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The effect of dietary fish oil-supplementation to healthy young men on oxidative burst measured by whole blood chemiluminescence

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Dietary long-chain n-3 PUFA (n-3 LCPUFA) are thought to have immune-modulating effects, but the specific effects and mechanisms are not fully elucidated. The aim of this study was to determine whether dietary n-3 LCPUFA could affect ex vivo oxidative burst in healthy young men. The study had a randomised 2×2-factorial design in which subjects were randomly assigned to 8-week supplementation with capsules containing fish oil (about 2.9 g n-3 LCPUFA/d) or olive oil (control). Subjects were also randomly assigned to household use of oils and fat spreads with a high or a low 18:2n-6 content. At baseline and at the end of the intervention, the fatty acid composition of peripheral blood mononuclear cells (PBMC) was analysed by GLC and oxidative burst was studied in whole blood stimulated with zymosan using luminol-enhanced chemiluminescence. The PBMC content of n-3 LCPUFA was markedly increased by the fish oil-supplementation (P<0.001, compared to the olive oil groups). No effect of the intervention was observed on neutrophil count, but one measure of the zymosan-induced oxidative burst was higher in the fish oil groups (P=0.03) compared to the olive oil groups. The fat intervention did not in itself affect oxidative burst neither did it change the effect of the fish-oil intervention. The measures of oxidative burst at the end of the intervention period were found to be associated with the DHA content of PBMC (r 0.44, P=0.016), suggesting a dose–response relationship. These results indicate that n-3 LCPUFA may have immuno-stimulating effects.

Respiratory burst: Dietary supplementation: Inflammation: Human intervention study

Dietary long-chain (≥C20) n-3 PUFA (n-3 LCPUFA) are generally considered to be immuno-suppressive¹ and have been shown to favourably affect a number of inflammatory diseases including rheumatoid arthritis (²⁵) and CVD (³⁻⁴). Neutrophils are central players in inflammatory diseases (⁵) as they form part of the innate immune response, which acts in the early host defence. Neutrophils kill microbes by releasing proteolytic enzymes, superoxide and related reactive oxygen species (ROS), a process referred to as oxidative burst. ROS are toxic to the surrounding tissue, and the damage provoked intensifies neutrophil activation, recruits more neutrophils and thereby increases the inflammatory response (⁶).

Feeding fish oil (FO), which contains n-3 LCPUFA, primarily DHA and EPA, to laboratory animals has been shown to decrease ex vivo neutrophil superoxide production measured in isolated neutrophils (⁷⁻⁸). A number of ex vivo studies have investigated the influence of FO supplements on oxidative burst in healthy human volunteers, but the results are conflicting. Some have found suppressive effects of FO⁹⁻¹² whereas others have not¹³⁻¹⁷.

The mechanisms by which n-3 LCPUFA exert their immune-modulating actions are unknown. From a classical point of view, the immuno-suppressive effect is thought to operate through changes in eicosanoid production. It has, however, also been suggested that n-3 LCPUFA mediate their effect by changes in gene expression or membrane structure (¹⁸⁻¹⁹). The NADPH oxidase, which is responsible for oxidative burst, is a large membrane-associated protein complex that assembles in lipid rafts (²⁰). Lipid rafts are generated by separation in the membrane bilayer of cholesterol and phospholipids containing unsaturated fatty acids, especially DHA (²¹). It is therefore plausible that the activity of the NADPH oxidase could be affected by incorporation of DHA into cellular membranes.

A common characteristic of many of the previous studies investigating the effects of n-3 LCPUFA on oxidative burst is that the oxidative burst assays were performed on isolated

Abbreviations: ANCOVA, analysis of covariance; AUC, area under the curve; Dex, desensitisation; FA%, fatty acid area percentage; FO, fish oil; n-3 LCPUFA, long-chain n-3 PUFA; OO, olive oil; PBMC, peripheral blood mononuclear cell; R/K, rapeseed oil/Kaergaarden; ROS, reactive oxygen species; S/B, sunflower oil/Becel; T²P, time to half peak height; α₀⁻₆₀, slope between 40 and 60% of peak height; Δ, changes in (endpoint value – baseline value).

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The present study was part of a larger intervention study investigating the effects of FO-supplementation in combination with a high or low 18:2n-6 intake on tissue PUFA-incorporation\(^{(28)}\), CVD risk markers and cytokine production in healthy young men. The study ran from October 2005 to February 2006. Healthy male subjects aged 18–40 years were recruited by notes at universities throughout Copenhagen. Subjects were invited to participate in the study if they matched the following inclusion criteria: BMI 18.5–27 kg/m\(^2\); non-smokers or infrequent smokers (≤ 5 cigarettes a week); daily consumption of butter, margarine and/or oil, and home-made cooking ≥5 times per week. Volunteers were excluded if they suffered from chronic diseases or serious allergic symptoms, were taking medication or dietary supplements, exercised more than 7 h per week or had donated blood within the last 2 months. Ethical permission for the study was obtained from the Ethical Committee of the Municipalities of Frederiksberg and Copenhagen (Journal No.KF 01 267804). The study was registered in the clinical trial database of the US National Institutes of Health (ClinicalTrials.gov, NCT00266292). Informed written consent was obtained from all subjects enrolled in the study.

The design of the study was as follows: a screening visit, a 2-week run-in period, a baseline visit, an 8-week intervention period, an endpoint visit and a washout visit 8 weeks post-intervention\(^{(28)}\). The sub-study described in the present paper ended after the subjects had completed the endpoint visit and therefore does not contain any washout data. The following sections describe only measurements relevant to this particular sub-study.

The study had a randomised-parallel 2 × 2-factorial design in which the subjects were randomly assigned to daily supplementation with capsules containing either FO or olive oil (OO, control) for 8 weeks. Subjects were also randomly assigned to household use of oil and fat spreads with either a high or a low 18:2n-6 content supplied by sunflower oil/Bece1 (S/B) and rapeseed oil/Kaergaarden (R/K), respectively. The randomisations were performed in two steps: first, notes were drawn from one envelope for the capsule intervention and second, within each capsule group notes were drawn for the oil and fat spread intervention. Thus, the subjects were randomly allocated in a double-blind fashion to one of four intervention groups: (1) OO-capsules and S/B, (2) OO-capsules and R/K, (3) FO-capsules and S/B or (4) FO-capsules and R/K.

Capsules (in a defined surplus), fat spreads and oils supplied by the randomisation were supplied to the subjects on the day of their baseline visit. The subjects were instructed to consume ten capsules per day (equal to 5 ml/d) with FO (Bio-Marin) or OO (both kindly provided by Pharma Nord, Vejle, Denmark). Each FO-capsule contained 333 mg n-3 PUFA (NEFA, approximately 166 mg EPA and 119 mg DHA). The OO was given as TAG, but the capsules were matched for fat content. The S/B groups were supplied with sunflower oil kindly provided by Aarhus United Denmark A/S, Aarhus, Denmark) and margarine (Becel 60, kindly provided by Unilever Denmark A/S Foods, Brøndby, Denmark) with a high content of 18:2n-6, whereas the R/K groups were supplied with rapeseed oil (also a gift from Aarhus United Denmark A/S) and a butter product containing two-thirds butter and one-third rapeseed oil (Kaergaarden Light, kindly provided by Arla Foods amba, Viby, Denmark) with a low content of 18:2n-6. Both the margarine and the butter products contained 60 g fat/100 g product.

Prior to the intervention period, all subjects were provided with butter (Lurpak, also a gift from Arla Foods amba) and OO (Oleificio R.M. s.p.a., Lucca, Italy) for a 2-week run-in period. The purpose of this was to reset the fatty acid composition of the tissue marker. There were no restrictions concerning the amount of fat spreads or oil ingested, but it was intended that the intake of fat and energy remained constant during the study. The average energy intake and the dietary composition were determined by 4d weighed dietary records: (1) prior to the run-in period and (2) at the end of the 8-week intervention period. The mean length of the intervention was 56 d (range 50–65 d).

Seventy-five subjects were invited to a screening visit at which height, weight, hip- and waist-circumference were measured and the subjects were instructed in how to perform the dietary records. Eight subjects dropped out during the run-in period due to the workload of attendance and one subject was rejected at the baseline visit due to intense discomfort during blood sampling. This left us with sixty-six men enrolled in the study, of which sixty-four completed the intervention period. The reasons for drop-out were discomfort swallowing the oil capsules (n 1) and the workload of attendance (n 1). Furthermore, six subjects were left out from this sub-study, four because no endpoint oxidative bursts were measured and two because the baseline neutrophil count failed (equipment error). Drop-outs and data collection failures were evenly distributed across the four intervention groups. Data are reported only for those subjects with a complete dataset (n 58). Furthermore, one analysis run stopped (for an unknown reason), which resulted in a loss of three subjects in all statistical tests of the area under the curve (AUC) and desensitisation (Des) oxidative burst variables (n 55). The baseline characteristics of the subjects in
each of the intervention groups are shown in Table 1. Age, height, weight, BMI and waist/hip ratio did not differ
between the groups.

Fasting blood samples were collected at the baseline and
endpoint visits. The subjects were asked to follow standar-
dised fasting conditions before each visit. These were: no
food for 12 h or more (except from 0·5 litre water), lack of
strenuous physical activity for 36 h, no smoking for 1 week
and no medicine for 24 h. Moreover, the subjects were told
to eat the same meal the evening before all visits and to re-
schedule their visit if they were ill or had a cold. These criteria
were checked at every visit.

Experimental procedures

On the days of examination, body weight and waist/hip ratio
were determined. The subject filled out a questionnaire on
dietary habits, illnesses, use of medication and smoking.
Blood (80–110 ml) was drawn from fasting males after
10 min of rest and analysed for: oxidative burst, cell count,
Hb concentration and the fatty acid composition of peripheral
blood mononuclear cells (PBMC).

Compliance was assessed by counting returned capsules
and measuring tissue fatty acid composition of the PBMC.
The mean daily consumption of capsule oils was 4·5 (SD 0·5)
ml/d (range 3·0–5·5 ml/d), equivalent to 89 % compliance. No
difference in compliance was observed between the interven-
tion groups. The fatty acid composition in the total lipid frac-
tion from PBMC was determined as a biomarker of the
resulting changes in tissue composition.

Measuring fatty acid composition of the tissue

PBMC were isolated from blood sampled with Na-heparin
and fatty acid composition was determined by GLC as described
elsewhere. Fatty acids are given as area percentage (FA%), comparable to g/100 g.

Table 1. Baseline characteristics of the subjects in the four intervention groups (n 58)

<table>
<thead>
<tr>
<th></th>
<th>Olive oil capsules</th>
<th>Fish oil capsules</th>
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<tbody>
<tr>
<td></td>
<td>S/B (n 14)</td>
<td>R/K (n 17)</td>
</tr>
<tr>
<td>Age (years)</td>
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<tr>
<td>Median</td>
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<td>25</td>
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<tr>
<td>Height (m)</td>
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</tr>
<tr>
<td>Mean</td>
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<td>1·83</td>
</tr>
<tr>
<td>SD</td>
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<td>0·07</td>
</tr>
<tr>
<td>Weight (kg)</td>
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<td></td>
</tr>
<tr>
<td>Median</td>
<td>74·4</td>
<td>72·9</td>
</tr>
<tr>
<td>Percentiles‡</td>
<td>69·6–79·0</td>
<td>65·8–81·3</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
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<td></td>
</tr>
<tr>
<td>Mean</td>
<td>3·1</td>
<td>2·2</td>
</tr>
<tr>
<td>SD</td>
<td>0·85</td>
<td>0·84</td>
</tr>
<tr>
<td>Waist/hip ratio</td>
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<td></td>
</tr>
<tr>
<td>Median</td>
<td>0·81–0·89</td>
<td>0·83–0·87</td>
</tr>
<tr>
<td>Percentiles‡</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

R/K, rapeseed oil/Kaergaarden; S/B, sunflower oil/Becel.
† Between-group comparisons were performed using one-way ANOVA and Kruskal–Wallis tests with significance established at
P<0·05. No significant differences were observed for any of the variables.
‡ Percentiles: 25th to 75th.

Neutrophil counts and Hb concentration

Neutrophil count was determined by leucocyte size diff-
erentiation carried out on a Sysmex KX-21 automated
haematology analyser (Sysmex Corporation, Kobe, Japan)
on venous blood collected in EDTA tubes using the appropri-
ate control. At the same time the Hb concentration was
measured. Intra- and interassay CV for Hb were 0·7 %
(n 12) and 0·6 % (n 25), respectively. Prior to the study
intra-individual variation in neutrophil count was determined
to 17·5 % measured on three independent days in five
individuals.

Measurement of oxidative burst

Prior to the study all stock solutions were portioned and frozen
at –20°C. Zymosan A, a proteoglycan isolated from Sacchar-
omycetes cerevisiae (Sigma-Aldrich, St Louis, MO, USA), was
suspended to 1 mg/ml in Hank’s Balanced Salt Solution with
CaCl₂ + MgCl₂ (Invitrogen, Carlsbad, CA, USA) and por-
tioned in aliquots of 300 µl, and luminol (5-amino-2,3-dihy-
dro-1,4-phthalazinedione; Across Organics, NJ, USA) was
dissolved to 5 mg/ml in dimethyl sulphoxide (Sigma-Aldrich)
and portioned in aliquots of 100 µl.

Blood, sampled in Na-heparin tubes for assessment of oxi-
dative burst, was stored at 37°C and oxidative burst was
determined within 2 h after sampling by chemiluminescence:
90 ml whole blood diluted 1:9 in pre-warmed (37°C) Hank’s
Balanced Salt Solution was loaded on ninety-six-well Lumi-
Nunc polystyrene plates (Nunc, Roskilde, Denmark), which
were preloaded with a 1:10 diluted stock solution of luminol
(10 µl) with Hank’s Balanced Salt Solution in a final concen-
tration of 142 µM/well. The diluted blood samples were
stimulated with 100 µl pre-warmed (37°C) zymosan at two
different final concentrations (5 and 0·5 µg/ml). Chemilumi-
nescence (relative light units/s) was measured in a thermo-
stat-regulated (37°C) ninety-six-well Orion II Microplate
luminometer (Berthold Detection Systems, Pforzheim, Germany). The microplate was read for 1 s/well with a 2 min delay, a cycle repeated for a total of seventy-five rounds. Unstimulated control samples (blanks) were analysed in duplicate and stimulated samples in sextuplicate.

Five variables were used to define the kinetics of the mean chemiluminescence response curve (Fig. 1): (1) time to half peak (T_{2P}), (2) peak height, (3) AUC, (4) slope between 40 and 60% of peak height (\(a_{40-60\%}\)) and (5) Des. All variables, except for AUC were calculated with GraphPad Prism version 4.00 (GraphPad USA) based on linear mathematical principles. AUC was determined with GraphPad Prism version 4.00 (GraphPad Prism, San Diego, CA, USA) based on integral calculation. Prior to the study, the inter-day variation of the oxidative burst variables was determined: blanks 2.4%, peak height 25.9%, AUC 29.6%, T_{2P} 3.8%, Des 48.5% and \(a_{40-60\%}\) 35.4% based on three independent days in five individuals. Data were analysed with the Statistical Package for the Social Sciences software version 13.0 (SPSS Inc., Chicago, IL, USA). Data are only shown for the 5 mg/ml zymosan-stimulated samples. The results of the 0.5 mg/ml zymosan-stimulated samples were similar, but the responses were smaller.

Statistical analyses

Data were checked for Gaussian distribution with Shapiro–Wilks test and visual inspection of histograms. Gaussian-distributed data are presented as means with their standard errors, whereas non-Gaussian-distributed data are presented as medians with 25th–75th percentiles. Baseline comparison between the four intervention groups were performed by one-way ANOVA and Kruskal–Wallis tests. Analyses of covariance (ANCOVA) were used for endpoint comparisons. The three statistical procedures were as follows:

1. The ANCOVA for the endpoint comparisons of PBMC fatty acid content (EPA, 22 : 5n-3, DHA and total n-3 PUFA) included fat type and capsule type as fixed factors and were adjusted for baseline values. All models were checked for interaction between the capsule and the oil/fat spread interventions.

2. All oxidative burst variables at endpoint were checked for bivariate Spearman’s correlation with the following covariates: neutrophil count (endpoint), changes in (\(\Delta\), endpoint value - baseline value) neutrophil count, smoking (yes/no) and \(\Delta\) Hb concentration. Correlating parameters (neutrophil count and \(\Delta\) neutrophil count) were tested as covariates in the ANCOVA endpoint comparison of the oxidative burst variables, which also included fat type and capsule type as fixed factors (including check for fat x capsule interaction) and adjustment for baseline values. Covariates were successively removed, but those affecting the outcome (\(P<0.05\)) were kept in the model (\(\Delta\) neutrophil count).

3. The dose–response relationship analysis of the oxidative burst variables were performed using multiple linear regression analysis including the following covariates: endpoint PBMC DHA content, baseline values and the covariates included in the oxidative burst variable ANCOVA.

Wilcoxon test and paired-sample t test were used to check for within-group changes during the intervention of non-Gaussian-distributed and Gaussian-distributed data, respectively. Associations between the estimated FO-consumption (from the count of returned capsules (ml/d)) and \(\Delta\) PBMC content of total n-3 PUFA, DHA, 22 : 5n-3 and EPA was analysed using Pearson correlation analysis. Significance was established at \(P<0.05\).

Results

The subjects had a mean energy intake of 12 531 (SD 2742) kJ/d (range 6709–18 942 kJ/d), of which fat constituted 32.4 (SD 5.4) % (range 20.0–43.0 %) of energy. No between-group differences in energy consumption or macronutrient distribution were observed at baseline or during the intervention(28). All subjects maintained a constant weight during the study.

Fatty acid composition of mononuclear cells

The effect of the interventions on PBMC fatty acid composition has been published recently(28). In short, no differences were observed in the PBMC content of total n-3 PUFA, DHA, 22 : 5n-3 or EPA between the treatment groups at baseline, after 2 weeks of run-in where oil and fat spreads were standardised. Endpoint comparison showed that 8 weeks of FO-supplementation increased total n-3 PUFA, DHA, 22 : 5n-3 and...
EPA; all of which were significantly higher (P<0·001, for all) in the FO groups. The effects were supported by paired-sample t-tests (P<0·001 for all fatty acid categories in both FO groups). The best FO compliance marker was EPA, which increased in the FO groups from 0·52 (SE 0·02) to 2·73 (SE 0·13) FA% in the PBMC compared to no changes (0·44 (SE 0·02) to 0·39 (SE 0·02) FA%) in the OO groups. The estimated FO-capsule consumption (count of returned FO-capsules (ml/d)) correlated with endpoint PBMC EPA content (r 0·51, P=0·006), total n-3 PUFA content (r 0·48, P=0·011) and the DHA content (r 0·38, P=0·049).

Neutrophils
The neutrophil counts at baseline did not differ between the four intervention groups. The median count was 2·4×10⁹ cells/l (25th–75th percentiles 2·0–3·2×10⁹ cells/l), ranging from 1·2 to 4·3×10⁹ cells/l. The interventions had no effect on the cell counts. In general, the neutrophil count of each subject was relatively constant (mean change of 0·66 (SD 0·88)×10⁹ cells/l), but in five of the subjects the cell count varied greatly (≥1·8×10⁹ cells/l) between the baseline and endpoint visit.

Oxidative burst
There were no differences between the four groups for any of the oxidative burst variables at baseline. Four of the five oxidative burst variables (with the exception of T₃P) correlated with one another at baseline (0·89 < r < 0·99, P<0·001) and after 8 weeks of intervention (0·35 < r < 0·52, P≤0·0007). Bivariate correlations were observed between four of the five oxidative burst endpoint variables (with the exception of T₃P; P=0·23) and endpoint neutrophil count (0·35 < r < 0·52, P=0·0007). The oxidative burst endpoint variables were checked for correlation with confounders, such as Hb concentration (possible quenching of emitted light) and if the subject smoked (yes/no). No relations were observed, except for a correlation between Δ Hb concentration and T₃P (r 0·29, P=0·03).

Eight weeks of capsule intervention were found to have a significant effect on one of the five oxidative burst variables, Des (P=0·03; Table 2). No effects of the fat intervention or of the capsules × fat interaction term were observed (data not shown). Two outliers were observed in the ANCOVA model for peak height, AUC and Des (Fig. 2). After exclusion of these two subjects, Des (P=0·01), peak height (P=0·02) and AUC (P=0·05) were significantly increased in the FO-supplemented groups.

Dose–response relationship
Endpoint comparison of the oxidative burst variables correlated significantly with the DHA content in the PBMC at the end of the intervention (r 0·44, P=0·016 for Des and AUC shown in Fig. 3). As for the capsule effect on oxidative burst, the two subjects were also outliers in the dose–response relationship regression models (for peak height, AUC and Des). Exclusion of the two subjects in the analyses decreased the P value for the dose–response relationships (Des (P=0·010), peak height (P=0·047) and AUC (P=0·02)). Similarly, a significant dose–response relationship was observed for EPA (Des (P=0·029) and peak height (P=0·047)), but to a lesser extent than the one observed for DHA.

Discussion
The present results indicate that FO may increase oxidative burst after 8 weeks of intervention, as one of five variables was significantly increased in relation to the control supplement. This was supported by outlier analysis and a dose–response relationship between the oxidative burst and the DHA content of immune cells (here PBMC). The effect was most evident for the oxidative burst variables: peak height, AUC and Des, but the oxidative burst variables were highly inter-correlated. The present result is in agreement with a recent study that found enhanced oxidative burst in healthy young men after 8 weeks of FO-supplementation[29]. However, both these studies contradict the results from previous similar studies that have found either no effects[10,13–17] or a reduction in the response[11,12]. The two early studies that observed a reduction in oxidative burst[11,12] were small (n 6 and n 22), had no control group, and gave little information on inclusion and exclusion criteria and participant characteristics (e.g. the mean age). In general, the more recent and larger studies[10,13–17] have all reported no effect of n-3 LCPUFA-supplementation on oxidative burst in healthy young men. All of these, except one[13], were randomised, placebo-controlled and double-blinded like the present study. The FO-dose used in the present study (about 2·9 g n-3 LCPUFA/d) was within the range provided in the previous studies[10–12,14]. We had only minor drop-outs and compliance was good as judged from both capsule count and PBMC fatty acid analysis. In agreement with results from a previous study[17], no effect was observed on neutrophil counts. Most of the subjects had very small changes in neutrophil count (comparable to the intra-individual variation measured in our pre-study), but a few varied more. Therefore, all analyses were adjusted for the changes in neutrophil count.

A study by Rees et al.[10] suggested that the effect of FO-supplementation may be age-related, since they found an immuno-suppressive effect on oxidative burst in a group of older subjects (aged 55–73 years), whereas no effect was observed in a group of younger subjects (aged 18–52 years). However, another study found no effect of FO-supplementation on oxidative burst in subjects aged 55–74 years[17]. The most striking difference between these two studies was the type of oils that they used. Thies et al.[17] used five different oils including a pure DHA oil (700 mg/d) and a FO that supplied 720 mg EPA/d and 280 mg DHA/d, whereas Rees et al.[10] used a pure EPA oil in four different doses. Thus, it might be of interest to investigate possible differences in the effects of EPA, DHA or a combination of the two on oxidative burst in elderly subjects. The opposing effects on the immune response in older versus young men might be related to health status. It has been shown that FO lowers cardiovascular risk markers, such as blood pressure[30] and heart rate[31], primarily in high-risk subjects, e.g. elderly or those with high baseline values. Whether the immuno-stimulating effect of FO in the present study has any long-term health effects is up for further investigation. Increased
Table 2. Oxidative burst variables at baseline and endpoint in the four intervention groups (n 58)

<table>
<thead>
<tr>
<th></th>
<th>Olive oil</th>
<th></th>
<th>Fish oil</th>
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<tr>
<td></td>
<td>S/B (n 14)</td>
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<td>S/B (n 15)</td>
<td>R/K (n 12)</td>
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<td>32</td>
<td>34</td>
<td>32</td>
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<tr>
<td>SD</td>
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<tr>
<td>Median</td>
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<td>1104*</td>
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<td>1155</td>
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<tr>
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<td>857–2005</td>
<td>768–2194</td>
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<td>AUC‡</td>
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<tr>
<td>Median</td>
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<td>300–1000</td>
<td>284–912</td>
<td>0·12</td>
</tr>
<tr>
<td>α_{40–60} (RLU/s per min)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>70·9</td>
<td>49·8*</td>
<td>60·1</td>
<td>57·4</td>
<td>0·10</td>
</tr>
<tr>
<td>Percentiles‡</td>
<td>56·0–123·0</td>
<td>27·6–89·3</td>
<td>27·2–114·3</td>
<td>25·0–103·5</td>
<td>48·0–81·8</td>
</tr>
</tbody>
</table>
| AUC, area under the curve; blank, non-stimulated control samples; Des, desensitisation; peak, peak height; RLU, relative light units; R/K, rapeseed oil/Kaergaarden; S/B, sunflower oil/Becel; TJP, time to half peak; α_{40–60}, slope between 40 and 60 % of peak height.
*Values were significantly different from those of the baseline (paired-sample t-test or Wilcoxon test): peak, P = 0·016; AUC, P = 0·016; Des, P = 0·026; α_{40–60}, P = 0·013.
† Percentiles: 25th to 75th.
‡ Analysis stop (unknown reason), values for three subjects were lost leaving the groups with n 14, 14, 15 and 12 subjects, respectively for the AUC and Des analyses (n 55).
§ Endpoint between-group comparisons were performed using analysis of covariance with fats and capsules as fixed factors, adjusting for baseline values and changes in neutrophil count (endpoint − baseline).
oxidative burst may have beneficial anti-bactericidal effects as shown in mice(32).

The inconsistent results, concerning the effect of n-3 LCPUFA on oxidative burst, may to some extent be due to the method of assessment. The more recent studies have used fluorescence-activated cell-sorting analyses for detection of ROS in isolated neutrophils(29) or whole blood(10,15,17). The earlier studies have used isolated neutrophils assayed by either luminol-enhanced chemiluminescence(11,12,16) or cytochrome c reduction(12 – 14). The isolation procedure has a mechanical impact on the neutrophils and may change the structure and function of the cells. All the methods rely on chemical trapping of ROS, which due to the extreme reactivity of superoxide is rapidly converted to other ROS. Some ROS are able to cross membranes, which makes it difficult to distinguish between intracellularly and extracellularly produced ROS.

Luminol-enhanced chemiluminescence is thought to measure intracellular as well as extracellular ROS, whereas the fluorescence-activated cell-sorting analyses only detect the intracellular ROS. More methods could be used in parallel in order to enhance the validity of the results. To our knowledge, this is the first study to use chemiluminescence to study the effect of FO-supplementation on oxidative burst ex vivo in whole blood. The advantage of the whole blood assay is that the neutrophils are kept in their physiological environment. We handled the blood for the assay as gently as possible throughout the entire procedure and performed the assay at 37°C. Furthermore, we used a sub-maximal stimulus (a zymosan–neutrophil ratio of 2·4) to induce an oxidative burst, which may also be more physiologically relevant than the use of surplus stimulus to ensure a 100 % cell activation as is done in most other studies(10,15,17).

We used OO as capsule control because it has a low content of PUFA, especially n-3 PUFA, and because 18 : 1 n-9 was regarded as a neutral fatty acid. The use of an OO control in studies of immune function has been questioned since some studies have reported effects on immune function after OO-supplementation(33,34). The observed effect of the capsule intervention appears to be due to a reduction in the oxidative burst, especially when assessed as Des, in the OO groups rather than an increase in the FO groups. However, due to the observed correlation between the oxidative burst effect and the PBMC content of DHA we find it unlikely that the effect is driven by the OO. The observed dose–response correlation was not so strong (r 0·27, P=0·04), but since this was interfered with by inter-day variation (biological as well as methodological, which for AUC amounted to 29·6 %) it may in reality represent a stronger true biological dose–response relationship. A dietary PUFA-induced increased membrane DHA content in Jurkat T-cells was shown to be paralleled by a decrease in MUFA(35). No inverse association between DHA and MUFA was observed in the present study (data not shown) and it is therefore not likely that the correlation between DHA and oxidative burst could reflect an indirect association with 18 : 1n-9.

In general, changes in the fatty acid composition of all immune cells are related to the dose(10,15) and the length of
intervention\(^{(26)}\). The DHA content of the PBMC was significantly increased after 8 weeks of FO-supplementation as expected\(^{(26)}\) and parallel changes have most likely also occurred in the neutrophils, as observed by Gibney & Hunter\(^{(26)}\). The active NADPH oxidase-complex has been shown to be located in membrane domains, which are rich in saturated acyl groups, and the so-called lipid rafts\(^{(20)}\). Lipid rafts are formed by segregation of cholesterol and sphingolipids from DHA-containing phospholipids\(^{(21)}\). It was therefore our baseline assumption, although we did not test this, that increased incorporation of DHA could lower the lipid raft area and have functional consequences equivalent to that of cholesterol extraction. Cholesterol depletion of HL-60 cells (a neutrophil-like cell line) has been shown to result in a dramatic reduction in NADPH oxidase activity\(^{(20)}\). Similar effects were observed in a number of other raft-associated proteins in Jurkat T cells and isolated human neutrophils\(^{(15,37)}\). Furthermore, cholesterol depletion has been linked to a decrease in raft-associated protein trafficking\(^{(37)}\), and thus may affect the assembly of the NADPH oxidase complex. An ex vivo study of PUFA-supplemented Jurkat T cells showed a selective modification of unsaturation in the cytoplasmic layer of the detergent-resistant membrane domains (lipid rafts) and an associated inhibition of T cell signal transduction\(^{(35)}\). We did not determine whether the increased intake of n-3 LCPUFA had any effect on the microdomain structure and, to our knowledge, this has not been done in any previous studies. The observed effect of the capsule intervention corresponds poorly with this baseline assumption. The effect of the FO supplement on neutrophil function may therefore be explained by other mechanisms. Several effects of n-3 LCPUFA in other cell types (monocyte, B lymphocyte and T lymphocyte) are thought to be exerted via alterations in gene expression and transcription factors such as PPAR\(^{(28,39)}\), Gorjão et al.\(^{(29)}\) showed modification of seventy-seven genes in human lymphocytes after FO-supplementation, most of which were related to signaling pathways. According to the classical hypotheses, the immuno-suppressive effect of n-3 LCPUFA is thought to be caused by changes in eicosanoid production (e.g. modulation of the LT\(_B_4\) levels\(^{(46)}\)). However, blocking of eicosanoid production by a cyclo-oxygenase inhibitor (indomethacin) has been shown not to affect oxidative burst\(^{(25)}\).

In conclusion, the present study suggests an immuno-stimulating effect on oxidative burst after 8 weeks of FO-supplementation in healthy young men. The effect seems to be related to the cellular DHA content, but the mechanism remains to be determined.

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References


