Case studies demonstrate that oral intake of omega-3 polyunsaturated fatty acids from pharmaceutical-grade fish oil supplements results in pain reduction and functional improvement in patients with neuropathic pain.

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The literature of the past twenty years contains numerous studies and clinical investigations that highlight the critical role played by omega-3 polyunsaturated fatty acids (PUFAs) in human health and disease. Omega-3 fatty acids are essential for growth and development and play a vital role in the prevention and treatment of cardiovascular disease, inflammatory and autoimmune disorders, cancer, diabetes, and depression.1

In this paper we will review the etiology, symptoms, and treatment of neuropathic pain; review the literature on the reported benefits of omega-3 fatty acids; present case studies of patients suffering from neuropathic pain treated with omega-3 FA; and provide some practical application recommendations for a clinical practice.

Neuropathic Pain Etiology
Neuropathic pain (NeP) is defined as pain caused by a lesion of the peripheral or central nervous system (or both) manifesting with sensory symptoms and signs. Current estimates suggest that this devastating condition may affect up to 3% of the population.2 NeP is costly to the health care system. In the U.S. alone, the costs associated with this disorder are estimated at $40 billion annually.3

NeP can be caused by trauma, inflammation, transection, nerve compression, ischemia or metabolic injury to neuronal cell bodies. NeP can also be caused by cancer, diabetes, multiple sclerosis (MS), Parkinson disease, infectious agents (e.g., HIV-1) or by the toxic side effects of various drug regimens.4,5

In states of NeP, sensory (nociceptive) neurons damaged by disease, injury or drugs discharge spontaneously and leads to sustained levels of excitability. These abnormal discharges “cross-talk” with adjacent uninjured nerve fibers, resulting in amplification of pain impulses which causes peripheral sensitization. In turn, central neurons innervated by such nociceptors undergo dramatic functional changes including a state of hyperexcitability termed central sensitization.6 This heightened activity is thought to result from increased neuronal expression and activation of ion channels, such as voltage-gated sodium channels (VGSCs), and receptors that initiate and mediate the abnormal generation of action potentials and synaptic transmission in pain pathways.7

To summarize, following injury to sensory nerves, nociceptor-driven activity in the spinal cord becomes divorced from normal physiology, so that pain is produced in the absence of any appropriate stimulus and results in NeP.8

Neuropathic Pain Symptoms and Treatment
Patients suffering from NeP experience a wide variety of symptoms. Some examples include spontaneous paresthesias and dysesthesias manifesting as abnormal sensations including crawling, numbness, itching, and tingling. Pain resulting from this disorder can be divided into two categories: stimulus-evoked pain and stimulus-independent (spontaneous) pain.9 Stimulus-evoked pain is associated with different types of hypersensitive
pain behavior, including allodynia and hyperalgesia. Spontaneous pain can be either constant (e.g., burning) or intermittent (shooting, electric shock-like) and most patients describe having both.

From a therapeutic standpoint, NeP is a difficult disorder to treat. Animal models of chronic NeP induced by spinal root ligation or sciatic nerve constriction show that prostaglandins are required to initiate the NeP process, but are not necessary for its maintenance. Therefore, nonsteroidal anti-inflammatory drugs (NSAIDs) such as aspirin and ibuprofen have limited efficacy in treating NeP. Furthermore, the use of opiates to alleviate NeP is challenging because of the high doses that are often required which effectively narrow the therapeutic index. Other drugs, such as anticonvulsants and tricyclic antidepressants, may have some limited use but they are associated with significant adverse effects.

**Omega-3 Fatty Acids**

**Structure, Sources and Dietary Intake.**

Omega-3 (or n-3; ω-3) fatty acids are long-chain PUFAs of plant and animal origin, that are typically 18, 20, or 22 carbon atoms in chain length. The term "ω-3" signifies that the first double bond in the molecule is located at the third carbon position counting from the ω-end of the fatty acid chain.

Fish oil from oily fish is a rich source of long chain n-3 PUFAs, consisting mainly of eicosapentaenoic acid (EPA; 20:5n-3) and docosahexaenoic acid (DHA; 22:6n-3) (see Figure 1). Vegetable oils are not a source of EPA or DHA although certain types (e.g., flaxseed and walnut oil) do contain varying amounts of an alternate form of omega-3 fatty acid known as alpha-linolenic acid (ALA; 18:3n-3). ALA can be metabolized by the body into the longer chain EPA and DHA via a series of desaturation/elongation reactions (see Figure 2) but, unlike rats, humans can only convert a small amount (5%) of ALA into EPA.

Omega-6 (or n-6; ω-6) fatty acids, such as linoleic acid (LA; 18:2n-6) can be found in abundance in many vegetable oils (e.g., safflower, sunflower, corn, cottonseed, soybean). LA can be converted by the body into the longer chain arachidonic acid (AA; 20:4n-6). Arachidonic acid is known to lead into pro-inflammatory eicosanoids.

Both n-3 and n-6 PUFAs are used for phospholipid production and are thus components of cell membranes throughout the body, contributing to the physical and functional properties of those membranes. In addition, n-3 and n-6 PUFAs serve as precursors to eicosanoids which are key mediators and regulators of various physiological processes such as inflammation, vascular tone, and hemostasis.

Since the dawn of the industrial age, the dietary intake of n-6 fatty acids has steadily increased—particularly in Western diets—to the point where they currently represent the principal dietary source of PUFAs. By contrast, the dietary intake of omega-3 fatty acids has dramatically declined in Western countries over the last 100 years (see Figure 3). This is due, in part, to the vast consumption of omega-6 rich vegetable oils and products from animals fed with grains containing n-6 PUFAs. It has been suggested that the ideal dietary ratio of n-6 to n-3 fatty acids be approximately 1:2:1. However, in the typical North American diet, the n-6:n-3 dietary ratio is about 8:1 and, in some instances, may be as high as 20:30:1. This great discrepancy between n-6 and n-3 PUFAs intake is not without consequence. It is thought that the elevated n-6:n-3 ratio most likely

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**Figure 1.** Molecular structures of Omega-3 fatty acids.

- **EPA, Eicosapentaenoic Acid, 20:5ω-3** (contains 20-carbon atoms and 5 double bonds/unsaturation sites)
- **DHA, Docosahexaenoic Acid, 22:6ω-3** (contains 22-carbon atoms and 6 double bonds)

**Figure 2.** Omega-3 fatty acid pathway.

- **a-Linolenic acid** 18:3ω-3 (canola, flaxseed, soybean)
- **Desaturation**
  - 18:4ω-3
  - 20:4ω-3
  - 20:5ω-3 (EPA)
  - 20:6ω-3 (DHA)
- **Elongation**
  - 22:5ω-3 (EPA)
  - 22:6ω-3 (DHA)

**Figure 3.** Changes in fat consumption throughout human history: note the dramatic rise in trans-fats, omega-6, saturated fats. From The Omega Plan by Artemis P. Simopoulos MD and Jo Robinson (with permission).
contributes to an increased incidence of cardiovascular disease (CVD), inflammatory disorders, autoimmune diseases, major depression, and cancer.  

**PUFAs and Eicosanoid Metabolism**

The n-3 and n-6 fatty acids are chemically and metabolically distinct and have contrasting physiological functions (see Figure 4). The eicosanoid metabolic products synthesized from AA, namely prostaglandins (PGs), thromboxanes (TXs), leukotrienes (LTs), hydroxy fatty acids and lipoxins are formed in larger quantities than those formed from n-3 PUFAs—such as EPA—because of the increased amounts of n-6 PUFAs in the Western diet. Eicosanoids derived from AA are biologically active in small quantities and, if they are formed in inordinate amounts, they contribute to the formation of atheromas and thrombi. In addition, they lead to the development of allergic and inflammatory disorders and to cell proliferation.

When humans increase their consumption of EPA and DHA, from oily fish or from fish oil supplements (liquids or capsules), an increased proportion of these fatty acids are found in the cell membranes of inflammatory cells in particular. EPA and DHA incorporation into these membranes occurs in a dose-response fashion and is partly at the expense of AA. Hence, since there is less AA available for eicosanoid biosynthesis by the cyclooxygenase (COX) and lipoxygenase (LOX) enzymes, fish oil supplementation of the diet results in a decreased production of the proinflammatory prostaglandin E2 (PGE2), thromboxane A2 (TXA2), TXB2, leukotriene B4 (LTB4), LTE4 and 5-hydroxyeicosatetraenoic acid, but not of prostacyclin (PGI2).

EPA can also act as a substrate for both COX and LOX enzymes and thereby compete with AA for prostaglandin and leukotriene synthesis. EPA gives rise to the 3-series PGs and TXs (such as TXA3, PGE3, PGF3) and to the 5-series LTs (LTB5, LTE5) and 5-hydroxyeicosapentaenoic acid, but not of prostacyclin 12 (PGI2).

Eicosapentaenoic acid (EPA) C20:5 n-3

Docosahexaenoic acid (DHA) C22:6 n-3

**FIGURE 4. Pathways to inflammation: omega-6 is pro-inflammatory and omega-3 is anti-inflammatory.**

TXA3 is a weaker platelet aggregator and vasoconstrictor than TXA2. Recent studies have shown that EPA and DHA also give rise to resolvins (from EPA and DHA) and docosanoids (from DHA) through pathways involving COX and LOX enzymes. In cell culture and animal-feeding studies, these novel mediators were demonstrated to be anti-inflammatory, inflammation resolving, and immunomodulatory.

Some of the effects of n-3 PUFAs are elicited by eicosanoid-independent mechanisms. Studies have shown, for example, that the intake of dietary fish oil results in decreased leukocyte chemotaxis, decreased expression of adhesion molecules and decreased generation of reactive oxygen species. n-3 PUFAs have also been shown to inhibit the production of proinflammatory cytokines such as tumor necrosis factor-α (TNF-α), interleukin-1β (IL-1β), IL-6 and IL-8, in both cell culture studies and in human trials. Additionally, n-3 PUFAs might exert their effects on inflammatory gene expression through direct action on intracellular signaling pathways which lead to activation of one or more transcription factors such as nuclear factor kappa B (NF-κB).

**n-3 PUFAs and Chronic Disease**

The benefits of n-3 PUFAs supplementation are well documented in the literature for the prevention and management of a wide variety of health conditions including inflammatory and autoimmune diseases, cardiovascular disease, and depression. Less is known, however, about the use and efficacy of these fatty acids in the treatment of other disorders such as fibromyalgia syndrome and neuropathic pain.

**n-3 PUFAs and Inflammatory/Autoimmune Diseases**

A number of clinical trials have been conducted assessing the benefits of dietary supplementation with n-3 PUFAs...
in inflammatory and autoimmune disease in humans, including RA, Crohn's diseases, ulcerative colitis, psoriasis, lupus erythematosus, MS, and migraine headaches. A recent meta-analysis of 17 randomized controlled trials (RCTs) was conducted by Goldberg et al. to assess the pain relieving effects of n-3 PUFA supplementation in patients with rheumatoid arthritis (RA) or joint pain secondary to inflammatory bowel disease and dysmenorrhea. Results from the analysis showed that n-3 PUFA supplementation for 3-4 months reduces patient-reported joint pain intensity, minutes of morning stiffness, number of painful and/or tender joints, and NSAID consumption.

Maroon et al. conducted a non-placebo controlled study to determine if patients could effectively substitute fish oil as an anti-inflammatory for NSAIDs. The study included 250 participants who had been seen by a neurosurgeon and were found to have chronic, nonsurgical, neck or back pain. After taking n-3 PUFA supplements for an average of 75 days, 59% discontinued their use of prescription NSAIDs and 60% stated that their overall pain was improved. Moreover, 88% stated they would continue to take the n-3 PUFA supplements. No significant adverse effects were reported. These results corroborate other controlled trials that compared ibuprofen and n-3 PUFA supplementation demonstrating equivalent effects in reducing arthritic pain. Furthermore, such findings suggest that n-3 PUFA supplements may be a safe alternative to NSAIDs, particularly the COX-2 inhibitors which have been associated with extreme complications such as gastric ulcers, bleeding, blood pressure increase, myocardial infarction, and even death.

n-3 PUFA and Cardiovascular Disease
Numerous intervention and clinical trials support the use of n-3 PUFA supplements in the reduction of various risk factors for CVD, myocardial infarction, and sudden cardiac death. Some of the cardiovascular benefits that have been associated with n-3 PUFA supplementation include reduction in ventricular arrhythmias, increase in heart rate variability, antithrombotic and other effects on the hemostatic system (moderately longer bleeding times, reduced plasma viscosity), lipid lowering, improved endothelial relaxation, inhibitory effect on athero-sclerosis and inflammation, and suppressed production of inflammatory cytokines (interleukins and TNF) and mitogens. The anti-arrhythmic effect of n-3 PUFA supplementation is thought to be the main benefit associated with their use. However, this effect has not been clearly demonstrated in clinical trials. A meta-analysis of 3 RCTs was conducted by Jenkins et al. to determine the effects of fish oil supplementation in patients with implantable cardioverter defibrillators who were at risk of ventricular arrhythmias. They also wanted to determine if there was significant heterogeneity between trials. Meta-analysis of data collected at one year showed no overall effect of fish oil on the relative risk of implantable cardioverter defibrillator discharge. Results also showed significant heterogeneity between trials. Furthermore, patient response to fish oil supplementation was heterogeneous with one study showing a significant benefit of fish oil and another an adverse tendency.

n-3 PUFA and Depression
Mounting evidence from laboratory, epidemiological, and clinical studies suggests that n-3 PUFA supplementation may play a role in the prevention and management of depression. For the time being, the bulk of clinical evidence indicates that EPA may be most important in mood stability and that relatively low levels are required for successful outcomes. The beneficial effects of n-3 PUFA supplementation in depression may be due in part to the modulation of brain-derived neurotrophic factor (BDNF). cAMP response element binding protein (CREB) upregulation, via n-3 PUFA mediated inhibition of PGE2 and IL-1β, may activate the downstream target BDNF. BDNF is known to be lower in depressed patients and thus may play an important role in major depression.

n-3 PUFA and Fibromyalgia Syndrome
Some clinical evidence suggests n-3 PUFA supplementation may be of use in the management of fibromyalgia. Ozgocmen et al. investigated the effect of n-3 PUFA supplementation in the management of fibromyalgia syndrome in an open, non-controlled single-blind study involving 12 female patients. The patients were treated for a period of 4 weeks with high doses of n-3 PUFA supplements. Results from the study showed statistically significant beneficial changes from baseline for tender point counts, chest expansion measurements and pain severity, fatigue, and depression scales that were evaluated using the Fibromyalgia Impact Questionnaire.

n-3 PUFA and Neuropathic Pain
To date, there are no clinical trials that have examined the effects of n-3 PUFA supplementation in the treatment of neuropathic pain patients. Moreover, very few studies have investigated the mechanism(s) whereby n-3 PUFA supplementation may modulate NeuP mechanisms. There are considerable differences between chronic NeuP and chronic inflammatory pain. The eicosanoid-dependent anti-inflammatory effects of n-3 PUFA supplementation may not be relevant to NeuP conditions. A significant factor in NeuP is the activation in the spinal cord of non-neural glial cells, macroglia, and astrocytes. Activated glia are characterized by proliferation, hyper trophy and increased production of inflammatory cytokines such as IL-1β, IL-6 and TNF-α. EPA and DHA could possibly reduce the production of these cytokines but this remains to be determined. Specifically, DHA has recently been shown to play a larger role in neurogenic inflammation which was not previously anticipated.

It has been suggested that n-3 PUFA supplementation may block pain neuron voltage-gated sodium channels (VGSCs) which underlie NeuP. The gene encoding one of the nociceptor specific VGSCs, SNS/PN3, shares a very similar genomic structure with the human cardiac VGSC gene. n-3 PUFA supplementation is known to potentially and reversibly bind to and block current through this cardiac VGSC. Outside of their neurogenic influences, some evidence suggests a role in modulation of the stress response through influence of plasma cortisol. Low plasma cortisol, which is required to blunt the inflammatory process and therefore influence the inflammatory component of NeuP, is associated with an increased incidence of neuropathic pain patients who improved with the use of omega-3 fatty acid supplements. This case series is the first ever published on using omega-3 fatty acids in the treatment of neuropathic pain (and using updated criteria in diagnosis such as the DN4 and Pain Detect Questionnaires).
Omega-3 Fatty Acids and Neuropathic Pain

Patient 1: C7 Radiculopathy
A 53-year-old left-handed police officer, married father of two, was diagnosed with right-sided cervical radiculopathy. He had developed neck symptoms in 2004. His symptoms worsened to the point where by December of 2005 he could no longer play hockey and had sleeping problems. MRI results showed evidence of a C6/C7 right lateral disc herniation compressing the right C7 nerve root with spinal stenosis and multi-level degenerative disc disease. Past medical history included anxiety/panic attacks, depression, gout, and vasectomy.

Previous treatments included: physiotherapy, which was passive and did not focus on core stability training; prescribed pain-killers (Naproxen), and other NSAIDs such as Advil. His other medications included Diltiazem, Lossec, ECASA (enteric coated acetylsalicylic acid) and Tylenol-3.

"He was started on a treatment of high doses of n-3 PUFAs (8 capsules/day with EPA-DHA total dose of 4800mg/day). After two and a half weeks his pain started to subside. Strength measurements were done and there was noticeable improvement in his C7 innervated muscles in the right arm."

Physical exam (July 2006) revealed a height of 6'1" with a weight of 250 lbs. Blood pressure (BP) was 106/77 mmHg. Biomechanical examination revealed a marked head-forward posture with tight pectorals and poor core stability. Neurological examination revealed a possible right rotator cuff impairment as well as vasculogenic thoracic outlet compression which also would perpetuate symptoms. Additionally, he had a positive Allen’s test and weakness in the C6/C7 myotomes.

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Prior to taking n-3 PUFAs his SF-MPQ (short form McGill pain questionnaire) rating was 30/45.

He was started on a treatment of high doses of n-3 PUFAs (8 capsules/day with EPA-DHA total dose of 4800mg/day). After two and a half weeks his pain started to subside. Strength measurements were done and there was noticeable improvement in his C7 innervated muscles in the right arm. In fact, his triceps strength on the right side was greater than the left side, measured at 125 lbs and 123 lbs respectively, with the Lafayette manual muscle tester. Jamar grip strength was also improved: Rt 39.9 lbs and Lt 42 lbs. After taking n-3 PUFAs for a little over eight months, his SF-MPQ decreased to 0/45 and he started playing hockey and working out again.

He returned back on Feb 21 2008. He was still benefiting from the n-3 PUFAs (his NRS pain score was 0/10 and SFMC-Gill 4/45). Weight and blood pressure were unchanged. He cited a time when he forgot to bring his n-3 PUFAs to a hockey tournament and noted recurrence of neck pain after 4 days. Otherwise, he continues to be pain free and able to play full equipment ice hockey.

Patient 2: Thoracic Outlet Syndrome
A 48-year-old right-handed registered nurse, married mother of 2, was diagnosed with left lateral epicondylalgia. Her work, in the continuing care department of the hospital, involves the transfer of heavy patients. She injured her left arm while transferring a 245 lb patient.

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Her physical exam revealed a height of 5'4" with a weight of 125 lbs. BP was 138/88 mmHg. Physical examination revealed left knee pain in both medial and lateral compartments of the knee, slightly more severe in lateral parts. Knee range of motion was 150° bilaterally with end range pain and flexion of the knee. Range of motion was adequate with mild effusion noted in the left knee and mild signs of synovial thickening. Neurological testing revealed normal motor strength at 5/5 and a DN4 score of 5/10 (burning pain, cold, paresthesia, numbness, mild brush allodynia, pinprick hypoaesthesia). His average pain was rated at 7/10 with a best of 3/10 and worst 8/10. His SF-MPQ was 11/45.

He was started on a treatment of high doses of n-3 PUFAs (8 capsules/day) and began supplementing with vitamin D and glucosamine. He reported dramatic improvement in range of motion, tolerance for physical activity, and reduced swelling after aggravating activities following taking n-3 PUFAs for six months.
Patient 4: Cervical Radiculopathy and Carpal Tunnel Syndrome

A 50-year-old right-handed Holter Monitor Company representative, married father of two, was diagnosed with chronic right C7 radiculopathy/carpal tunnel syndrome. In 2003, he was involved in a motor vehicle accident. MRI results revealed evidence of right central disc protrusion at C6/C7. There was also evidence of severe spinal stenosis at C5/C6 and moderate stenosis at C4/C5 and C6/C7. He reported limitations in such areas as, self-care, household responsibilities, social activity, recreation, sports, grip, lifting from floor to waist, and lifting overhead.

Past treatments included physiotherapy, chiropractic treatment, massage therapy and occasional NSAIDS. He also supplemented with B-vitamins and Coenzyme Q10.

Physical exam revealed a height of 5'9 1/2" with a weight of 240 lbs. BP was 142/95 mmHg. His jamar grip was: Lt 10/35 and Rt 10/35. He showed evidence, both electrodiagnostically (moderately prolonged median sensory and motor latencies, 2+ denervation in C7 myotomes) and on clinical exam, of weakness in the right arm. His SF-MPQ 5'9 1/2" with a weight of 240 lbs. BP was 142/95 mmHg. His jamar grip was: Lt 10/35 and Rt 10/35.

He was started on a treatment of high calories and patients are advised to adjust their diet and insulin accordingly (long-term use of omega-3, however, does reduce insulin resistance and improves diabetic control).

Because of the "blood-thinning" effects of omega-3, we usually advise patients to stop such supplements, as well as herbal products such as ginkgo, curcumin, ginger, two weeks prior to any surgery, dental work, and invasive procedures such as a colonoscopy.

Lab work should be done to follow patients on high dose omega-3. This includes markers of "silent inflammation" and include the Arachidonic acid to Eicosapentaenoic acid ratio (AA:EPA ratio). The average North American ratio is 12:1. An optimal ratio for cardiovascular health is 1.5:3:1. Excess intake of omega-3 with a ratio of 0.5:1 is associated with an increased risk for hemorrhagic stroke.

Unfortunately, such lab testing is expensive (about $150) and most of our patients did not undergo such testing unless they were taking extremely high doses—7.5-11 gm EPA/DHA or more per day. Testing is available at a lab such as Nutrasure Diagnostics, Inc., Ontario, Canada. This lab measures the serum phospholipid levels which is more accurate, and more studied, than red blood cell (RBC) levels. Other useful lab tests to detect silent inflammation include the HS-CRP (optimal levels are < 1.0), fasting insulin (optimal is < 10 uIU/ml) and TG-HDL ratio (optimal is < 2). The references and research for this are summarized nicely in chapters 4 and 7 in the "Anti-inflammation Zone" book by Barry Sears, Ph.D.

It is important to recommend a high quality brand of omega-3. Patients are taught to read labels and ensure that products have good potency—one gets more for the money in a capsule or teaspoon that has a higher concentration of EPA/DHA—and has been tested for impurities. Websites such as the www.ifosprogram.com will list omega-3 products that have been independently studied and have been independently found to be beneficial.
We often combine omega-3 FA and other pregabalin. For opioid-resistant neurotrauma. For severe neuropathic pain (NRS pain >6/10), we combine omega-3 FA with pregabalin. For opioid-resistant neuropathic pain, pharmaceutical cannabinoids are also helpful (nabilone, sativex spray).

Long-lasting lifestyle changes need to be adopted to ensure long-term relief of pain. This includes appropriate exercise, both cardio and core strengthening, weight-loss, stress reduction (prayer, meditation, humor), and good sleep hygiene. Efforts to detoxify the body of unhealthy "toxic" substances, such as trans-fats, and unhealthy "talk-sick" attitudes and behaviour are all important.

Conclusion

Case studies using omega-3 FA supplementation for neuropathic pain—in a variety of patient presentations—has demonstrated the efficacy of this modality. While pain questionnaires were utilized in documenting outcome measures, further research in the way of randomized double-blind controlled trials would be needed to validate the use of omega-3 fatty acids for neuropathic pain. We hope this article will stimulate such research and lead to greater pain-free wellness in our patients.

References

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5. Leigh Arsenault, BSc is a naturopathic doctor and consultant at the Canadian Centre for Integrative Medicine.

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Lab tested for contaminants such as heavy metals—including mercury—PCBs, and dioxin. The standards set by IFSO for ultra-refined EPA/DHA concentrate are very rigorous with upper limits set as follows: mercury <10 parts per billion (ppb), PCBs <45 ppb, dioxin <1 part per trillion, total oxidation <13 meq/L.

Our preferred products, based on purity and patient compliance (taste), are the SuperEFA liquid (Seroyal Inc) and the See Yourself Well capsules (Nutratec). A recommended conservative dose is 2700mg of EPA + DHA, based on the Goldberg meta-analysis. However, a more aggressive approach for more severe pain can be up to 7500mg EPA + DHA. The latter approach will require serum lab tests to monitor the AA-EPA ratio.

For patients who experience stomach difficulties or nausea from the use of omega-3, we usually advise them to try freezing the capsules. A better response occurs with enteric coated capsules for which we recommend the Metagenics' EPA-DHA extra strength product. Digestion is better when omega-3 are taken with food. It is also useful to split the dosage through several meals instead of all at once.

Instruct patients clearly to take only omega-3 and not omega-3-6-9. As noted previously, omega-3, omega-6's are help in ameliorating pain. Omega 6's are deficient in vitamin B6, magnesium, zinc and have excessive trans-fatty acids, and unhealthy "talk-sick" attitudes and behaviour are all important.

Remember that omega-3 FA are just one component of an overall integrative medical approach in treating pain and optimizing wellness. Patients must learn to improve their diets and reduce arachidonic acid sources such as too much red meat and fried foods. Diets that are deficient in vitamin B6, magnesium, zinc and have excessive trans-fatty acids, caffeine will have impaired delta-6 desaturation activity (needed to convert alpha-linolenic acid in the pathway towards EPA). We often combine omega-3 FA and other nutraceuticals with judicious courses of anti-inflammatory drugs (cyclooxygenase 2 inhibitors) for post-surgical and post-musculoskeletal trauma. For severe neuropathic pain (NRS pain >6/10), we combine omega-3 FA with pregabalin. For opioid-resistant neuropathic pain, pharmaceutical cannabinoids are also helpful (nabilone, sativex spray).

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