text{Fig. 1})\).

CAT activity was significantly higher \((P<0.001)\) in the kidneys of the 1:1 and 2:1 EPA/DHA groups, and lower in the brain of the 1:1 EPA/DHA group \((P<0.001)\) (Fig. 1). The activity of the glutathione-associated enzymes GR and GPxs was measured in erythrocytes and kidneys, the compartments where SOD and CAT were most affected in comparison with the control group (Fig. 2). In erythrocytes, only GPx activity was detected \((P<0.01)\) in the 1:1 and 2:1 EPA/DHA groups, and in the kidneys, both enzymes were significantly activated \((P<0.01)\) in the 1:1 and 2:1 EPA/DHA groups (Fig. 2). The ratio of the reduced-oxidised form of glutathione \((\text{GSH:GSSG})\) was lower \((P<0.05)\) in the kidneys of the 1:1 and 2:1 EPA/DHA groups (30·94 \((\text{SEM 3·09})\) and 36·73 \((\text{SEM 3·77})\)), respectively) than that found in the kidneys of both the control group (70·96 \((\text{SEM 18·02})\)) and the 1:2 EPA/DHA group (76·70 \((\text{SEM 7·58})\)).

**Discussion**

The present study focused on aspects of the action of \(n\)-3 PUFA in SHROB rats that were not addressed before, specifically their role in modulating OS and the influence of the relative proportions of EPA and DHA on both OS and significant markers of the MetS. The results of the present study indicate that EPA/DHA mixtures at different proportions clearly improve the altered lipid profile in SHROB rats; the three proportions tested (1:1, 2:1 and 1:2) decreased LDL-C concentration and the proportions 1:1 and 2:1 decreased plasma TAG concentration. It has been reported that the effects of \(n\)-3 PUFA on the lipid profile were derived from their capacity to up-regulate the expression of genes encoding proteins involved in fatty acid oxidation, while simultaneously down-regulating genes encoding proteins involved in lipid synthesis\(^{25}\). According to our data, mixtures of EPA and DHA that include more or equal amounts of shorter-chain EPA may be a better option than an excess of DHA for the control of dyslipidaemia in the MetS. A meta-analysis of clinical studies has revealed that both PUFA were similarly effective in decreasing TAG concentration\(^{26}\); however, the results of the SHROB model appeared to differ from the outcome of human studies.

Chronic inflammation is closely linked to obesity\(^{27}\), and it is another relevant risk factor for CVD\(^{28}\). In the present study, we found a significant improvement in plasma CRP concentration in obese SHROB rats fed EPA/DHA, indicative of an amelioration of systemic inflammation. It has been suggested that \(n\)-3 PUFA show anti-inflammatory effects because they cause a partial displacement of the pro-inflammatory \(n\)-6 PUFA pathway and simultaneously generate anti-inflammatory resolvins (from EPA and DHA) and protectins (from DHA)\(^{29,30}\). Also, recent results have shown that \(n\)-3 PUFA inhibit NLRP3 inflammasome activation\(^{31}\). Again, in the results of the present study, those mixtures with more or equal amounts of EPA appear to be more effective than the DHA-rich 1:2 mixture. Indeed, the 1:2 mixture increased the concentration of vascular cell adhesion molecule-1, a marker of further inflammation, as previously reported in healthy WKY rats\(^{9}\). None of the mixtures had any effect on either intercellular adhesion molecule-1 or on the marker of thrombocytic activity plasminogen activator-inhibitor-1. The effects of \(n\)-3 PUFA on adhesion molecules appear to be controversial\(^{32}\).

Regarding glucose metabolism, none of the EPA/DHA mixtures modified postprandial glycaemia, but all of them significantly decreased plasma glycated Hb levels. This is a beneficial effect of \(n\)-3 PUFA that has also been observed in normal WKY rats\(^{9}\). Meanwhile, we observed that only the mixture with a higher proportion of EPA tended to increase insulin secretion. This seems to agree with a previous study in which DHA, but not EPA, prevented the increase in circulating insulin, induced by conjugated linoleic acid\(^{30}\). In contrast, other authors have reported that PUFA do not affect insulin secretion in healthy pigs\(^{34}\). Whereas the overall effect of PUFA on glucose metabolism is still controversial, there is evidence that EPA and DHA may act at different sites and involve different
mechanisms related to insulin resistance (35). Consequently, mixtures at different proportions yield different results.

As OS is another factor that has been closely associated with the MetS (3), we examined the effect of EPA/DHA supplementation on endogenous antioxidant systems and IsoP as the ‘gold’ standard marker of OS (36,37). We found that EPA/DHA at the ratio of 1:1 increases SOD activity in the adipose tissue, erythrocytes, heart and brain of SHROB rats. In a previous study, we described that fat and erythrocytes of SHROB rats exhibited decreased SOD activity compared with WKY rats (14). Herein, we show that EPA/DHA supplementation at the ratio of 1:1 can increase SOD activities to values close to those found in healthy WKY rats. We also previously reported that antioxidant enzyme activities in the heart and brain of SHROB and normal WKY rats were similar (14). In these organs, n-3 PUFA appear to activate this

Fig. 3. Urine isoprostanes ((a) 15-F2t-IsoP, (b) 2,3-dinor-15-F2t-IsoP, (c) 5-F2t-IsoP, (d) 5-F2c-IsoP and (e) 8-F2t-IsoP) in spontaneously hypertensive obese rats supplemented with EPA/DHA at different proportions (1:1, 2:1 and 1:2) and soyabean oil as the control diet. Values are means, with their standard errors represented by vertical bars. *Mean value tended to be different from that of the control group (P<0·1; Kruskal–Wallis and Mann–Whitney U tests). The structures of the respective compounds are also shown. 15-F3t-IsoP was not detected in any of the samples.
protective system in SHROB rats. This activation has also been observed in the erythrocytes of healthy WKY rats supplemented with EPA/DHA at the ratio of 1:1\(^{[9]}\). Moreover, EPA/DHA mixtures also increase the activity of other enzymes, such as CAT and GPx in erythrocytes, with this effect being more marked in rats supplemented with EPA/DHA at the ratio of 2:1. In the kidneys, an organ particularly affected in the present rat model\(^{[12]}\), EPA/DHA supplementation at the ratio of 2:1 was more effective than that at 1:1.

We have also previously reported that endogenous antioxidant systems in the kidneys were not altered in SHROB rats compared with WKY rats\(^{[14]}\). However, in the present study, EPA/DHA at the ratio of 2:1 caused an increase in the activity of different antioxidant enzymes (SOD, CAT, GR and GPx), and a displacement of the glutathione system towards accumulation of GSSG (a decreased GSH/GSSG ratio). All these results suggest that EPA/DHA supplementation triggers some OS with subsequent activation of defence mechanisms. The changes in the markers of OS (plasma LDL-ox and urine IsoP) examined in the present study are consistent with these aforementioned observations.

EPA/DHA mixtures at all the three ratios significantly increased plasma LDL-ox in SHROB rats. A similar increase in LDL-ox after supplementation with n-3 PUFA has previously been described in an animal model of the MetS\(^{[38]}\). These authors hypothesised that because n-3 PUFA are incorporated into cell membranes, they cause an increase in the peroxidability index, i.e. the number of sites susceptible to lipid peroxidation. However, at this point, it should be noted that the PUFA supplemented in the present study did not directly contribute to the circulating lipid peroxides, as demonstrated by the absence of 15-F\(_{2}\)-IsoP, which would have been derived from EPA and DHA. In the apparent contradiction of the results for plasma LDL-ox, the supplemented rats presented lower levels of urine IsoP (Fig. 3), particularly the 1:2 EPA/DHA group. This means that the overall effect of EPA and DHA in the organism, as evidenced by the levels of endogenous lipid oxidation, appears to be antioxidant. This overall antioxidant effect may be caused by the activation of defence systems, which would also agree with our previous observation that plasma antioxidant capacity increases in WKY rats after supplementation with n-3 PUFA\(^{[9]}\).

Taken together, it seems that the results of the present study indicate that EPA is especially relevant to the activation of endogenous antioxidant systems and the decrease in circulating TAG and CRP levels, while DHA is more effective in lowering IsoP. These differential effects may be due to the different effects of EPA and DHA on signalling pathways, or on the expression of certain genes, particularly on those related to the activation of antioxidant enzymes. It has been reported that EPA and DHA differentially influence transduction pathways related to cell growth, thereby affecting different biological processes\(^{[39]}\), the release of inflammatory cytokines by peripheral blood mononuclear cells\(^{[40]}\), and the expression of genes involved in lipid metabolism\(^{[41]}\).

Also, the incorporation of either EPA or DHA into cell membranes has different effects on their structure, as shown for endothelial cells, lymphocytes and in ion channel effects, as observed in several cellular models\(^{[42]}\). Another relevant factor is that EPA and DHA may exhibit specific interactions with other food components such as vitamin E that influence oxidative status, and act together with DHA in the expression of certain genes\(^{[42]}\). Certainly, our observations indicate that the proportions of EPA and DHA should be carefully considered when carrying out studies with n-3 PUFA or when comparing results from different studies.

In short, supplementation of SHROB rats, a model of the MetS, with EPA/DHA mixtures at different proportions significantly lowered plasma total cholesterol and LDL-C concentrations, compared with the control group, given a standard diet. Specific mixtures significantly decreased inflammation (CRP levels, 1:1 and 2:1 ratios), increased the activity of antioxidant enzymes (SOD in erythrocytes and abdominal fat, 1:1 ratio; SOD, CAT, GR, GPx in kidneys, 1:2 ratio), and showed a tendency towards decreasing excretion of urinary IsoP (mainly 1:2 ratio). More LDL-ox was observed in all the supplemented groups. These results are compatible with the stimulation of antioxidant defences by EPA/DHA mixtures, leading to lower OS, together with a reduction in CVD risk factors such as LDL-C and inflammation in SHROB rats.

**Supplementary material**

To view supplementary material for this article, please visit http://dx.doi.org/10.1017/S0007114514004437

**Acknowledgements**

The authors thank Lorena Barros and Mª Jesus Gonzalez for technical support, and Christopher Evans for language revision. The authors also thank AFAMSA (Vigo, Spain) for the provision of the commercial fish oil APAMPES 121 EPA.

The present study work was supported by the Spanish Ministry of Science and Innovation and of Economy and Competitiveness (grant no. AGL2009-12374-C03-01, -02 and -03; AGL2013-49079-C2-1,2-R, respectively). The authors acknowledge the Panamanian Government (SENACYT/IFARHU) and the Spanish Ministry of Science and Innovation for their predoctoral fellowships to E. M.-T. and L. M., respectively. They also acknowledge ISCIII for the postdoctoral contract ‘Sara Borrell’ to J. P.-J. (CD09/00068). None of the funders had any role in the design, analysis and findings of the study or in the writing of this article.

The contributions of the authors are as follows: I. M. and J. L. T. designed the research; E. M.-T., J. P.-J., M. R., M. G., N. T., M. M.-C., O. J. and L. M. carried out the experimental work; E. M.-T., J. P.-J., S. R.-R. and J. L. T. analysed the data; E. M.-T., J. P.-J. and S. R.-R. wrote the first draft of the manuscript; J. P.-J. and J. L. T. had primary responsibility for the final content. All the authors read and approved the final version of the manuscript.

The authors declare there are no conflicts of interests.
References

33. Vemuri M, Kelley DS, Mackey BD, et al. (2007) Docosahexaenoic acid (DHA) but not eicosapentaenoic acid (EPA) prevents trans-10, cis-12 conjugated linoleic acid (CLA)-induced insulin resistance in mice. Metab Syndr Relat Disord 5, 315–322.


