The effect of EPA and DHA on metabolic syndrome patients: a systematic review of randomised controlled trials

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Abstract
Metabolic syndrome (MS) is characterised by accumulation of CVD risk factors. The use of very long chain n-3 polyunsaturated fatty acids (VLC n3 PUFA) could potentially benefit MS by reducing risk factors. To better understand the possible VLC n3 PUFA benefits, the literature was systematically reviewed for randomised controlled trials (RCT) that published effects of VLC n3 PUFA on MS patients. 17 RCT fulfilled the inclusion criteria and were analysed for relevance to the research question. The available RCT convincingly show that the administration of VLC n3 PUFA doses > 1 g for at least 3 months produces a significant reduction of triglycerides ranging from 7% to 25%. These results confirm the hypotriglyceridemic effect of VLC n3 PUFA in MS patients. The triglyceride lowering may produce further benefits by reducing the % of pro-atherogenic small dense LDL particles (sdLDL) and also perhaps by ameliorating the inflammatory process associated with MS. High doses of VLC n3 PUFA (≥3 g/day) may produce further TAG reductions but could raise other risk factors such as LDL-C. No clear effects were found on other MS markers. The combination of VLC n3 PUFA plus a statin may be useful to prevent the occurrence of coronary events. More studies are needed using different amounts of VLC n3 PUFA, time lengths, dietary backgrounds and different profiles of MS patients before clear recommendations can be made.

Key words: systematic review: metabolic syndrome: very long chain n3 fatty acids: cardiovascular disease

Introduction
Metabolic syndrome (MS), also named syndrome X or metabolic syndrome X, is defined by the National Cholesterol Education Program—Adult Treatment Panel III as the presence of at least 3 cardiovascular disease (CVD) risk factors from a list of the following 5: abdominal obesity (waist circumference > 102 cm for males or > 88 cm for females), impaired fasting glucose (≥6.1 mmol/L), elevated fasting triacylglycerides (TAG ≥1.7 mmol/L), decreased high-density lipoprotein cholesterol (HDLC, < 1 mmol/L for males and < 1.3 mmol/L for females), and high blood pressure (BP ≥130/85 mm Hg) (3). The last two decades, the number of people with MS has increased considerably. This increase is directly associated with the global epidemic of obesity and diabetes (2). MS affects approximately 10–25% of adults worldwide, in some countries and age groups the percentage can be as high as 50–60% (4). MS increases the risk of suffering from type 2 diabetes and CVD by 5- and 2-fold, respectively (4). There is a wealth of evidence from epidemiologic and clinical studies suggesting that modifications in dietary fat composition affect the risk of CVD. Although diet is not specifically identified as a risk factor for MS, there is good evidence to suggest that hypercaloric–hyperlipidic diets, particularly rich in SFA promote obesity, insulin resistance and MS (5–7). Very long chain n-3 polyunsaturated fatty acids (VLC n3 PUFA) in the diet (namely EPA and DHA) have a positive effect on blood lipid levels. The pleiotropic, cardioprotective effects VLC n3 PUFA have been extensively reported. They specifically reduce plasma TAG levels and thereby the risk of CVD. Other potential mechanisms of cardiovascular protection may include lowering of BP, reduced thrombotic tendency, antiinflammatory and antiarrhythmic effects, improved vascular endothelial function, increased plaque stability, increased paraoxonase levels and improved insulin sensitivity (8).

Abbreviations: AHA, American Heart Association; BP, blood pressure; CAD, coronary artery disease; CAVI, cardio-ankle vascular index; CETP, cholesteryl ester transfer protein; CRP, C-reactive protein; CV, cardiovascular; EFSA, European Food Safety Agency; FA, fatty acids; HDLC, high-density lipoprotein cholesterol; HMUFA, high monounsaturated fatty acids; HOSO, high oleic sunflower oil; HSFA, high saturated fatty acids; Hsp27, heat shock protein 27; IMA, Ischemia Modified Albumin; VLC n3 PUFA, very long chain n-3 polyunsaturated fatty acids; LDL-C, low-density lipoprotein cholesterol; PFHCC, low fat high complex carbohydrate; LFHCC n3, low fat high complex carbohydrate supplemented with VLC n3 PUFA; MS, metabolic syndrome; PAI-1, plasminogen activator inhibitor-1; PWV, pulse wave velocity; TC, total cholesterol; t-PA, tissue plasminogen activator; TRL, triacylglycerol rich lipoproteins; RCT, Randomized Controlled Trial; SAA-LDL, Serum Amyloid A–LDL; sdLDL, small dense LDL.

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As a consequence, health authorities and scientific associations have made recommendations of VLC n3 PUFA intake, like the European Food Safety Authority (EFSA) that recommends 250 mg/day\(^9\) for the general population. Only recently, WHO has established new intake recommendations for adults of 250 mg of EPA + DHA per day and 300 mg of EPA + DHA for pregnant or lactating women (of which 200 mg should be DHA)\(^10\). For hypertriglyceridemic subjects, the American Heart Association (AHA) recommends: 2–4 g EPA + DHA/day\(^11\), and for patients with CVD, the AHA recommends 1 g/day\(^11\) to maintain cardiac function, reduce TAG levels and the risk of CHD.

Therapeutic lifestyle modification to manage risk factors is the first-line therapy for MS patients who are not at high risk of suffering a CVD event\(^5\). However, from a clinical perspective, it is difficult to define the optimum diet with which to treat MS. The use of VLC n3 PUFA could potentially benefit MS by reducing risk factors. However, not many studies have investigated the effects of EPA + DHA in this type of patients and there are no recommendations available. To better understand the possible VLC n3 PUFA benefits, the literature was systematically reviewed for randomised controlled trials (RCT) that published effects of VLC n3 PUFA in MS patients.

Materials and Methods

Literature search

The research question applied to the systematic review was “what are the effects of VLC n3 PUFA in MS patients?”. The VLC n3 PUFA of particular interest with respect to MS included EPA and DHA from fish oils or their purified forms. The review included English, Spanish, French, Italian, Portuguese and German articles, without limits on time frame or country, published before April 2011. We searched for publications using the electronic databases MEDLINE and ISI Web of Science. The MeSH terms used in the general search and the search strings used were “metabolic syndrome X” AND “metabolic syndrome” AND “fatty acids, omega-3” NOT “alpha-linolenic acid”. The search was then limited to “RCT” and “humans” only.

Eligibility criteria

To qualify, RCT could only include subjects diagnosed with MS who had three, four or five characteristics of the MS, without history of CVD. RCT reporting chronic effects of EPA + DHA (as either supplements or dietary components) and postprandial studies were considered. The studies had to quantify the EPA + DHA supplementation. Studies reporting effects of alpha-linolenic acid in MS patients were excluded. Also, \textit{in vitro}, animal studies and non-RCT were excluded. 138 articles were firstly identified in the search, 92 original articles, 26 reviews, 12 proceedings papers, 7 meeting abstracts and 1 editorial material. When we limited for RCT and humans (as mentioned above), the number of publications was reduced to 14. The titles and abstracts were analysed by 2 independent scientists of the field. Three studies were excluded as they did not meet the search criteria: an animal study\(^12\), a RCT using alpha-linolenic acid as omega 3 source instead of EPA + DHA\(^13\), and a study describing and evaluating a food-exchange model in MS patients for further research trials\(^14\). The 11 remaining trials were relevant to the research question and were included\(^15–25\). The references sections of those 11 studies revealed 6 additional RCT with MS patients that met the search criteria, and were also included\(^20–31\).

Quality and applicability assessment

All RCT were assessed for both study quality and applicability (Tables 1 and 2). Methodological quality refers to the design, conduct, and reporting of the clinical study. Because studies with a variety of design types were evaluated, a 3-level classification of study quality was used\(^32\), as follows: A) least bias, when the study mostly adhered to the commonly held concepts of good quality, including formal randomized study, clear description of the population, setting, interventions and comparison groups, appropriate measurement of outcomes, appropriate statistical and analytic methods and reporting, no reporting errors, <20% dropout, clear reporting of drop-outs, and no obvious bias. B) Susceptible to some bias. The study had some deficiencies but none likely to cause major bias or may be missing information making assessment of the limitations and potential problems difficult. C) Significant bias. Study has serious errors in design, analysis, or reporting or may have large amount of missing information or discrepancies in reporting. Regarding applicability, a 3-level classification was also used. I: if the sample was representative of the target population sufficiently large to cover both sexes, a wide age range, and where dietary intake was controlled. II: if the sample was representative of a relevant subgroup of the target population and food intake was not controlled but the subjects received dietary advice; III: if sample was representative of a narrow subgroup of subjects only, not well applicable to other subgroups and/or where dietary intake was not controlled at all.

Results

The studies that fulfilled the inclusion criteria and were analysed for relevance to the research question were 17 in total\(^15–31\). Information on each study is summarised in Tables 1 and 2. Twelve publications showed results from European countries (Spain, Norway, Ireland, France, Poland, The Nederlands, Sweden, Denmark, United Kingdom), two from Canada, three from Japan, one from Brazil and one from Iran. Seven European publications were results of the LIPGENE project, a large scale European Union Project conducted by 25 research centres across Europe focused on diet, genomics and the MS (http://www.ucd.ie/lipgene/). The number of participants varied from 8\(^19,21\) to 957\(^17\) in the selected studies. Eleven studies reported chronic effects of VLC n3 PUFA by administering n3 rich diets or supplements for a period of time and analysed parameters of MS patients in fasting conditions, compared with controls (Table 1). Six RCT published recently have evaluated the postprandial response of MS patients to fat.
### Table 1. RCT describing chronic effects produced by the supplementation with EPA + DHA in MS patients (1)

<table>
<thead>
<tr>
<th>Author, year (reference number)</th>
<th>n</th>
<th>Dose (g/day)</th>
<th>Duration</th>
<th>Quality(2)</th>
<th>Changes observed in the VLC n3 PUFA Group</th>
<th>Conclusion of the study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brady et al., 2004(15)</td>
<td>30</td>
<td>2.5</td>
<td>6 weeks</td>
<td>B</td>
<td>TAG -25%, small dense LDL -11% Greater TAG reductions in this group</td>
<td>Cardioprotective TAG lowering effects of the EPA + DHA supplement not interfered by the high n-6 PUFA diet</td>
</tr>
<tr>
<td>Benito et al., 2006(16)</td>
<td>72</td>
<td>0.186</td>
<td>3 months</td>
<td>A</td>
<td>Systolic BP -9.7 mm Hg, diastolic BP -7.3 mm Hg, TC -6.2%, LDL-C -7.5%, TAG -13.3%, fasting glucose -5.3%</td>
<td>Low dose of EPA + DHA administered with oleic acid in a dairy drink improved the CV risk profile in MS patients</td>
</tr>
<tr>
<td>Satoh et al., 2007(17)</td>
<td>44</td>
<td>1.8</td>
<td>3 months</td>
<td>B</td>
<td>CRP -63%, TAG of remnant lipoprotein particles -40%, cholesteryl ester transfer protein activity -2.5%, sdLDL particles -1.3%</td>
<td>The biomarkers modified by the EPA supplementation potentially reduce development of atherosclerosis and CVD in MS</td>
</tr>
<tr>
<td>Saito et al., 2008(18)</td>
<td>957</td>
<td>1.8</td>
<td>5 years</td>
<td>A</td>
<td>TAG -25%. Risk for major coronary events -53% vs control</td>
<td>EPA may be especially beneficial in patients who with abnormal TAG and HDL-C levels</td>
</tr>
<tr>
<td>Ebrahimi et al., 2009(19)</td>
<td>89</td>
<td>0.3</td>
<td>6 months</td>
<td>B</td>
<td>Weight -3.8%, systolic BP -5%, TC -13%, LDL-C -21%, TAG -18%, CRP -66% and Hsp27 -68%</td>
<td>The low dose of VLC-omega 3 PUFA could be useful to improve the CV risk profile of the MS patients, but to be confirmed</td>
</tr>
<tr>
<td>Satoh et al., 2009(20)</td>
<td>92</td>
<td>1.8</td>
<td>3 months</td>
<td>A</td>
<td>TAG -25%, SAA-LDL -17%, CRP -26%, PWV -5.6%, CAVI -3.5%, adiponectin +8%.</td>
<td>EPA improves arterial stiffness through suppression of SAA-LDL, which could be an important step for the EPA anti-atherogenic effects</td>
</tr>
<tr>
<td>Hartwich et al., 2009(21)</td>
<td>78</td>
<td>1.24</td>
<td>12 weeks</td>
<td>A</td>
<td>Favourably altered LDL phenotype from B (sdLDL) to A, decreased density of LDL, no changed in oxidative stress markers</td>
<td>The study showed efficacy of dietary n-3 PUFA to modify pro-atherogenic to less atherogenic LDL phenotype in MS patients.</td>
</tr>
<tr>
<td>Gulseth et al., 2010(22)</td>
<td>391</td>
<td>1.24</td>
<td>12 weeks</td>
<td>A</td>
<td>No changes in systolic BP, diastolic BP or pulse pressure</td>
<td>Lack of effect in BP. n3 supplement dose may be too low</td>
</tr>
<tr>
<td>Tierney et al., 2010(23)</td>
<td>317</td>
<td>1.24</td>
<td>12 weeks</td>
<td>A</td>
<td>no effect in insulin sensitivity, TC, LDL-C, BP or inflammation. TAG -7%, NEFA -10%, TC:HDL-C -4%, atherogenic index -19%, ApoCII -3.6%</td>
<td>The VLC n-3 PUFA diet improved TAG-related MS risk profiles</td>
</tr>
<tr>
<td>Petersson et al., 2010(24)</td>
<td>317</td>
<td>1.24</td>
<td>12 weeks</td>
<td>A</td>
<td>No effects of oxidative stress and inflammation markers (8-iso-PGF2α, 15-keto-13,14-dihydro-PGF2α, CRP)</td>
<td>Lack of effect explained by the low n3 dose administered or high n3 endogenous levels at baseline</td>
</tr>
<tr>
<td>Simão et al., 2010(25)</td>
<td>40</td>
<td>3</td>
<td>90 days</td>
<td>B</td>
<td>TAG -23%, TC +7.9%, LDL-C +18.8% Plasma Antioxidant capacity +16%, blood glucose +4%, insulin resistance index +24%</td>
<td>The intake of fish oil favourably reduced TAG and increased antioxidant capacity but increased TC, LDL-C and insulin resistance</td>
</tr>
</tbody>
</table>

1. Abbreviations: Atherogenic index, log (TAG/HDL-C); BP, blood pressure; CAVI, cardio-ankle vascular index; CETP, cholesteryl ester transfer protein; CRP, C-reactive protein; CV, cardiovascular; HDL-C, high-density lipoprotein cholesterol; Hsp27, heat shock protein 27; VLC n3 PUFA, long chain n-3 polyunsaturated fatty acids; LDL-C, low-density lipoprotein cholesterol; MS, metabolic syndrome; n, number of subjects in the study; NEFA, non-esterified fatty acid; PWV, pulse wave velocity; TC, total cholesterol; RCT, Randomized Controlled Trial; SAA-LDL, Serum Amyloid A–LDL; sdLDL, small dense LDL.

2. The criteria used to assess quality and applicability of the studies in the Materials and Methods section.
loads enriched with VLC omega 3 PUFA, the most recent ones are results from the LIPGENE project (Table 2). The results of the trials analysing chronic effects and biomarkers in fasting conditions are presented first.

**Studies reporting chronic effects of VLC-n3 PUFA**

The first RCT in MS patients reporting effects of VLC n3 PUFA was published in 2004 by Brady et al. This study examined whether a higher intake of n-6 PUFA in combination with a low intake of VLC n3 PUFA played a role in markers of MS in Indian Asians. The results demonstrated that supplementation with VLC n3 PUFA had no impact on insulin action but showed significant reductions in fasting TAG and a reduction in the percentage of denser, proatherogenic LDL particles. Another RCT with MS patients investigated the effects of an EPA + DHA and oleic acid enriched milk compared with a control milk. At the end of the 3-month study, MS patients of the enriched milk group improved fasting glucose levels and significantly reduced BP, TAG, total and LDL-cholesterol. In a study carried out in Japan, the supplementation with 1.8 g of EPA for 3 months showed reductions in the TAG of the remnant lipoprotein particles, cholesteryl ester transfer protein (CETP) activity (that may reduce the atherogenic process), the proportion of small dense LDL particles (sdLDL) and C-reactive protein (CRP). The JELIS study was a large-scale clinical trial examining the effects of EPA on coronary artery disease (CAD) (n = 14,981). The patients were assigned to an EPA + statin group, administering 1.8 g of EPA/d, or a control group that received only the statin, for a period of 5 years. EPA did not affect TC, LDL-C or HDL-C, but significantly reduced TAG. The EPA treatment lowered the risk for major coronary events in the context of a HSFA intake, a lower capacity for fibrinolysis was produced by the high n-3 fat load consumed.

<table>
<thead>
<tr>
<th>First author/year (reference)</th>
<th>VLC n3 PUFA dose in post prandial test (g)</th>
<th>Quality</th>
<th>Applicability</th>
<th>Changes observed in the n3 Group</th>
<th>Conclusion of the study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tulk and Robinson, 2009(29)</td>
<td>3-3</td>
<td>A</td>
<td>II</td>
<td>No changes in postprandial TAG or inflammatory response (IL-6, soluble IL-6 receptor and CRP)</td>
<td>Increasing the n-3 PUFA content of a HSFA diet had no effects on postprandial TAG or inflammatory responses</td>
</tr>
<tr>
<td>Montegaard et al., 2009(21)</td>
<td>3-3</td>
<td>A</td>
<td>II</td>
<td>Reduction of PAI-1 concentration and activity reductions of t-PA activity. No effects on t-PA concentration and platelet aggregation.</td>
<td>In the context of a HSFA intake, a lower capacity for fibrinolysis was produced by the high n-3 fat load in postprandial TAG</td>
</tr>
</tbody>
</table>
following four isocaloric diets (LIPGENE diets): (A) High SFA diet; (B) High MUFA diet; (C) low-fat diet with a daily supplement of 1.24 g/d high oleic sunflower oil; and (D) low-fat with a supplement of 1.24 g/d VLC n3 PUFA (n3 diet). The diet and supplements were administered for 12 weeks and the biomarkers were analysed at the beginning and at the end of the study period.

In the first study(22), the n3 diet resulted in favorable alteration of LDL phenotype from small, dense proatherogenic to large LDL particles and also decreased LDL density. In the second study(24), no significant differences on systolic or diastolic BP in the entire cohort. Regarding oxidative stress and inflammation, the results obtained by Peterson et al.(28) showed no effects of the LIPGENE diets on markers of oxidative stress or inflammation. This lack of effects was explained by the low dose of omega 3 used in the study (1.2 g/day) could have explained the lack of effect as only supplements of high doses (>3 g/d) have been reported to lower BP in hypertensive patients(39). In the third study(27), the LIPGENE diets showed no effect in insulin sensitivity, TC, LDL-C, inflammation or BP in the entire cohort. Regarding oxidative stress and inflammation, the results obtained by Peterson et al.(28) showed no effects of the LIPGENE diets on markers of oxidative stress or inflammation. This lack of effects was explained by the low dose of omega 3 compared with previous studies(15,16) or the endogenous levels of n3 fatty acids at baseline being too high to be modified by the diets administered.

Finally, one study carried out in Brazil(25) investigated the effects produced by the administration of 1.8 g of EPA + 1.2 g DHA per day (10 g of fish oil) to a group of MS patients. The fish oil group decreased TAG but increased total cholesterol, LDL-C and the antioxidant capacity of plasma. In addition, the fish oil group worsened the glucose control in line with other previous studies(40,41).

Postprandial studies with VLC-omega 3 PUFA

Elevated postprandial lipemia has recently emerged as a risk factor for CVD, indicating that this disease is in part a postprandial phenomenon(42,43). Postprandial hyperlipidemia is characteristic of MS patients, with increased postprandial TAG-rich lipoprotein (TRL) concentrations(44,45).

Tulk and Robinson(19) showed that increasing the VLC n3 PUFA content of a high-SFA oral fat load did not change postprandial TAG or inflammatory responses. Using a similar design, the same research group showed that a high n3 fat load increased PAI-1 reduced t-PA activity compared with the low n-3 fat load. The authors concluded that in the context of a high SFA intake, a lower capacity for fibrinolysis was produced by the high VLC n3 PUFA fat load.

From the LIPGENE project, four RCT have evaluated postprandial effects of VLC n3 PUFA, all of them using a similar design, as follows. MS patients were divided into 4 groups that received the LIPGENE diets A-D (see above) for 12 weeks. A fat challenge with the same fat composition as the LIPGENE diets, but administering 0.7 g of fat load/kg of body weight, was conducted pre- and postintervention. Jimenez-Gomez et al.(25) examined the postprandial effects of the fat loads in the lipoprotein profile of 117 MS patients. The n3 diet group had a lower postprandial TAG concentration than the other diet groups. In contrast, long-term ingestion of the n3 diet did not have any effect on postprandial TAG and TRL. The same research group reported no effects of the n3 diet on the endothelium-dependent vasomotor function and plasma levels of cellular adhesion molecules in 75 MS patients(29). Another study reported that the n3 diet group could favourably modify the LDL particle profile and plasma Ischemia Modified Albumin (IMA)(31). Finally, the administration of the LIPGENE fat loads did not affect forearm muscle FA handling in men with MS, but the VLC n3 PUFA group decreased their postprandial TAG concentrations(30).

Discussion

Biomarkers of intake

The use of biomarkers of intake is essential to show compliance with the intake of the supplement or test food of the study and to correlate effects produced by the dietary fat. However, there is little consensus about which biomarker of dietary fat modification should be used in RCT. Of the studies showing chronic effects of VLC n3 PUFA, only 7 studies reported biomarkers of intake in blood, including plasma concentration of EPA/DHA fatty acids(20–28,17), increases in the n3/n6 ratio of plasma FA(22), percentage of n3 FA in serum phospholipids(16) and the ratio of n6:n3 platelet membrane phospholipids(15). All the VLC n3 PUFA supplements used in the trials were either fish oils high in EPA (as triglycerides) or purified EPA (as ethyl esters) with the exception of one study(16) using DHA-rich oil (from unknown origin). The intervention periods ranged from 6 weeks(16) to 5 years(37) and the amounts of VLC n-3 PUFA administered ranged from 186 mg/day(16) to 3 g/d(25). The increases in plasma EPA or n3/n6 ratio ranged 60%-120% and were reported after at least 6 weeks of intake. The available RCT showed that in order to observe a significant increase in plasma EPA in MS patients, a minimum amount of 1.2 g or 4 g of fish oil has to be administered for at least 3 months or 6 weeks, respectively. The study using the low dose (186 mg/day) of DHA oil for 2 months did not show any significant increase in plasma EPA or DHA. The use of one standardised biomarker of dietary fat modification would be desirable for this type of interventions. In this sense, erythrocyte EPA + DHA could be an option as it has been suggested to be a useful biomarker for CHD that may also predict the risk of cardiac events(40).

Effects of VLC n3 PUFA on MS

a) VLC n3 PUFA and blood lipids. Elevated fasting TAG increases the risk of CVD(31). The TAG lowering effects of VLC n3 PUFA have been extensively described(37). High doses (2–4 g/d) of EPA + DHA decrease serum triglycerides in both normo- and hyperlipidaemic individuals but low doses have also reported reductions when substitute SFA(48,49). The magnitude of the effects is related both to the dose of EPA + DHA and to the baseline concentrations of triglycerides(50). Harris(51) observed a mean reduction of 35% in subjects with hypertriglycercidaemia and of 24% in those with serum triglycerides < 2 mmol/L. In the meta-analysis by Balk et al.(32),

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a mean reduction of 27% in serum triglyceride concentrations was observed. In the diet and lifestyle recommendations given by the AHA, 2–4 g EPA + DHA per day provided in capsules under physician’s supervision are recommended for individuals with hyper-triglyceridaemia to lower TAG 20–40% \(^{(1)}\).

All RCT reporting blood lipid effects upon supplementation with VLC n-3 PUFA showed fasting TAG reductions, ranging from 7% to 25% (Fig. 1). One study also reported 40% reductions of TAG in remnant lipoprotein particles \(^{(26)}\). The mechanisms that explain the TAG lowering effect include inhibition of triglyceride synthesis, stimulation of fatty acid beta-oxidation, and increased lipoprotein lipase-mediated clearance of VLDL triglycerides \(^{(50)}\). The reported reductions in apo CII levels \(^{(27)}\) would facilitate TRL catabolism and apo-E dependent hepatic uptake of TRL remnants. Increased postprandial lipemia has recently emerged as a CVD risk factor, indicating that this disease is, at least in part, a postprandial phenomenon. Patients with MS usually have increased postprandial lipemia. However, the 5 postprandial studies administering VLC n-3 PUFA showed little improvements in the postprandial TAG response. Only one study concluded that the administration of VLC n-3 PUFA could reduce the adverse postprandial TAG-raising effects of long-term LFHCC diets \(^{(23)}\).

None of the considered studies reported modifications of HDL-C by administration of VLC n-3 PUFA. One study showed an activity reduction of CETP, but without effect of HDL-C \(^{(20)}\). Regarding TC and LDL-C, the RCT supplementing VLC n-3 PUFA to MS patients showed conflicting results: an increase in TC and LDL cholesterol was reported when high doses (3 g/d) were used for 3 months \(^{(25)}\), whilst administration of 1 g of fish oil containing 300 mg EPA + DHA for 6 months resulted in TC and LDL-C reductions \(^{(18)}\). In this latter study, the volunteers were not instructed to maintain their diet and lifestyle, the dietary intake was not controlled and a 3.8% reduction of body weight was obtained at the end of the study, which could have played a part in the reductions observed. Benito et al. \(^{(16)}\) also described reductions in TC and LDL-C upon 3-month consumption of a dairy drink where SFA were substituted by EPA + DHA (186 mg/day) and oleic acid. EPA plus DHA at high doses (2–4 g/d) have multiple effects on blood lipids \(^{(50)}\). TC and LDL-C concentrations, are generally not affected by this VLC n-3 PUFA supplementation, but in subjects with hypertriglyceridaemia, LDL-cholesterol concentrations may be increased by 5–10% \(^{(22,51,52)}\). Whilst the TAG reduction produced by VLC n-3 PUFA produces clear CV benefits, the concomitant LDL-C increases reported at moderate to high doses represent a risk to these patients. The daily dose of EPA + DHA and length of time needed to produce a relevant TAG reduction without increasing LDL-C or TC is yet to be determined for MS patients. More studies are required to clarify this point.

**b) VLC n-3 PUFA and LDL phenotype.** LDL particle size is an important predictor of CV events and progression of CHD \(^{(53,54)}\). The sdLDL pro-atherogenic particles have been associated with obesity, insulin resistance, high blood TAG and oxidative stress \(^{(55)}\). Supplementation of MS patients with VLC n-3 PUFA consistently reduce the percentage of sdLDL particles in favour of the large, buoyant, less atherogenic LDL phenotype \(^{(15,22,29)}\) which potentially reduces the CV risk. sdLDL concentrations are very much influenced by plasma TAG \(^{(50)}\). In MS, slow clearance of TAG from TRL lipoproteins allows for longer periods of circulating TAG, which can directly affect the composition of LDL. Increased transfer of TAG to LDL via CETP during neutral-lipid exchange and subsequent hydrolysis of LDL-TAG by lipases results in a preponderance of sdLDL \(^{(50)}\). Suppression of TAG production in the liver by EPA and reduction of CETP activity may explain the beneficial phenotype shift.

**c) VLC n-3 PUFA and BP.** Data from the available studies indicate that supplementation of MS patients with VLC n-3 PUFA does not produce significant effects on BP with the doses used. The large and well designed trial by Gulseth et al. \(^{(24)}\) from the LIPGENE project addressed the effects of VLC n-3 PUFA on the BP of MS patients. This trial showed no effects on BP of a dose of 1-24 g/day for three months. In line with this, a trial by Murphy et al. \(^{(57)}\), 86 overweight subjects with high serum TAG were randomised to 1 g/day of EPA + DHA (from enriched foods) or placebo, for 6 months, obtained no significant effects on BP. However, other two RCT trials described above \(^{(16,18)}\) reported significant decreases

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**Fig. 1.** TAG reductions \(^{(1,2)}\) reported in RCT with MS patients upon VLC n-3 PUFA supplementation. (1) Expressed as percentages calculated from baseline values. (2) All reductions were statistically significant \((P<0.05)\).
of BP using much lower amounts (300 mg and 186 mg/day). In those trials however, the patients from the intervention group lost a significant amount of weight during the study period which may indicate that the BP effect may not be derived from the n3 supplementation.

Indeed, EFSA, after a thorough revision of the scientific evidence, indicated that much higher doses of EPA + DHA (≥ 3 grams/day) are needed to have a short-term effect on systolic BP in subjects with untreated hypertension (52,58) (~3–5 mmHg decrease in systolic BP), which may also have smaller effects in normotensives (~1 mmHg decrease in systolic BP). Studies with MS patients using those high doses of EPA + DHA are currently lacking. Interestingly, the recent study by Satoh et al. (20) in MS patients showed that n3 supplementation of 1.8 g/day for 3 months improved wave pulse velocity and cardio-ankle vascular index, two biomarkers of arterial stiffness, the latter being less influenced by BP changes. Perhaps the use of these new biomarkers would be useful to better characterise the magnitude of the effects. It should be stated here that the above RCT did not differentiate between MS patients with normal or high BP at baseline, and antihypertensive medication was sometimes allowed (24). These factors make more difficult the interpretation of the data. Whether MS patients, all suffering from high BP as risk factor, may benefit from VLC n3 PUFA supplementation remains unanswered.

d) VLC n-3 PUFA and glucose control. Epidemiologic studies have reported a lower prevalence of impaired glucose tolerance and type-2 diabetes in populations consuming large amounts of fish (59). VLC n5 PUFA may decrease insulin resistance through a number of mechanisms including decrease in circulating TG and small dense LDL particles, increasing membrane fluidity and signal transduction. Further, substituting saturated fat with unsaturated fat, such as n-3 PUFAs, may have beneficial effects on insulin sensitivity (60). This change in dietary fat may reduce the risk of progression to type-2 diabetes from impaired glucose tolerance by improving hemostasis, albuminuria, subclinical inflammation, oxidative stress and slowing progression of artery narrowing (61). Clinical studies have shown that consumption of n-3 PUFAs has cardioprotective effects in persons with type-2 diabetes without adverse effects on glucose control and insulin activity. 3 different meta-analysis concluded that the use of moderate to high amounts of fish oil in diabetics had no effect on glycemic control (52,62,63). More recently published reviews also failed to show effects of VLC n-3 PUFA on the glucose control of type-2 diabetes patients (64–66).

In MS patients, effects of fasting glucose, insulin and insulin resistance (shown as HOMA) were reported in 5 RCT using different doses of n-3 PUFA. 3 g/day of EPA + DHA for 3 months increased blood glucose and insulin resistance by 4% and 24% respectively (25). Three studies administering daily doses of 2.5 g for 6 weeks, 1.8 g and 1.2 g both for 3 months did not show any effects (15,26,27). No changes in glycosylated haemoglobin were reported (28). The 3-month study with the low dose of EPA + DHA (186 mg/day) administered in a dairy drink showed that fasting glucose was reduced by 5.3% in the n3 group but the effect of the oleic acid and the weight reduction observed cannot be ruled out. Therefore, there is no evidence of beneficial effects of VLC n3 PUFA on the blood glucose control of MS patients and may even be harmful at high doses. Future studies with different EPA + DHA doses are needed to determine if increased consumption of n-3 PUFA will delay the conversion of insulin-resistant subjects (MS) to diabetics.

e) VLC n-3 PUFA and inflammation. Several features of MS are associated with increased inflammatory and oxidative stress markers (67–69). For example, MS patients usually have high CRP as a manifestation of proinflammatory status. In theory, supplementation with EPA + DHA should originate production of less-potent eicosanoids with anti-inflammatory and antithrombotic effects (compared with the n6 arachidonic acid) but the reports available are controversial. The LIPGENE intervention studies showed no significant effect of dietary fat modification (quantity and quality) and VLC n3 PUFA on several markers of inflammation (CRP, 15-keto-13,14-dihydro-PGF2α, IL-6, TNFα, sICAM, sVCAM, resistin, adiponectin, leptin) and oxidative stress (8-iso-PGF2α) (27,28). The postprandial studies only showed modest effects: higher endothelial cell function, decrease of sICAM-1 (29) and reduced postprandial elevation of the IMA which is related to oxidative stress (31). In contrast, other three RCT administering daily doses of 1.8 g, 0.3 g and 0.186 g of EPA + DHA showed significant reductions of CRP and of the inflammatory status (26,16,18). Satoh et al. (20) also reported an increase in adiponectin, an anti-atherogenic adipocytokine, and reduced serum amyloid A-LDL complex, an oxidised form of LDL generated under inflammatory conditions. In the LIPGENE trials, the n3 PUFA supplementation was carried out in the context of a low fat diet, where other factors may play a role. The study by Benito et al. (18) were poorly controlled and produced BP and body weight reductions. The only controlled study with weight stable MS patients in which both groups received the same diet showed CRP reductions in the n3 supplemented group (20) but was performed with a low number of participants (n = 44, 22 per group). More controlled studies with different doses of VLC n3 PUFA and higher number of participants are necessary to test the effects of VLC n3 PUFA on the inflammation associated with MS.

d) VLC n-3 PUFA and CHD risk. One study addressed this point (17) by analysing the high TAG/low HDL-C subgroup of patients (administered with 1.8g EPA/day + a statin) of the JELIS primary prevention trial. The results showed that in the high TAG/low HDL-C group, the risk of major coronary events was particularly high, and EPA was shown to potentially reduce the risk by 53%. In the high TAG/low HDL-C group, level of plasma EPA at the time of registration was low, but EPA administration increased plasma levels, suggesting some correlation between plasma EPA level and major coronary events risk. Fish intake has been associated with decreased risk of CVD (70). A few epidemiological studies have reported that men who ate at least some fish weekly had a lower coronary heart disease (CHD) mortality rate than that of men who ate none (59,71,72). A 30-year follow-up of the Chicago Western Electric Study showed that men who consumed 35 g or more of fish daily compared with those who consumed
none had a relative risk of death from CHD of 0.62 and a relative risk of nonsudden death from MI of 0.33. For these reasons and on the basis of the available data, the regular intake of fish, at least twice a week (preferably oily fish), is recommended to patients without documented CHD like the MS patients.

Concluding remarks

MS is characterised by a cluster of CVD risk factors. The available RCT on MS patients convincingly show that the administration of VLC n3 PUFA doses > 1 g for at least 3 months produced a significant reduction of triglycerides ranging from 7% to 25%. These results confirm the hypotriglyceridemic effect of VLC n3 PUFA in MS patients. The triglyceride lowering may produce further benefits on % sdLDL and perhaps ameliorate the inflammatory process associated with MS. No clear effects were found on other MS markers. High doses of VLC n3 PUFA (≥3 g/day) may produce further TAG reductions but could raise other risk factors such as LDL-C. The combination of VLC n3 PUFA plus a statin may be useful to prevent the occurrence of coronary events. More studies are needed using different amounts of VLC n3 PUFA, time lengths, dietary backgrounds and different profiles of MS patients before clear recommendations can be made.

Although both EPA and DHA are part of the fat composition of seafood products, their percentages in the oils differ depending on the species and other factors. Whilst fish oils in general usually contain about 30% of EPA + DHA, tuna oil can be as high as 25% in DHA and herring oil usually contains 18% of EPA. In recent years, differential effects on gene expression, cell function and physiology have been reported for EPA and DHA in their purified forms, reviewed in (74). Indeed, purified EPA and DHA administered in capsules seem to have differential effects on serum lipids and lipoproteins, LDL particle size, glucose and insulin release (40, 75). In addition, DHA and EPA possess different affinity for organs. DHA generally exceeds EPA 5- to 30-fold in most organs (76). Future research should explore these differential effects in MS patients. In addition, nutrigenetics and nutrigenomics are exceptional tools to identify possible EPA + DHA gene targets related to MS risk factors like obesity and insulin resistance.

Finally, in the last decade we have witnessed an extraordinary increase in the number of functional foods targeted to the ever-growing health conscious population. Several categories among these foods can be found according to their health related to labelling reference intake values for n-3 and n-6 polyunsaturated fatty acids. EFSA J 1176, 1–11.

Conflict of interest

The author declares no conflict of interest.

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