Omega-3 Fatty Acids for Nutrition and Medicine: Considering Microalgal Oil as a Vegetarian Source of EPA and DHA

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Abstract: Long-chain EPA/DHA omega-3 fatty acid supplementation can be co-preventative and co-therapeutic. Current research suggests increased accumulated long chain omega-3s for health benefits and as natural medicine in several major diseases. But many believe plant omega-3 sources are nutritionally and therapeutically equivalent to the EPA/DHA omega-3 in fish oil. Although healthy, precursor ALA bioconversion to EPA is inefficient and production of DHA is nearly absent, limiting the protective value of ALA supplementation from flax-oil, for example. Along with pollutants certain fish acquire high levels of EPA/DHA as predatory species. However, the origin of EPA/DHA in aquatic ecosystems is algae. Certain microalgae produce high levels of EPA or DHA. Now, organically produced DHA-rich microalgae oil is available. Clinical trials with DHA-rich oil indicate comparable efficacies to fish oil for protection from cardiovascular risk factors by lowering plasma triglycerides and oxidative stress. This review discusses 1) omega-3 fatty acids in nutrition and medicine; 2) omega-3s in physiology and gene regulation; 3) possible protective mechanisms of EPA/DHA in major diseases such as coronary heart disease, atherosclerosis, cancer and type 2 diabetes; 4) EPA and DHA requirements considering fish oil safety; and 5) microalgae EPA and DHA-rich oils and recent clinical results.

Keywords: Omega-3s, Schizochytrium, Fish oil, EPA, DHA.

INTRODUCTION

The foundation of Ayurveda (natural Indian medicine) is diet [1, 2]. Similarly, Hippocrates stated ‘Let food be thy medicine and medicine be thy food’ [3]. To achieve optimum health, contemporary lifestyles must consider food choices, eating habits and strategies for building up omega-3 levels in the body [4, 5]. For individuals ‘at risk’ for diet induced diseases everywhere, even dietary limitations, vitamin supplementation, prescription remedies, alternative medicine, and physical exercise may not be fully protective, preventative, or therapeutic without addressing inherent omega-3 fatty acid deficiencies [6, 7].

The Indian paradox of diet-induced diseases, whether vegetarian or non-vegetarian, is not isolated from the global dilemma facing modern society. The paradox is often simultaneous overeating and undernutrition within the same individual, leading to the initiation of major diseases [5]. Although various origins are proposed for some cases of obesity in India [8], urbanization and modernization increase the numbers consuming foods associated with convenience diets [9]. These include high fat foods, processed foods, snacks, and drinks made available for instantaneous consumption, resulting in a habit of overeating with overly-frequent caloric intake. Too much fatty, oil fried, and processed food often leads to insulin resistance, obesity, type-2 diabetes, high blood pressure, atherosclerosis, and heart disease [10]. To the extent that a modern diet departs from traditional dishes with their rich compliments of fresh herbs and spices - neglecting uncooked fruits, vegetables, and leafy greens as functional foods - there may be increased risk of micronutrient undernutrition [9, 11], particularly in essential omega-3 fatty acids [12]. Thus, many processed foods are nonfunctional and dysfunctional foods and result in diets containing little omega-3 nutritional complement.

OMEGA-3 FATTY ACIDS IN NUTRITION AND MEDICINE

Omega-3 fatty acids focused on throughout this review are the bioactive linoleic acid (LA) is the main omega-6 ‘precursor’ in plant/vegetable oils. The omega-6 fatty acid arachidonic acid (AA) is bioactive and found in red meat.

The balance of omega fatty acids is important to consider. The so-called omega-3:omega-6 ratio has become a model for gauging the proper balance of these fats in oils and the diet [13]. Diets with greater than a 1:10 ratio of omega-3 to omega-6 are not recommended, whereas a 1:1 ratio is considered perfect. Very unhealthy ratios of 1:25 and 1:50 are common, especially with regular consumption of ‘fast-food’, high amounts of fried food, and low intake of fresh whole foods. Thus, ‘Eating to live and not living to eat’ becomes an important consideration with increases in modern, convenient, non-functional food choices.

Fig. (1) shows the fatty acids of major interest in the diet where n-3 indicates omega-3 and n-6 indicates omega-6 (Fig. 1A-B). Notably, EPA n-3 and AA n-6 are the same length with 20 carbons each. However, EPA is unsaturated at the n-3 position providing signaling and metabolic specificity between AA and EPA. AA is converted into inflammatory compounds, but the unique n-3 bond means EPA is converted into dramatically different bioactive products. In addition, EPA may compete directly with AA for many of the same enzymes during the inflammatory process [14]. The other important omega-3, DHA n-3, has 22 carbons, is unsaturated at n-3 and is significant for affecting triglyceride levels, by as yet unknown mechanisms [15]. DHA is also incorporated into membrane phospholipids in the brain [16].

In fact, EPA/DHA omega-3 status may be one of the single most important nutritional requirements left unaddressed in nutritional medicine, but emphasis on omega-3 supplementation for health is rapidly increasing [17]. At the same time awareness is improving regarding the pro-inflammatory properties of omega-6 fatty acids, [18, 19]. Insufficient versus sufficient omega-3 EPA/DHA status is now clinically linked to several diseases as a co-cause or co-treatment, respectively [20]. In addition, clinical studies are now concluding significant benefits associated with EPA/DHA from fish oil and DHA-rich oil from microalgae [15, 20-25,]. High doses of oil from both fish and microalgal sources are considered safe, result in increased circulation of both EPA and DHA omega-3s, and both oil sources are protective against cardiovascular risk factors [20, 24, 25]. Significantly, DHA-rich oil formulations are equally protective compared to fish oil in nearly all human trials conducted.
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cal studies indicate beneficial effects of DHA-rich oil for cardiovascular risk prevention in healthy men and women [15], producing significant decreases in plasma triglyceride levels [21]. Results were comparable to fish oil in effects, bioavailability and safety profiles [22, 23]. In direct studies, algae oil cardio-protective effects were similar to fish oil [24, 25]. Furthermore, DHA-rich oil supplementation increased DHA levels in lactating women, in breast milk and in nursing infants [26]. DHA is particularly important for fetus development, pregnancy outcomes, cognitive development and maintenance, learning and memory, visual function, the immune system, and more [27]. Finally, no reports are currently available for microalgal DHA-rich oil treatment in diabetic patients.

SYNTHESIS AND METABOLISM

The vegetarian diet largely depends upon ALA to be synthesized into EPA and DHA in the body, according to the pathway shown in Fig. (1C). Important findings now show that ALA conversion into EPA and DHA is rate limiting [28, 29]. The first metabolic step in EPA synthesis by the enzyme Δ6 desaturase is kinetically slow (shown in Fig. (1C) as dashed arrow), so most ALA undergoes oxidative metabolism for energy and is not converted to EPA. That is, omega-3 oil supplementation from plant sources may produce only 7% of the EPA levels compared to the EPA levels gained through fish oil supplementation [29]. DHA accumulation from ALA only occurs at trace levels due to the additional steps required to convert EPA into DHA. Similarly, EPA concentrations increase with dietary EPA, but DHA does not [29]. Also, conversion of omega-6 to omega-3 does not occur. In contrast, dietary DHA supplementation will result in steady state DHA concentrations and modest increases in EPA concentrations through DHA retro-conversion, producing EPA at 5-11% of accumulated DHA levels [28, 29]. With continuous supplementation, EPA/DHA levels plateau, suggesting the body first needs to incorporate omega-3s directly through the diet before limiting omega-3 fatty acid accumulations by oxidative metabolism.

With few ALA rich plant sources for vegetarians, omega-3 supplementation is an important consideration. Beyond flax, walnut and mustard seed oils, ALA supplementation is limited because stearidonic acid (STA) is poorly formed as the second product in the EPA/DHA synthetic pathway [30]. Bypassing Δ6 desaturase at the first step with STA supplementation allows for EPA accumulation up to 30% of levels achieved by fish oil regimens [30]. However, plant sources of STA are found most abundantly in the invasive plant species *Echium* and seldom used herb *Borage*.

As a staple in the Indian diet, mustard seeds contain just 1-4% ALA omega-3 (~200 mg/tablespoon). Yet mustard oil supplementation does not produce significant effects compared to EPA/DHA omega-3s [31, 32]. Also, as a food it is not clear what effects pre-roasting or heating of mustard seeds have on ALA stability. As with all unsaturated fats, heating and cooking causes a percentage of the fat to undergo air oxidation. Polyunsaturated fats such as omega-3s and omega-6s are particularly sensitive to oxidation and must be consumed in relatively fresh foods. This suggests conversion of ALA to longer chain omega-3s may be further limited by oxidation of available omega-3 precursors in pre-cooked and/or processed foods. Proper information is particularly important as awareness of the benefits of omega-3 fatty acids increases. Also, because DHA is not readily produced from plant omega-3s, other truly alternative vegetarian sources of DHA are needed. At present, direct EPA/DHA delivery is believed dependent on certain marine fish species, which many fear may not be a pure or sustainable resource. Reputable data suggest wild marine fish populations are already becoming depleted [33].

Fig. (1). Omega-3 and Omega-6 Fatty Acid Structures and EPA/DHA Synthesis Pathway.
A) Omega-3 fatty acids include ALA (alpha-linolenic acid), STA (stearidonic acid), EPA (eicosapentaenoic acid), DPA n-3 (docosapentaenoic acid n-3), DHA (docosahexaenoic acid). B) Omega-6 fatty acids include LA (linoleic acid), GLA (gamma-linolenic acid), DHGLA (dihomo-gamma-linolenic acid), AA (arachidonic acid), DPA n-6 (docosapentaenoic acid n-6). The conventional designations are as follows # of carbons and # of double bonds with the first double bond from the end, the omega, stated as n-3 or n-6; thus for DHA n-3 the fatty acid is given as (22:6 n-3) and for EPA n-3 it is given as (20:5 n-3). Additionally, the arrows in A and B point to the omega-3 and omega-6 carbon, respectively. C) The EPA/DHA synthesis pathway is given with fatty acids and respective enzymes for each step. The omega-6 synthesis of LA (18:2 n-6) to DPA n-6 (22:5 n-6) uses the same enzymes.
However, microalgal oil products now provide an organic, vegetarian, sustainable alternative source of EPA/DHA. Yet these are still relatively unknown products.

**OMEGA-3 EPA/DHA IN PHYSIOLOGY AND GENE REGULATION**

Why are the longer chain omega-3 fatty acids different from other fatty acids in terms of their biological effects? Likely, the higher number of double bonds starting at carbon 3 from the end gives these fatty acids uniqueness and bioactive properties that were harnessed by organisms over evolutionary time. For instance, the brain is enriched in long-chain omega-3 fatty acids, particularly DHA, in phosphatidycholine and phosphatidylethanol phospholipids concentrated in plasma membranes at the neuronal synapse [16]. Plasma membranes contain various amounts of DHA or EPA in phospholipids. Biophysical studies of omega-3 membrane-protein interactions and functions are emerging areas of research.

In addition, long chain omega-3 fatty acids are definite biological regulators. Overall, reduction of chronic inflammation and improvement of lipid metabolism are positive effects that originate through omega-3 effects on gene expression [34]. The bioinformatics of omega-3 gene regulation has been provided here in summary, and because of recent reviews and research articles these data are only briefly covered here [34-38]. Two main groups of genes are affected by EPA/DHA omega-3s. These contain inflammatory genes and energy metabolism genes. First, inflammatory genes are suppressed by EPA/DHA and are as follows: nuclear factor κB; inhibitory κB kinase; inducible nitric oxide synthase; interferon γ; Interleukin-1β, 2, 6, 8, & 12; E-selectin; intercellular adhesion molecule; vascular cell adhesion molecule; monocytic chemoattractant protein 1; C-reactive protein; von Willebrand factor; matrix metalloproteinase 9; tumor necrosis factor α; and cyclooxygenase 2. Besides inflammation, endothelial and angiogenesis processes are implicated by these data. Second, energy metabolism genes are increased by EPA/DHA and are as follows: peroxisome proliferator-activated receptors; sterol regulatory element-binding proteins; adipocyte fatty acid binding protein; acyl CoA oxidase; uncoupling protein 1; carnitine palmitoyltransferase 1; leptin; pyruvate dehydrogenase kinase 4; glucose transporter 4; Caveolin-1; Caveolin-2; fatty acid transporter protein CD36; stearoyl CoA desaturase 1: ATP binding cassette transporter A1; lipoprotein lipase; liver-X-receptors; and apolipoprotein E. Because lipid metabolism is linked to insulin resistance, treatment of type 2 diabetes by omega-3s is implicated, but likely indirect. Notably, in treated hypertensive type 2 diabetic patients either EPA or DHA independently reduced oxidative stress but not markers of inflammation [39], yet EPA and DHA reduced triglyceride levels in these patients [40].

EPA and DHA may also act as free and/or acyl-CoA conjugated fatty acids, implicating Long-chain Acyl-CoA Synthetases (ACSLs) in their activation [41]. Because ACSLs are regulated by peroxisome proliferator-activated receptors [42], by analyzing specific ACSL isoforms via RT-PCR [43], ACSL expression patterns in tissues could provide some insight into omega-3 metabolism. Also, metabolized forms of EPA and DHA bind to the PPAR family of transcription factors [44], which may differentially regulate ACSL isoforms in different tissues. Therefore, bioactive PPAR ligands derived from EPA/DHA potentially signal through ACSLs to activate genes in a positive feedback loop that could make the effects of EPA/DHA increasingly effectual in tissues as regulators of lipid homeostasis. Genes influencing innate and acquired immunity in type 1 diabetes is also an area of active research [45]. How omega-3s affect these would be interesting to know.

**OMEGA-3 FATTY ACIDS FOR COMBATING MAJOR DISEASES**

**Coronary Heart Disease:** In the 1970’s researchers noted that Eskimo populations consumed extremely high levels of fat from fish and blubber, but this contradicted hypotheses of coronary heart disease at the time because the indigenous populations had no signs of cardiovascular disease. High levels of EPA/DHA are thought to be the protective complement. Recently, DART and GISSI-P clinical studies of fish oil supplementation revealed 15-45% reduction in mortality in ‘at risk’ patients for coronary heart disease [46, 47]. Reductions in sudden death were particularly significant. One small fish oil supplementation study in Norway did not show significant improvements, probably because of habitual incorporation of cold water fish as a regular part of the diet [46]. The latter result can be explained by assuming that omega-3 status was optimum to begin with, suggesting that with full EPA/DHA essentiality in the background this meant coronary heart diseases in Norway were likely due to additional factors such as obesity or genetic factors that played dominant roles in participating subjects. However, most of the risks of coronary heart disease globally are associated with diet. A high incidence of diet induced coronary heart disease occurs in many countries, including India.

**Arrhythmias:** The benefits of omega-3s were originally thought to be due to their antiarrhythmic effects, but recent evidence has indicated that the predominant effect may be antiarrhythmic. Omega-3 supplementation decreased heart rate variability in patients after myocardial infarction, which correlated with a lower risk of mortality and malignant arrhythmia [48]. In fact, direct addition of EPA/DHA into media with cultured cardiomyocytes prevents or terminates pharmacologically induced or electrically clamped arrhythmias. The modulation of plasma membrane permeability and the stabilization of ion channel functions are suggested to be acute protective properties of EPA/DHA on heart muscle cells.

**Atherosclerosis and Inflammation:** Omega 3 fatty acids may also influence the atherosclerotic process. Again, in patients with coronary heart disease EPA/DHA supplementation versus placebo for two years resulted in modest improvements in atherosclerosis as assessed by angiography [7]. An important recent study of patients awaiting carotid artery surgery randomized cohorts to fish oil capsules, sunflower oil capsules, or controls up until surgery and then assessed morphology of the plaque [49]. Omega 3 fatty acids incorporated into atherosclerotic plaques in the fish oil group, and these plaques were more likely to have reduced mass with less inflammatory infiltrate and increases in thickness of fibrous caps from protective responses. These features imply a plaque that is less vulnerable to rupture and indicates EPA/DHA may help to establish plaque stability. Additional improvements in overall endothelial function and decreases in pro-inflammatory signals have also been noted. The fundamental cellular processes activated or suppressed by omega-3 supplementation and their potential impact on coronary heart disease are active areas of research.

**Cancer:** Epidemiologic studies indicate populations that habitually consume high amounts of EPA/DHA fatty acids also have lower incidences of breast, prostate and colon cancers than those that consume less of these fatty acids in their diets. Many of the mechanisms that are thought to slow or prevent the growth of cancers may also slow or prevent the growth of residual metastatic cancer cells as well [4, 6]. Therefore, increasing the consumption of EPA/DHA from food or supplementation can naturally augment cancer therapy. However, clinical research is not complete in humans. The results of animal studies have demonstrated that the consumption of EPA/DHA can slow the growth of cancer xenografts, increase the efficacy of chemotherapy, and reduce the side effects of chemotherapy [4]. Mechanisms that may be involved include the suppression of cyclooxygenase-2 expression in tumors, decreased AP-1 and ras oncogene levels, and decreased NF-κB activation and bcl-2 expression [34]. Suppressing these would reduce proliferation and angiogenesis and increase apoptosis.

**Type-2 Diabetes:** EPA/DHA supplements may indirectly help prevent the development of type 2 diabetes through modulation of...
lipid metabolism [6]. These effects are likely mediated through transcription factors by decreasing inflammatory NF-kB activity and increasing pro-metabolic PPAR activities [34]. Weight reduction by restriction of total calories, increasing physical activity, and deriving total intake of fats from healthy sources is always advisable. Nutritional causes seem to be the main culprit in this wide-spread epidemic. Nutritional therapy appears to be the main option for treatment [50]. Again, omega-3s may be significant co-therapeutic treatments for lowering triglyceride levels in pre-diabetic and type 2 diabetic patients [50, 51]. However, omega-3 supplements may not directly affect glucose homeostasis [40], yet these essential fatty acids are protective against lipid oxidative stress in diabetic patients [39].

**PREVENTION AND SAFETY**

Current analysis reveals red blood cell levels of EPA/DHA omega-3s in urban Indian populations, and urban populations elsewhere, average between 3% and 4% [5]. Protective, preventative, and therapeutic EPA/DHA levels may require nutritional accumulation to between 6% and 8% [32]. Thus, to achieve nutritional essentiality for omega-3 fatty acids many individuals may need to nearly double their circulating EPA/DHA status.

The topic of benefit versus risk of regular fish consumption and fish oil supplementation is still hotly debated. Various and prevalent concerns exist regarding the safety, sustainability, and predatory species sources of EPA and DHA omega-3 fatty acids from fish. Although fish oil from reputable companies is regarded as safe, long term exposure through supplementation is often feared since trace pollutants from ocean ecosystems contaminate both fresh caught and farm-raised fish that feed on or are fed marine organisms [52, 53]. Some types of fish contain relatively high levels of mercury, polychlorinated biphenyls [PCBs], dioxins and other environmental contaminants [54]. In general, older, larger predatory fish contain the highest level of contaminants. Fish can also contain significant levels of methyl mercury, considered one of the more dangerous food contaminants today [55, 56]. PCBs and methyl mercury are believed to have long half-lives in the body and can accumulate in people who consume fish on a frequent basis. Recommendations currently suggest limiting intake of fish to twice per week.

A study in the journal Diabetes Care links persistent organic pesticide circulation from pollutants in fish to insulin resistance and type-2 diabetes [57]. The study reports the action of these pesticides may be critical during the early stages of diabetes. Similarly, contaminants in fish oil supplements should not be ignored, especially in light of recent mass market trends towards high level daily consumption. Heavy metals, dioxins and PCBs in fish oil supplements and in cod liver oil supplements are documented to occur at persistent low levels [58, 59]. More toxicology studies are needed, particularly with respect to fish based supplements. The possible effects of these trace toxins in the human population over years of exposure from omega-3 fish oil supplementation is not known and a cause for cautious concern. Even though clinically tractable effects of the contaminants found in fish require long term exposure at higher levels than found in supplements, pre-clinical effects cannot be determined or ruled out with confidence at this time.

For DHA-rich microalgae oil produced from culture by the Martek Biosciences Corporation (USA), independent quality control analysis of heavy metals or other pollutants indicate these cannot be detected (Martek unpublished reports, 2006). Thus, one safe and sustainable solution for supplementing long chain omega-3 may be to derive omega-3s from microalgal sources, which are currently the only well developed EPA and DHA omega-3 fish oil alternatives. Microalgae as a safe vegetarian source of EPA and DHA Schizochytrium as an Alternative Source of Dietary Omega-3s: Analysis of vegetarian sources of omega-3 fatty acids inevitably leads to microalgae. Overall, algae are a unique branch of life on planet earth. Most marine EPA and DHA do not originate with fish, but accumulate up the marine food chain from sources like microalgae. Certain non-toxic algal phyla contain high levels of EPA compared to DHA, but commercial development of these has not taken place, possibly due to aquaculture limitations for raising photosynthetic organisms, and higher AA levels may be another concern (Table 1). In comparison, the microalgae grown for making DHA-rich oil has low AA and low EPA levels, but very high DHA levels.

The discussion that follows focuses on the Schizochytrium microalgal strain, which makes high levels of DHA and some EPA (Table 1) [60]. Schizochytrium is a Thraustochytrid, a member of the kingdom Chromista. Schizochytrium is an ancient non-photosynthetic detritus feeding organism that does not assemble into higher ordered structures, as do some photosynthetic green, red, and yellow-green algae. Thraustochytrids form a part of the coastal food chains as a food source for shellfish, which form a significant part of the human diet in coastal regions around the world. Chromista are not related to toxic algae forms, such as some blue-green algae and dinoflagellates, which are in completely separate Kingdoms (Table 1).

Schizochytrium DHA-rich oil has no unpleasant flavor, no detectable environmental pollutants, and may be supplied as oil or in starch powder formulations for cooking, encapsulation, infant milk formula, rice powder, and as additives to cereal and other products (Martek unpublished reports, 2006). This microalgae is rapidly grown in culture where tons of carbon dioxide is removed from the atmosphere for each ton of oil produced, making it environmentally friendly. Martek currently owns the patents to the Schizochytrium production strain and the oil extraction process for making DHA-rich oil. Martek acquired these patents by their purchase of OmegaTech. Trademarked as ‘Life’sDHA’, this product is available on the market in the form of supplements. DHA-rich oil could become available in countries like India with increasing market demand due to its large vegetarian population. India’s current unmet needs for omega-3 supplementation may include preventative treatment for conditions such as heart disease [5], and dyslipidemia in pre-diabetic or type 2 diabetes patients [51, 52], but omega-3 supplements may not affect, or may only indirectly affect, insulin signaling and glucose homeostasis [40].

**Schizochytrium Oil Safety and Efficacy:** DHA-rich oil produced from Schizochytrium has undergone extensive analysis, showing that the individual components of the extracted oil are all present elsewhere in normal food consumed by communities. Thus, DHA-rich oil is inherently safe in its fatty acid and sterol components (Table 2) (OmegaTech unpublished reports, 1997). Safety is further supported by the historically safe use of fish oils of similar composition. In addition, its safety is also based on the small quantities expected to be consumed per dose, the knowledge of the metabolism of individual lipid components and the lack of published reports of inherent toxic effects or thyroid problems, or any adverse allergic reactions. However, there have been no reports in the literature of allergic responses to any members of the kingdom Chromista, including the Thraustochytrids. Allergic re-

**Table 1. Long Chain Polyunsaturated Fatty Acids of Some Non-Toxic Algae Versus Fish Oil Supplements**

<table>
<thead>
<tr>
<th>Kingdom</th>
<th>Phyla</th>
<th>Species Name (Common)</th>
<th>DHA</th>
<th>EPA</th>
<th>AA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plantae</td>
<td>Chlorophyceae</td>
<td>(Green Algae)</td>
<td>0%</td>
<td>2.5%</td>
<td>0.1%</td>
</tr>
<tr>
<td>Plantae</td>
<td>Rhodophyceae</td>
<td>(Red Algae)</td>
<td>0%</td>
<td>20%</td>
<td>4%</td>
</tr>
<tr>
<td>Chromista</td>
<td>Heterokontae</td>
<td>(Yellow-Green)</td>
<td>0.5%</td>
<td>27%</td>
<td>6%</td>
</tr>
<tr>
<td>Chromista</td>
<td>Heterokontae</td>
<td>Schizochytrium</td>
<td>37.4%</td>
<td>2.8%</td>
<td>1.0%</td>
</tr>
</tbody>
</table>

**Microalgae as Alternative Omega-3 Source**

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Table 2. Fatty Acid Species and Sterol Content of Microalgae Schizochytrium DHA-Rich Oil

<table>
<thead>
<tr>
<th>Species</th>
<th>Omega</th>
<th>% Total Oil</th>
<th>% Sterols in Oil</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myristate</td>
<td></td>
<td>10.8</td>
<td></td>
</tr>
<tr>
<td>Palmitate</td>
<td></td>
<td>25.2</td>
<td></td>
</tr>
<tr>
<td>Palmitoleate</td>
<td></td>
<td>1.9</td>
<td></td>
</tr>
<tr>
<td>DH-GLA</td>
<td>ω 6</td>
<td>2.4</td>
<td></td>
</tr>
<tr>
<td>Arachidonate</td>
<td>ω 6</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>EPA</td>
<td>ω 3</td>
<td>2.8</td>
<td></td>
</tr>
<tr>
<td>DPA</td>
<td>ω 6</td>
<td>14.4</td>
<td></td>
</tr>
<tr>
<td>DHA</td>
<td>ω 3</td>
<td>37.7</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>ω 6</td>
<td>17.8</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>ω 3</td>
<td>40.2</td>
<td></td>
</tr>
<tr>
<td>Ratio</td>
<td>ω 3/ω 6 = 2.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cholesterol</td>
<td></td>
<td>0.8 (8 mg/g)</td>
<td></td>
</tr>
<tr>
<td>Total Sterols</td>
<td></td>
<td>3.1 (31 mg/g)</td>
<td></td>
</tr>
</tbody>
</table>

responses by humans to microorganisms can sometimes be related to microbial toxins, but only certain types of algae are allergenic.

It has been shown that individual ratios of polyunsaturated fatty acids vary according to food source. For DHA-rich Schizochytrium oil the key points to note are very high levels of DHA, the low levels of EPA and moderate content of docosapentaenoic acid (DPA n-6) (Table 2) [60]. DPA n-6 is an omega-6 fatty acid that does not have the same bioactivity as AA. It is present in a wide variety of foods and is relatively abundant in eggs and breast milk. The ratio of DPA n-6 to DHA n-3 in human breast milk is reported to range normally from 1:1 to 1:6 [61]. The ratio of DPA n-6 to DHA n-3 in DHA-rich Schizochytrium oil is 1:3, the median range of breast milk. Schizochytrium microalgae strains have been developed by conventional techniques and no Genetic Modifications were used or needed. An independent panel of experts in the US has concluded that DHA-rich oil from Schizochytrium microalgae can be "Generally Regarded as Safe" as a nutritional food ingredient (OmegaTech unpublished reports, 1997). When DHA-rich Schizochytrium oil and fish oil were used in cell viability and proliferation tests with human colon adenocarcinoma Caco-2 cells, tests showed no differences between algal oil and fish oil, indicating safety and potency [62].

One question is whether DHA-rich microalgae oil can function as a universal fish oil alternative? Based on the effects and benefits of fish oil, it will be important to know how well DHA-rich oil compares in clinical efficacy. Early clinical indications strongly support the efficacy of microalgae oil compared to fish oil.

FUTURE DIRECTIONS

Although advances in our knowledge of the protective effects of EPA/DHA have increased, many issues remain. Additional biochemical understanding of the individual and/or overlapping roles of EPA and DHA are important scientifically. At the same time, long-term risk factors associated with fish oil supplementation are not known regarding pollutant and heavy metal accumulations in the body. Safe, sustainable alternatives to fish oil EPA/DHA capsules are available, but education and awareness of this option is limited. Double blind, placebo controlled trials of microalgae oils are also needed in more studies, particularly with life threatening heart conditions. Oil formulations of at least two microalgae classes may be considered as Schizochytrium mainly provides DHA while other microalgae classes are enriched with EPA. However, DHA retro-conversion in the body is a notable topic for making comparisons to fish oil supplements. Additionally, a standardized blood test for EPA/DHA levels is needed for diagnostic purposes to determine inherent deficiencies in individuals and to assign proper regimens. For general use, EPA/DHA fatty acids should be named vitamin-F, as was the practice with fish oil formulations in the early part of the 1900s. Currently, pharmaceutical and agricultural industries are rapidly increasing their research and development investments in omega-3 fatty acids for production of genetically modified plant organisms and for prescription of highly purified fish oil tablets [63]. All of these efforts are likely helpful, but controversial. Finally, individuals must carefully choose omega-3 fatty acid sources to suit their needs, keeping in mind that plant sources are healthy, but not fully preventative or therapeutic. Beyond omega-3s, supplementation as a single treatment of any disease is unwise. Always consult a physician and definitely include weight loss and a healthy diet as needed. In general, omega-3s are at minimum essential, but these may even be co-therapeutic. Since many individuals may be deficient in their total accumulated omega-3 status, global health education about these important nutrients and their sources will benefit public health.

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