Solvent	Saturated	Unsaturated			ບ	nidentified
		Mono	Di	Poly	H-Poly	
chio-meth	26,18	22,54	1,91	4,31	26,34	18,72
acetone	21,4	22,18	1,75	4,67	24,52	25,49
acetone	19,09	22,11	2,03	4,79	30,24	21,72
ethanol	45,93	22,96	1,23	2,72	11,11	16,05 (500 µg/mL)
	45,96	22,98	1,24	2,48	11,18	16,15 (200 µg/mL)

Table 10: Fatty acid composition E. pacifica

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Data expressed in percentage of total fatty acids (%).

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TABLE 11. OPTIMAL CONDITIONS FOR LIPID EXTRACTION OF AQUATIC ANIMAL TISSUES (suggested procedure)

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STEP	CONDITIONS
Grinding (if particles > 5mm)	4°C
Lipid extraction	sample-acetone ratio of 1:6 (w/v) 2h (including swirling 20 min) 4°C
Filtration	organic solvent resistant filter under reduced pressure
Washing	sample-acetone ratio of 1:2 (w/v) pure and cold acetone
Filtration	organic solvent resistant filter under reduced pressure
Evaporation	under reduced pressure
Oil-water separation	4°C
Lipid extraction	sample-ethanol ratio of 1:2 (w/v) pure ethanol 30 min 4°C
Filtration	organic solvent resistant filter under reduced pressure
Evaporation .	under reduced pressure

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Although the present invention has been described herein above by way of preferred embodiments thereof, it can be modified, without departing from the spirit and nature of the subject invention as defined in the appended claims.

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WHAT IS CLAIMED IS:

and

1. A method for extracting lipids from an aquatic animal tissue comprising the steps of:

a) suspending said animal aquatic tissue in an organic solvent;

b) extracting lipids by successive organic solvent treatment;

c) collecting said lipids in a first fraction and an organic insoluble fraction.

2. The method of claim 1, wherein said organic solvent of a) is acetone.

3. The method of claim 1 or 2, wherein said organic solvent of b) is selected from at least one of acetone and alcohol.

4. The method of claim 1, 2 or 3, wherein said organic insoluble fraction comprises a dry residue fraction which is enriched in protein.

5. The method of claim 1, 2, 3 or 4, wherein said aquatic animal tissue is at least one tissue selected from the group consisting of krill tissue, *Calanus* tissue and fish tissue.

6. A lipid extract obtained by the method of claim 2, 3, 4 or 5.

- 7. A protein rich fraction obtained by the method of claim 4 or 5.
- 8. A lipid extract having the properties in accordance with the present invention.

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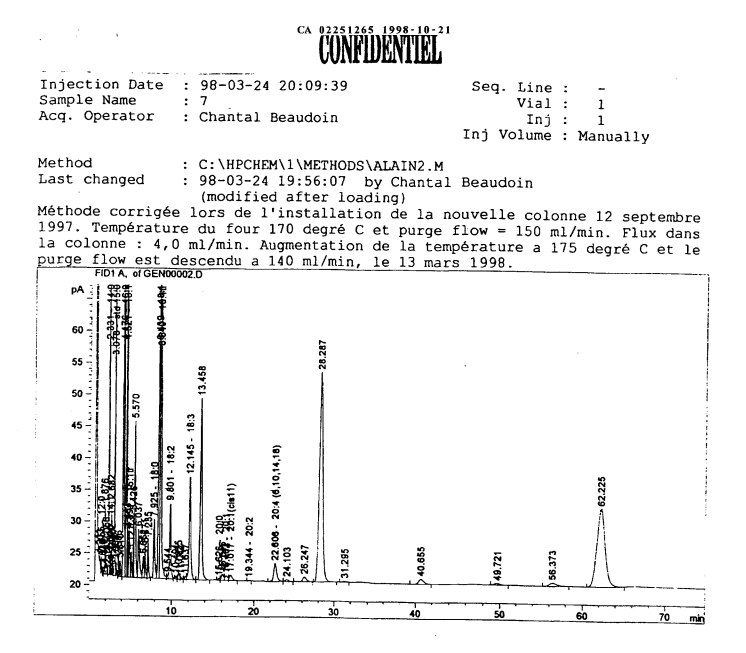


Figure 1: Gas-liquid chromatography of fatty acids from dry krill (chloroformmethanol).

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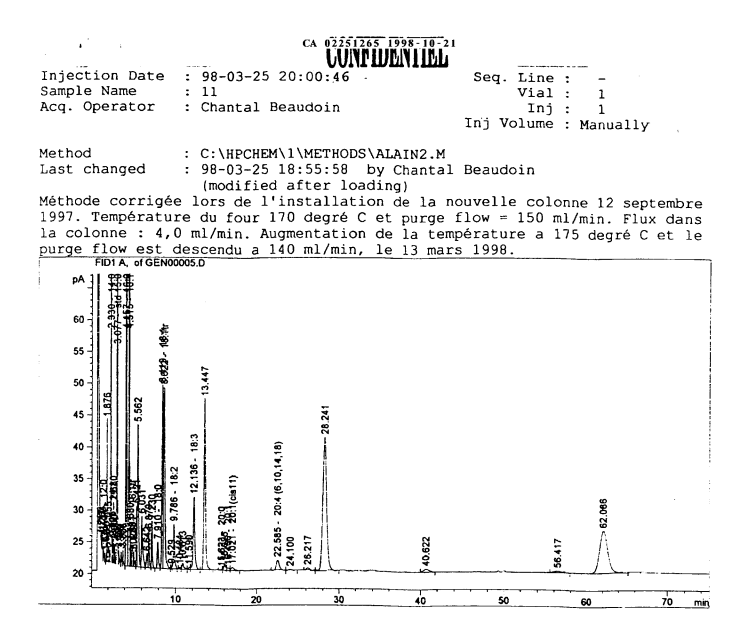


Figure 2: Gas-liquid chromatography of fatty acids from dry krill (acetone).

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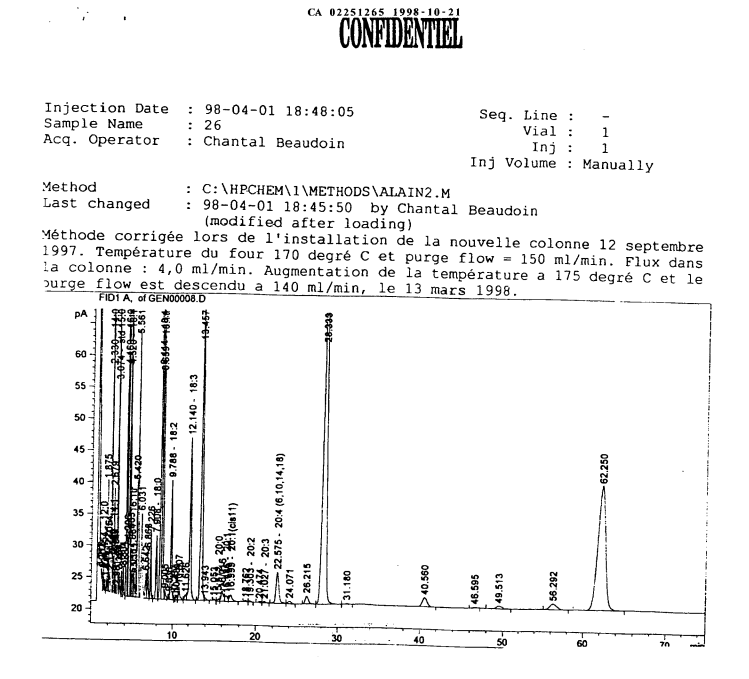


Figure 3: Gas-liquid chromatography of fatty acids from frozen krill (acetone).

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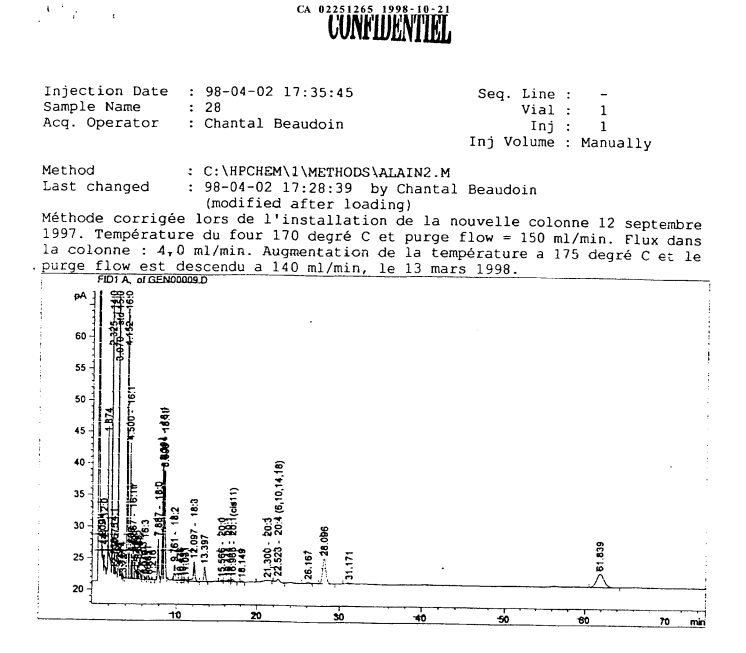
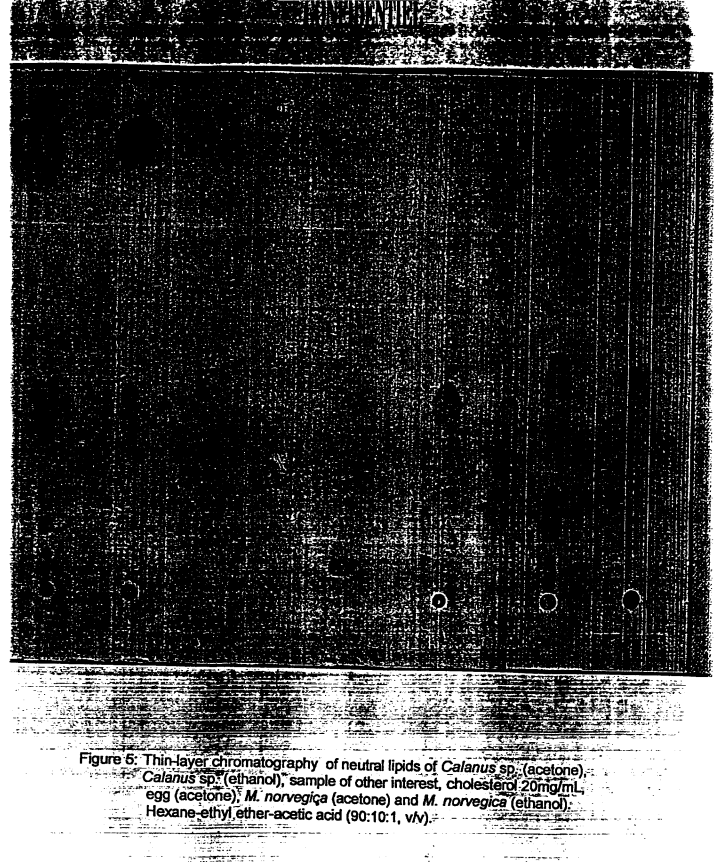


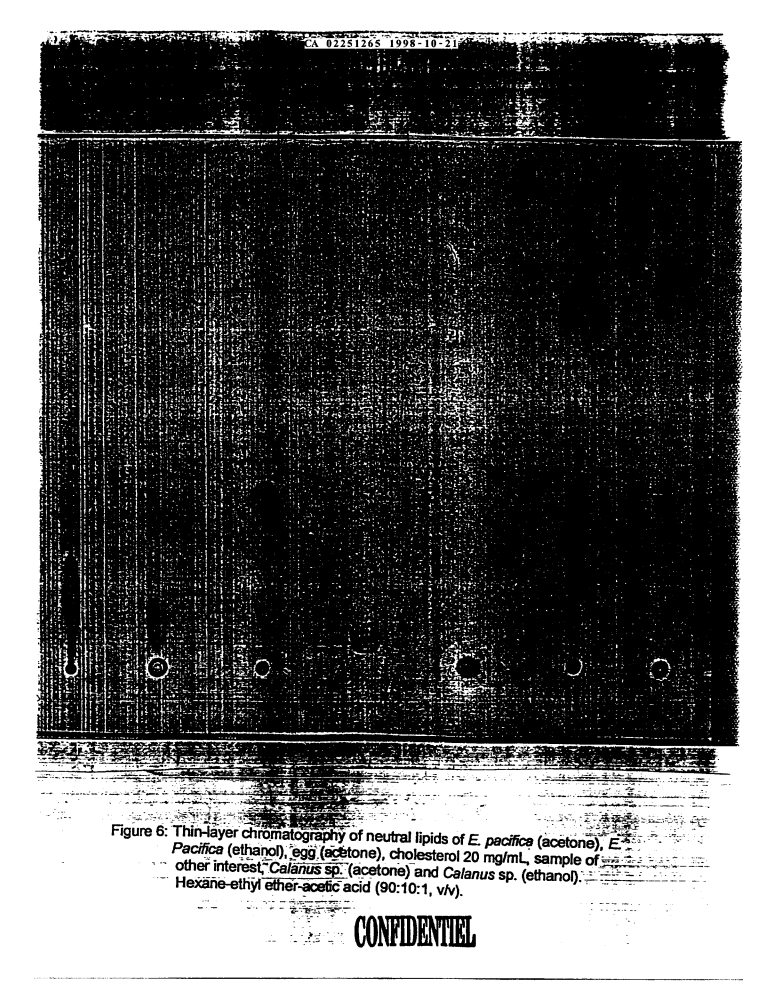
Figure 4: Gas-liquid chromatography of fatty acids from frozen krill (ethanol).

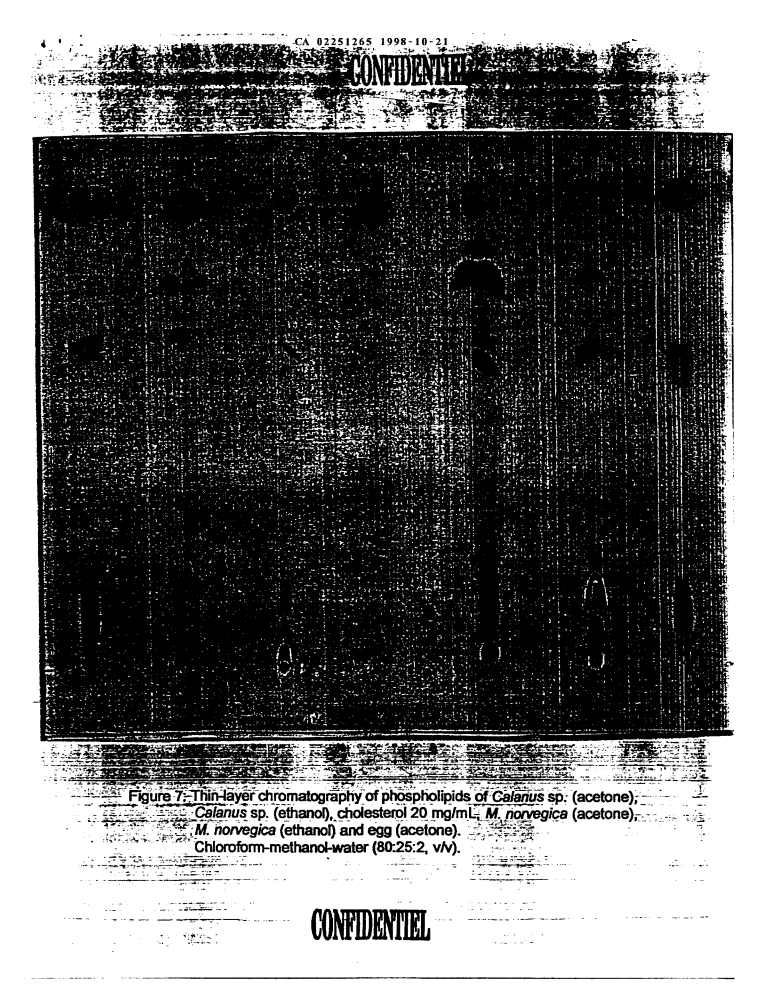
CONFIDENTIEL

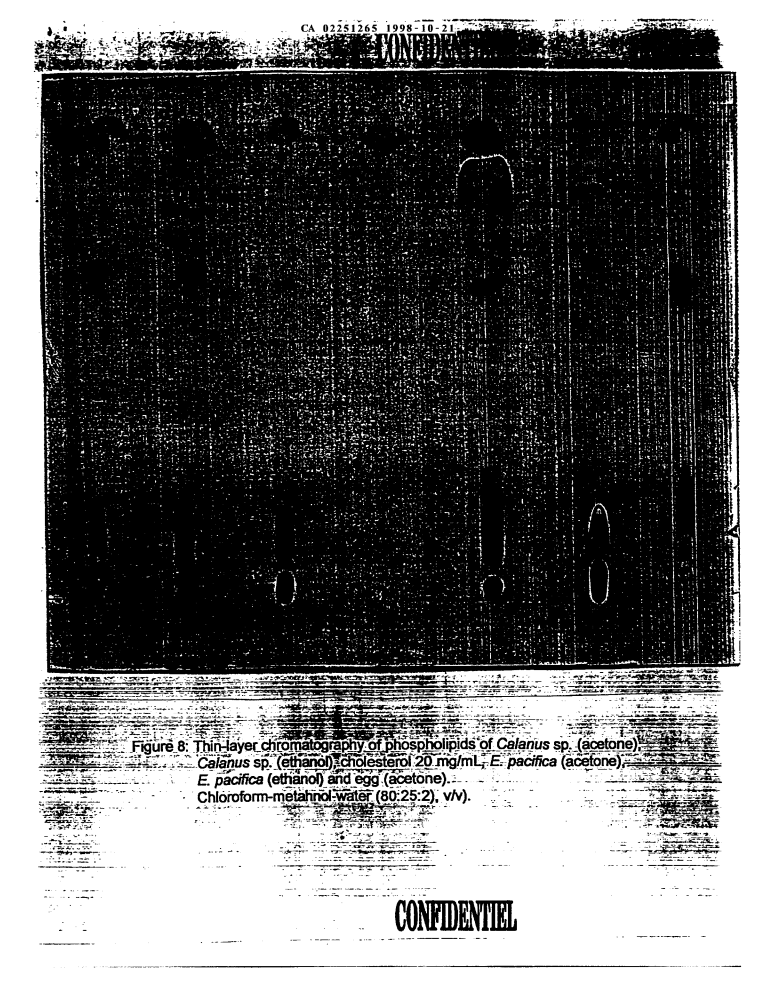




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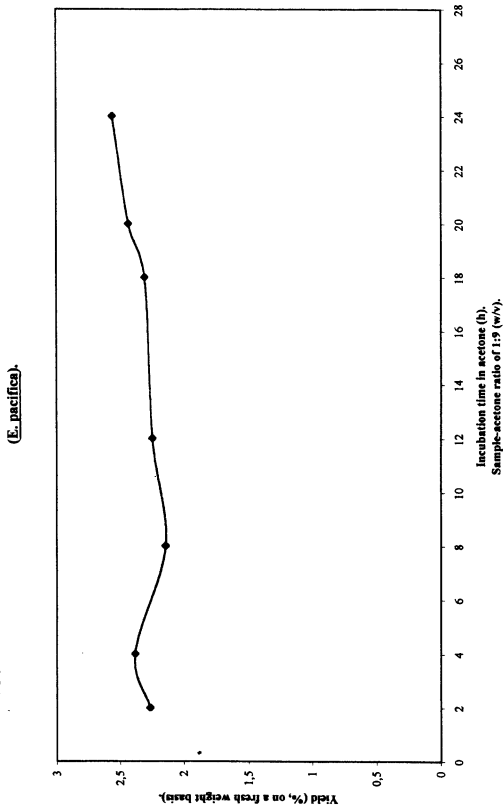


FIGURE 9. INFLUENCE OF INCUBATION TIME IN ACETONE ON LIPID EXTRACTION

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Determinations in triplicates (variation less than 5 %).

FIGURE 10. INFLUENCE OF THE VOLUME OF ACETONE ON LIPID EXTRACTION (E. pacifica)

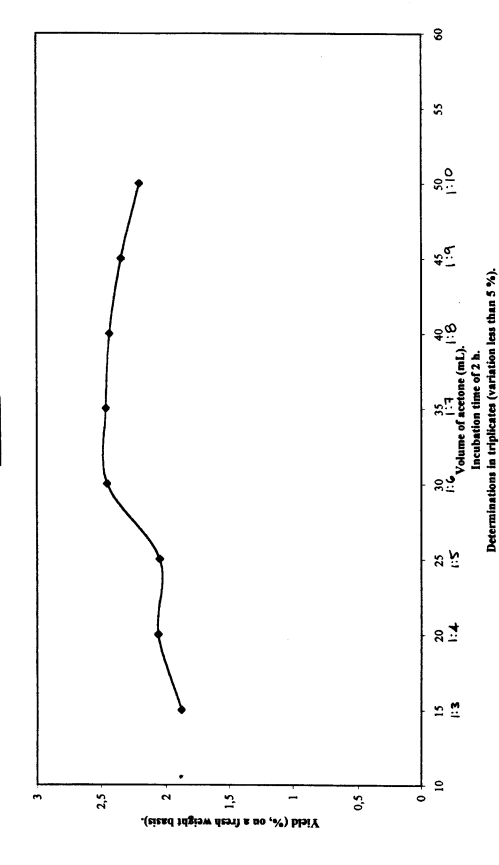
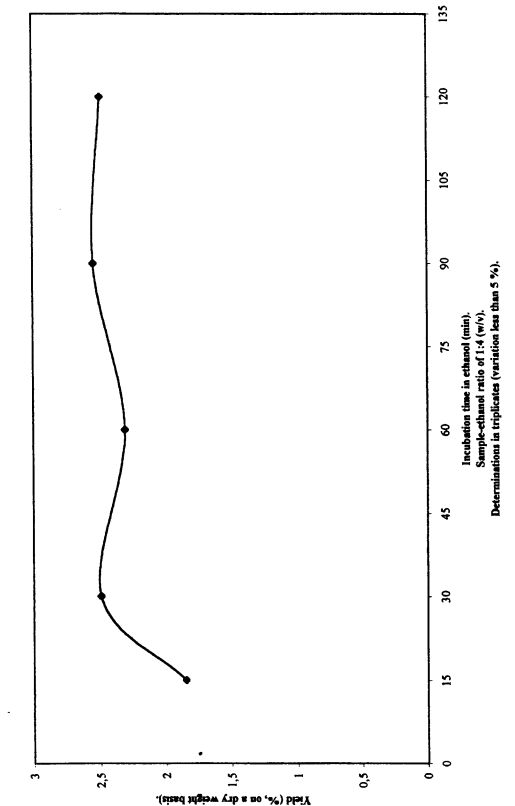


FIGURE 11. INFLUENCE OF INCUBATION TIME IN ETHANOL ON LIPID EXTRACTION (T. raschil).

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125 <u>8 ∓</u> Determinations in triplicates (variation less than 5 %). 52 <u>55</u> 50 |:2 Volume of ethanol (mL). ^{|..} ····· of 30 min. Incubation time of 30 min. 53 th 0 0 2 e 2 ø σ 00 5 Ś Yield (%, on a dry weight basis).

FIGURE 12. INFLUENCE OF THE VOLUME OF ETHANOL ON LIPID EXTRACTION (E. pacifica).

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①特許出願公開^{*} ⑫ 公 開 特 許 公 報 (A) 昭60-153779 @Int_Cl_4 識別記号 庁内整理番号 @公開 昭和60年(1985)8月13日 A 23 L A 61 K 8412-4B 6742-4C 1/42 9/48 7330-4C 31/355 ADL 審査請求 未請求 発明の数 1 (全2頁) 69発明の名称 栄養補助食品 ②特 顧 昭59-10625 ❷出 面 昭59(1984)1月24日 東京都中野区上鷺宮4丁目9番6号 @発 明 月 贷 一去 옆 治 明 岡 ⑦発 者 福 脩 多摩市永山4の4の21の304

東京都千代田区大手町1丁目2番3号

19日本国特許庁(JP)

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豊年製油株式会社

 発明の名称 栄養補助食品

⑦出

顧人

- 2. 特許請求の範囲
 - (1) ビタミンBおよび大豆レシチンをエイコサペンタ エン激含量の高い油に格解してなる混合液状物をゼ ラチンのカブセル内に封入した栄養補助食品。 (2) エイコサペンタエン酸含量の高い油が、イワシ油、
 - サパ油、イカ油、オキアミ油、ミンク鯨油等のごと き水産動物油である特許請求の範囲第(1)項記載の栄 **發補助食品。**

ふ 発明の詳細な説明

本発明はビタミンB、大豆レシチンおよびエイコサペ ンタエン酸含量の高い油を主成分とする新規な栄養補助 食品に関するものである。 近年、①豊かな食生活がもたらす栄養ペランスの偏り、

②階好優先の食生活なもたらす偏食、過剰摂取、③運動、 休息、栄養の健康保持パランスのくずれによる栄養損失、 ④高齢化社会に対応し得る補助栄養の必然性、等の要素 を背景として栄養補助食品の需要が急激に増加しており、 特に成人病は食生活の改善によって予防せんとする思想 が強いため栄養補助食品が好まれて食されている。

本発明はこのような食生活上のニーズから導かれたもの てもり、①細胞の老化を防ぐ、②コレステロール値を下 けて動脈硬化を防止する、③過酸化脂質の発生をおさえ て細胞の活性化を促す、④血管を浄化して脳卒中や心筋 標志を防止する等の効能を有するビタミンEと、①ビタ ミンEならびにエイヨサベンタエン酸の吸収を促進する、 (2)コレステロールを低下させて動脈硬化を防止する等の 効能を有する大豆レシチンと、抗血小板凝集作用に基づ く抗血栓、抗動脈硬化等の効能を有するエイコサペンタ エン酸含量の高い油を組合せた新規な栄養補助食品を提 供せんとするものである。 すなわち、本発明は、ビタミンEおよび大豆レシチンを エイコサベンタエン酸含量の高い油に溶解してたる混合 液状物をセラチンのカブセル内に封入した栄養指助食品 てある。 本発明において使用するビタミンBは、公知の製造法、 例えば、植物油の不ケン化物を分子蒸留あるいはクロマ

グラフイー等によって機縮する方法で得られたものが 適当であるが、その製造法は限定されるものではなく、 また、その起源も限定されない。 小麦胚芽油、サフラワー油、米油、コーン油等の液状植

物油中にはビタミンBが多く含まれているが、この含有 量はせいぜい Q3 劣以下であるためとれをそのまま使用 するととは好ましくない。

本発明におけるビタミンBの配合量は、カブセル内に封





特開昭60-153779(2)

入する液状物全体中に占める割合が少なくとも1%必要 であり、とれ以下では生体内での生理活性作用が劣り、 前配のごときビタミンBの効能が十分得られない。 また、ビタミンBと併用する大豆レシチンは、通常、大 豆曲の脱ガム工程で創生するガム質を脱水、乾燥して得 られる大豆油を含んだ大豆リン脂質(所開大豆レシチン)が通当であるが、ナセトン、アルコール等により精製 または繊縮されたレシチンを用いてもよく、また、ケフ アリン含量の少ないもしくはケファリン含量のない分別 レシチンを使用することもできる。

との大豆レシチンが、カブセル内に對入する液状物全体 中に占める割合は少なくとも1%必要であり、これ以下 では前配のごとを大豆レシチンの効能が十分得られない。 さらに、ビタミンEと大豆レシチンを浴解するエイコサ ペンタエン酸(C20:5)含量の高い油は、イワシ油、イ カ油、オキアミ油、ミンク酸油等のごときエイコサペン タエン酸を8%以上含有する水産動物油の積製油、ある いはこれらの油から分別法の手段によってエイコサペン タエン酸を酸縮して得られる油等が使用できる。 本発明におけるエイコサペンタエン酸含量の高い油の配

合量は、カブセル内に封入する液状物全体中に占める割 合が少なくとも10%必要であり、これ以下ではエイコ サペンタエン酸の生体内での生理活性作用が劣り、前記 のごときエイコサペンタエン酸の効能が十分得られない。 ビタミンBおよび大豆レシチンをエイコサペンタエン酸 含量の高い油に溶解してなる混合液状物は、必要に応じ、 とれら各成分の有効機度を維持できる範囲内において、 小麦胚芽油、サフラワー油、米油、コーン油、大豆油、 菜種油等の液状植物油で希釈することができる。 このようにして得られた混合液状物は、次いで、常法に

従ってゼラチンのカブセル内に封入する。

この封入方法の一例としては、組合液状物をゼラチン、 グリセリン、および水を溶験後射出成型したゼラチンカ プセルに所定量注入し、その後、注入口を加熱密封して 本発明の栄養補助食品を製造する。

セラチンカブセルの形状は球形、ラクビーボール形勢任 意である。

このようにして得られた本発明の栄養補助食品は、細胞 の老化を防ぎ、コレステロール値を下げ、過酸化脂質の 発生をおさえ、血管を浄化する等の作用を有するビタミ ンBと、ビタミンBおよびエイコサペンタエン酸の吸収 を促進し、コレステロールを低下させる等の作用を有す る大豆レシチンと、抗血小板要集による抗血栓および抗 動脈硬化等の作用を有するエイコサペンタエン酸を含有 するものであるから、これら各生理活性成分の相互作用 によって、血中コレステロールを下げ、高血圧を防ぎ、 細胞を若返らせて活性化するほか、心筋梗塞、脳梗塞の ごとき血栓性疾患等、循環器系成人病の治療および予防 に効果を有する等、健康食品としての機能を発揮し得る ものである。

次に本発明の実施例を示す。

実施例 1.

ビタミンE25重量部および大豆レシチン25重量 部を稍製イワシ油(エイコサペンタエン酸含量約16 %)50重量部に混合し、約60℃に加温、攪拌して 均一に溶解した。

一方、ゼラチン60重量部、グリセリン30重量部、 水10重量部を均一に混合し、フイルム状にした後、 容量約300mgのカブセル状に射出成型してゼラチン 容器を製造した。

この容器に前配の混合液状物を注入し、しかる後、注入口を加熱密封して本発明の栄養補助食品を得た。 実施例2.

ビタミンE30重量部および大豆レシチン30重量部 を、市販のエイコサペンタエン酸機能油(日本油脂KK 製、サンオメガ、エイコサペンタエン酸含量約25%))とサフラワー油を1:1の重量割合で混合したエイ コサペンタエン酸含量の高い油40重量部に混合し、 約60℃に加强、搅拌して均一に溶解した。 一方、ビラチン60重量部、グリセリン30重量部、水 10重量部を均一に混合し、フイルム状にした後、容 量約300mgのカブセル状に射出成型してゼラチン容器 を製造した。 との容器に前配の混合液状物を注入し、しかる後、注 入口を加熱密動して本発明の栄養補助食品を得た。

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A 61 K 9 / 48	6742	- 4C
31/355	ADL 7330	- 4C
	Request for	Review Unrequested Number of Claims 1 (Total 2 Pages)
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	(21) Application Number	Sho 59 – 10625
	(22) Application Date	January 24, 1984
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Specification

1. Title of Invention

Nutritional supplement

2. Scope of Claims

(1) A nutritional supplement being contained inside a gelatin capsule and comprising a nutritional liquid being formed by melting vitamin E, soybean lecithin in an oil having high eicosapentaenoic acid content.

(2) A nutritional supplement of claim 1 wherein the oil having high eicosapentaenoic acid content is a sea animal oil such as pilchard oil, mackerel oil, cuttlefish oil, krill oil, or mink oil.

3. Detailed Explanation of the Invention

The present invention relates to a new nutritional supplement having as its primary ingredients vitamin E, soybean lecithin, and an oil having high eicosapentaenoic acid content. In recent years, the demand for nutritional supplements has greatly increased due to factors such as (1) lack of nutritional balance brought about by a rich food culture, (2) pickiness and lack of food diversity caused by a selective food palette, (3) lack of nutrition brought about by a breakdown in balance of nutrition and health, exercise, and energy, and (4) the necessity for supplementary nutrition to correspond to an aging population. In particular, the desire to stem adult disease by the improvement of food life is strong, so people enjoy eating nutritional supplements.

The present invention arises out of the aforementioned nutritional needs and its object is to provide a new nutritional supplement from the combination of vitamin E, which (1) prevents the aging of cells, (2) lowers cholesterol and prevents the hardening of arteries, (3) prevents the occurrence of liquid fatty deposits and encourages cell life, (4) cleans and washes blood vessels and prevents brain or heart blood clots; soybean lecithin, which (1) encourages the absorption of vitamin E and eicosapentaenoic acid, (2) reduces cholesterol and prevents hardening of blood vessels and oil having high eicosapentaenoic acid content, which is effective anti-platelet aggregation, making it an anti-coagulant and acts to prevent the hardening of blood vessels.

In other words, the present invention is a nutritional supplement consisting of vitamin E and soybean lecithin melted in an oil having high eicosapentaenoic acid content in liquid form inserted into a gelatin capsule.

As for the vitamin E used in the present invention, items obtained by publicly known methods such as concentration by chromatography or molecular distillation of plant oils are appropriate, but the method of production thereof is not limited, and the source of energy is also not limited.

Vitamin E is included in great amounts in safflower oil, rice oil, corn oil, and other oil from water dwelling plants. However, the concentration

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of the oil is at most 0.3%, so using any of these as-is is not desirable.

The amount of vitamin E included in the present invention must be at least 1% of the liquid included in the capsule. If less than this concentration is achieved the activity of the vitamin in the body is reduced, and the benefits of vitamin E cannot be achieved.

Also, soybean fatty substance containing soybean oil obtained by dehydrating and drying gum produced by the production of soybean gum is generally acceptable for the soybean lecithin used in conjunction with vitamin E, but it is also acceptable to use lecithin that has been sugared and concentrated via acetone or alcohol, and it is also possible to use separated lecithin that contains little of no keratin.

This soybean lecithin must be present in the liquid that is inserted into the capsule in a concentration of at least 1%. As noted before, if the concentration falls below this level the effects of the substance will not be sufficient. In addition, for oil with a high concentration of eicosapentaenoic acid (C20 : 5) that has vitamin E and soybean lecithin dissolved within it, an oil with a eicosapentaenoic acid content of 8% or higher may be used, such as that found in sea animal oils such as pilchard oil, mackerel oil, cuttlefish oil, krill oil, or mink oil, or an oil that was derived from one of the above and had was concentrated with respect to eicosapentaenoic acid content.

The concentration of oil having a high concentration of eicosapentaenoic acid in the present invention must be at least 10% of the volume of liquid in the capsule. If this concentration is not achieved then the effect of the oil in the body will be lacking and as noted before the

Below, the embodiments of the present invention are explained. Embodiment 1.

25 parts vitamin E and 25 parts soybean lecithin are dissolved into 50 parts fish oil (containing 16% eicosapentaenoic acid), heated to around 60 degrees Celsius, and dried.

Then 60 parts gelatin, 30 parts glycerin, and 10 parts water are uniformly mixed, are made to take a film like form, are poured into a 300 milligram capsule and put into a gelatin container. The aforementioned compound is poured into this container, the entry point is heat sealed, and the nutritional supplement of the present invention is formed.

Embodiment 2

30 parts vitamin E by weight and 30 parts soybean lecithin are dissolved into a 1:1 mixture of safflower oil and market concentrated eicosapentaenoic acid oil (produced by Nippon Yushi Co. Ltd., Sun Omega, and containing 25% eicosapentaenoic acid), the mixture being 40 parts. The liquid is heated to around 60 degrees Celsius and dried.

Then 60 parts gelatin, 30 parts glycerin, and 10 parts water are uniformly mixed, are made to take a film like form, are poured into a 300 milligram capsule and put into a gelatin container. The aforementioned compound is poured into this container, the entry point is heat sealed, and the nutritional supplement of the present invention is formed. effects will not be sufficient. A nutritional supplement formed by melting vitamin E and soybean lecithin in oil with a high concentration of eicosapentaenoic acid, within the scope wherein it is possible to maintain the relative levels of the various substances, may be diluted with safflower oil, rice oil, corn oil, soybean oil, or seaweed oil or other sea life oil. Mixed oil obtained in this way is inserted into a gelatin capsule under normal circumstances.

As one example of the method of sealing the substance in a capsule, the mixed liquid will be sealed in a gelatin capsule in a certain amount, the gelatin formed by melting gelatin, glycerin, and water. After that, the entry hole will be heated and sealed to create the nutritional substance of the present invention.

The form of the gelatin capsules may be spherical or in the form of a rugby ball.

Because the nutritional supplement of the present invention obtained in this way contains vitamin E which prevents the aging of cells, lowers cholesterol numbers, prevents excess fat deposits, and has other positive side effects, as well as contains soybean lecithin, which acts to lower cholesterol and promote the absorption of eicosapentaenoic acid, and contains eicosapentaenoic acid, which acts to prevent the hardening of blood vessels, the combined effects of the various ingredients act to reduce the cholesterol levels in the blood, prevent high blood pressure, reduce fat, and promote heart and brain function in brain and heart medical patients, and acts to treat and prevent adult circulatory diseases, and as such can be said to function as a nutritional supplement.

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(54)【発明の名称】 痴呆症状改善薬

(57)【要約】

【目的】 痴呆症状を速やかに改善する、副作用のない 痴呆症状改善薬を提供する。

【構成】 ドコサヘキサエン酸を有効成分として含有す ることを特徴とする痴呆症状改善薬。

【効果】 痴呆症による意欲の低下、せん妄、対人関係の悪化、徘徊、落ちつきのなさ等の精神症状、および/ または痴呆症による計算能力の低下、判断力の低下、高 次機能の低下等の知的機能の低下を改善する。

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【特許請求の範囲】

【請求項1】 ドコサヘキサエン酸を有効成分として含 有することを特徴とする痴呆症状改善薬。

【請求項2】 痴呆症状が痴呆症による精神症状である、請求項1に記載の痴呆症状改善薬。

【請求項3】 精神症状が、痴呆症による意欲の低下、 せん妄、対人関係の悪化、落ちつきのなさ、および/ま たは徘徊である、請求項2に記載の痴呆症状改善薬。

【請求項4】 痴呆症状が痴呆症による知的機能の低下 である、請求項1に記載の痴呆症状改善薬。

【請求項5】 知的機能の低下が、痴呆症による計算能 力の低下、判断力の低下、および/または高次機能の低 下である、請求項1に記載の痴呆症状改善薬。

【発明の詳細な説明】

[0001]

【産業上の利用分野】本発明は痴呆症に伴う痴呆症状を 改善する薬剤に関し、更に詳しくは、ドコサヘキサエン 酸を有効成分として含有することを特徴とする痴呆症状 改善薬に関するものである。

[0002]

【従来技術】近年の人口の高齢化に伴い、痴呆症に対す る薬剤の開発は医学的にも社会的にもますます重要な課 題となっている。例えば、痴呆症患者の意欲の低下およ びせん妄、あるいは対人関係でのトラブル等により、家 族との関係が悪化して家庭介護が困難となることは最も 大きな問題として指摘されている。従来種々の薬剤が痴 呆症状改善薬として開発されてきたが、その効果は必ず しも満足しうるものではないばかりか、頭痛、めまい、 自発性低下、感情障害、胃腸障害等の副作用を伴うこと が多く、より優れた痴呆症状改善薬に対する期待は大き い。

【0003】ドコサヘキサエン酸は脳や網膜の興奮性膜 に多く含まれている不飽和脂肪酸で、アラキドン酸カス ケードを阻害する作用を有していることが知られてい る。またこのほかに、幾つかの有用な生理作用を有する ことが知られており、例えば、脳機能改善組成物、学習 能力增強剤、記憶力増強剤、痴呆予防剤、痴呆治療剤、 または脳機能改善効果を有する機能性食品(特開平2-49723号)、コリン作動性薬剤(特開平1-279 830号)、血栓症治療剤(特開昭57-35512) 号)等の特許出願がなされている。これらの中で、特開 平2-49723号はドコサヘキサエン酸による学習能 力や記憶力の増強及び血小板凝集の抑制作用を明らかに しているにすぎず、痴呆症状の改善については具体的開 示は全くない。また、特開平1-279830号はドコ サヘキサエン酸によりコリンエステラーゼ阻害剤である フイゾスチグミンの脳への送達量が増加することに関す るものである。

[0004]

【発明が解決しようとする課題】本発明は、痴呆症状を

速やかに改善する、副作用のない痴呆症状改善薬を提供 することにある。

【0005】

【課題を解決するための手段】本発明者らは、健康食品 として広く知られているドコサヘキサエン酸を痴呆症患 者に投与すると、その痴呆症状が速やかに改善されると いう新たな知見に基づき、本発明を完成するに至った。

【0006】すなわち、本発明は、ドコサヘキサエン酸 を有効成分として含有することを特徴とする痴呆症状改 善薬を提供する。

【0007】本発明の痴呆症状改善薬は、多発梗塞性痴 呆、脳血管性痴呆、脳機能障害による痴呆、ならびにア ルツハイマー型痴呆等の痴呆症に随伴する精神症状(例 えば、意欲の低下、せん妄、対人関係の悪化、落ちつき のなさ、徘徊等)あるいは知的機能の低下(例えば、計 算能力の低下、判断力の低下、高次機能の低下等)など に適用される。

【0008】本発明に用いるドコサへキサエン酸とは、 遊離酸をはじめ、その塩、エステル、グリセリド、リン 脂質、コリン化合物、アスコルビン酸化合物、アミノ酸 化合物等を意味するものである。このドコサヘキサエン 酸を含む油としては、好ましくは総脂肪酸中のドコサへ キサエン酸(遊離酸として)の占める割合が10%以上 のものが良く、このようなものの例を上げるとイワシ、 サバ、アジ、サケ、サンマなどの青背魚より抽出した魚 油、マグロやカツオなどの大型海産魚の眼窩脂肪由来の 魚油、微生物由来の油脂、オキアミ油、タラやイカ肝臓 より抽出した海産物由来の油脂などが好ましい例として 挙げられる。

【0009】本発明の痴呆症状改善薬は治療のために経 口的あるいは非経口的に投与することができる。経口投 与剤とては散剤、顆粒剤、カプセル剤、錠剤などの固形 製剤あるいはシロップ剤、エリキシル剤などの液状製剤 とすることができる。また、非経口投与剤として注射剤 とすることができる。これらの製剤は活性成分に薬理学 的、製剤学的に認容される製造助剤を加えることにより 常法に従って製造される。更に公知の技術により持続性 製剤とすることも可能である。当該製造助剤を用いる場 合は、本発明の痴呆症状改善薬中のドコサヘキサエン酸 (遊離酸として)の配合量は通常は10~100重量 %、好ましくは50~100重量%である。

【0010】上記製造助剤としては、内服用製剤(経口 剤)、注射用製材(注射剤)、粘膜投与剤(バッカル、 トローチ、坐剤等)、外用剤(軟膏、貼付剤等)などの 投与経路に応じた適当な製剤用成分から使用される。

【0011】例えば、経口剤および粘膜投与剤にあって は、賦形剤(例:澱粉、乳糖、結晶セルロース、乳糖カ ルシウム、メタケイ酸アルミン酸マグネシウム、無水ケ イ酸)、崩壊剤(例:カルボキシメチルセルロース、カ ルボキシメチルセルロースカルシウム)、滑沢剤(例: ステアリン酸マグネシム、タルク)、コーテング剤

(例:ヒドロキシエチルセルロース、白糖、ヒドロキシ プロピルセルロース、ポリビニルピロリドン)、矯味剤 などの製剤用成分が使用される。

【0012】顆粒剤を製造するには湿式又は乾式造粒

し、錠剤を製造するにはこれらの散剤及び顆粒剤をその ままあるいはステアリン酸マグネシウム、タルクなどの 滑沢剤を加えて打錠すればよい。これらの顆粒又は錠剤 はヒドロキシプロピルメチルセルロースフタレート、メ タアクリル酸、メタアクリル酸メチルコポリマーなどの 腸溶性基剤で被覆して腸溶性製剤、あるいはエチルセル ロース、カルナウバロウ、硬化油などで被覆して持続性 製剤とすることもできる。また、カプセル剤を製造する には散剤又は顆粒剤を硬カプセルに充填するか、活性成 分をそのままあるいはグリセリン、ポリエチレングリコ ール、ゴマ油、オリーブ油などに溶解したのちゼラチン 膜で被覆し軟カプセル剤とすることができる。

【0013】経口投与用の液状製剤を製造するには活性 成分と白糖、ソルビトール、グリセリンなどの甘味剤と を水に溶解して透明なシロップ剤、更に精油、エタノー ルなどを加えてエリキシル剤とするか、アラビアゴム、 トラガント、ポリソルベート80、カルボキシメチルセ ルロースナトリウムなどを加えて乳剤又は懸濁剤として もよい。これらの液状製剤には所望により矯味剤、着色 剤、保存剤などを加えてもよい。

【0014】また注射剤にあっては、水性注射剤を構成 し得る溶解剤ないし溶解補助剤(例:注射用蒸留水、生 理食塩水、プロピレングリコール)、懸濁化剤(例:ポ リソルベート80などの界面活性剤)、pH調整剤

(例: 有機酸またはその金属塩)、安定剤などの製剤用 成分が使用される。

【0015】注射剤を製造するには活性成分を必要に応 じ塩酸、水酸化ナトリウム、乳剤、乳酸ナトリウム、リ ン酸一水素ナトリウム、リン酸二水素ナトリウムなどの pH調整剤、塩化ナトリウム、ブドウ糖などの等張化剤 とともに注射用蒸留水に溶解し、無菌沪過してアンプル に充填するか、更にマンニトール、デキストリン、シク ロデキストリン、ゼラチンなどを加えて真空下凍結乾燥 し、用時溶解型の注射剤としてもよいし、活性成分にレ シチン、ポリソルベート80、ポリオキシエチレン硬化 ヒマシ油などを加えて水中で乳化せしめ注射用乳剤とす ることもできる。

【0016】さらに外用剤にあっては、水性ないし油性 の溶解剤ないし溶解補助剤(例:アルコール、脂肪酸エ ステル類)、粘着剤(例:カルボキシビニルボリマー、 多糖類)、乳化剤(例:界面活性剤)などの製剤用成分 が使用される。直腸投与剤を製造するには活性成分及び カカオ脂、脂肪酸のトリ、ジ及びモノグリセリド、ポリ エチレングリコールなどの坐剤用基剤とを加湿して溶融 し型に流しこんで冷却するか、活性成分をポリエチレン グリコール、大豆油などに溶解したのちゼラチン膜で被 覆すればよい。

【0017】その他、上記構成を有する本発明の痴呆症 状改善薬は、公知の製造法、例えば日本薬局方第10版 製剤総則記載の方法ないし適当な改良を加えた方法によ っても製造することができる。

【0018】とくに本発明の痴呆症状改善薬は、高純度 の(例えば90%以上)のドコサヘキサエン酸を軟カプ セル剤の形態で投与するのが、投与が簡便な点で好まし い。

【0019】本発明の痴呆症状改善薬の有効成分である ドコサヘキサエン酸の投与量は、患者の体重、症状等に より異なるが、一般には一日当たり、100~2000 mg/人程度であり、一日1回~数回に分けて投与す る。

【0020】以下、本発明を実施例により詳細に説明す る。

[0021]

【実施例】

試験例1. 精神症状の改善度評価試験

脳血管性痴呆患者13名、アルツハイマー型痴呆患者 (アルツハイマー病と老年痴呆)5名の対象者に対し て、従来の薬剤に加えてDHA70mg入りカプセルを 10錠~20錠投与(以下、DHA投与群と略)し、D HAの投与前と投与6ヶ月後の検査結果を比較検討し た。併せて、従来の薬剤治療を継続した群(以下、投与 不変群と略;24名)を対照群として同様の検査を施行 し、DHA投与群との変化を比較した。結果を表1に示 す。

[0022]

【表1】

表1. 精神症状の改善度

	改善	やや改善	不変	悪化
脳血管性痴呆	9	1	2	1
アルツハイマー型痴呆	0	5	0	0

【0023】なお、脳血管性痴呆における改善の内容

は、主にせん妄が改善された症例が2例、主に意欲が改

【0024】試験例2. 知的機能低下の改善度評価試 験 試験例1と同じ対象者に対して、計算力、判断力及び高 次機能の3項目を知的機能の簡易評価検査とし、コース 立方体組合せテストを動作性知能の簡易評価検査とし た。検査は、投与前と投与後6ヶ月後の計2回行い、結 果は統計学的に処理をした。結果を表2に示す。 【0025】 【表2】

	معليه بالبلار العبر بين مالة فالأله الروحية	
表2.	知的機能の改善度	

	DHA	没与群	投与不変群	
知的機能	投与前 (試験開始時)	67月後	試験開始時	6ヶ月後
計算力総点	6.2±3.3	6.9±3.0	3.4±2.9	3.1±3.3
判断力総点	4.6±3.0	5.5±3.3	4.1±2.9	2.6±2.4
高次機能	5.1±2.9	6.0±2.9	3.6±3.3	2.4±2.8

【0026】なお、コース立方体およびIQ(動作

性)試験においても、投与前後で改善が認められた。

フロントページの続き

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(54) [Title of the Invention] Medicine for Improvement of Dementia Symptoms

(57) [Abstract]

[Objective] To smoothly improve the symptoms of dementia and provide a medicine for said improvement without side effects.

[Structure] A medicine for improvement of dementia symptoms that has as a characteristic the inclusion of docosahexaenoic acid (DHA).

[Effect] The medicine improves the following ailments caused by dementia: loss of will, delirium, worsening of human relationships, loitering, manic psychological episodes and/or the reduction of powers of calculation, reduction of judgment, and reduction in the intellectual capacities and functioning of the higher functions.

000002

[Scope of Claims]

[Claim 1] A medicine for improvement of dementia symptoms being characterized by including as an active ingredient DHA. [Claim 2] A medicine for improvement of dementia symptoms of claim 1 that treats an adverse psychological state that is dementia.

[Claim 3] A medicine for improvement of dementia symptoms of claim 2 working to reduce will loss, delirium, worsening of human relationships, manic states, and/or loitering.

[Claim 4] A medicine for improvement of dementia symptoms of claim 1 working to reduce the loss of higher functions and of judgment brought about by dementia.

[Claim 5] A medicine for improvement of dementia symptoms of claim 1 working to reduce the loss of intellectual capacity, loss of facilities of calculation, loss of judgment, and/or loss of higher function due to dementia.

[Detailed Description of the Invention] [0001]

[Industrial Field of Use] The present invention is in relation to a medicine for the betterment of mental symptoms that accompany dementia, and in particular relates to a medicine for the improvement of dementia symptoms that includes as an active ingredient DHA.

[0002]

[Related Art] With the aging of society in recent years, the development of medicine for the treatment of dementia has become more important both medically and socially. For example, a dementia patient may suffer worsening family relationships as a result of loss of will, delirium, or trouble in interpersonal relationships, and the looking after of the patient within the family becomes difficult. This has been pointed out as the most serious cause for concern. In past years many medicines have been developed for dementia, but the results haven't always been satisfactory. Furthermore, the traditional medicine can cause headache, dizziness, reduction in sex drive, emotional disturbances, and other side effects such as damage to the stomach. It is with this that there has been great expectation for the development of a new medicine for dementia.

[0003] DHA is present in abundance in the brain and the thick mucus membranes. DHA is known to stop the functioning of arachidonic acid. Also, in addition to this, it is known that DHA contains several useful biological functions. For example, the following patent applications have been made: substance for the increase of brain function, medicine for the improvement of academic performance, medicine for improvement of memory, dementia prevention substance, substance for the treatment of dementia, and functional food that improves brain function (Hei 2-49723), cholinergic agent (Hei 1-279830), agent for the treatment of thrombosis (Sho 57 - 35512), among others. Among these, patent application Hei 2 - 49723 shows that DHA can aid in the improvement of academic ability as well as increasing memory performance, and also acts to prevent the formation of platelet aggregation. However, this application said nothing more and did not hint at the specific application of DHA to dementia. Also, Application Hei 1 - 279830 is in relation to the increase of transmission volume to the brain of physostigmine, a cholinesterase antagonist, via the DHA. [0004]

[Problem Solved by the Invention] The present invention provides a medicine to improve with the symptoms of dementia without providing side effects. [0005]

[Method of Solving the Problem] The inventors of the present invention gave DHA, widely known for being a health food, to dementia patients, whereupon the symptoms of the dementia were immediately lessened, and based on that discovery gathered to file this application.

[0006] In other words, the present invention provides a medicine for the improvement of dementia symptoms that includes DHA.

[0007] The medicine for the improvement of dementia symptoms of the present invention is applied to psychological states accompanying dementia from multiple infarction, brain blood vessel function, brain damage, or Alzheimer's disease (such as loss of will, delirium, worsening of human relationships, mania, loitering, etc.) or the reduction in intellectual capabilities (for example a reduction in the powers of calculation, a reduction in judgment, or a reduction in higher order functions).

[0008] The DHA used in the present invention is an isolated acid, and refers to salt, ester, glyceride, phospholipids, choline compounds, ascorbic acid compounds, amino acid compounds. As for the oil that includes the DHA, an inclusion ratio of 10% or more DHA (as an isolated acid) within general fatty acids. As an example of such an oil, the fish oil extracted from blue backed fish such as Japanese pilchard, mackerel, horse mackerel, salmon, and Pacific saury, the fish oil from large ocean fish eye oil, such as that of the tuna or the shipjack tuna, oil coming from microorganisms, krill oil, and oil from industrial products extracted from the livers of Pacific cod and dolphins.

[0009] The medicine for the improvement of the symptoms of dementia of the present invention may be administered either orally or non-orally. For oral administration, powder, granule, capsule, lozenge, and other solid forms of administration are acceptable. Alternatively, the medicine may be administered as syrup, elixir, and other liquid forms. Also, for non-oral administration an injection can be given. By adding these forms of manufacturing to the approved medicine that is the active portion of the drug the medicine may be manufactured in the normal fashion. Furthermore, it is also possible to turn the medicine into extended release tablets via publicly known methods. When using those manufacturing helper substances the DHA levels within the medicine for the improvement of the symptoms of dementia of the present invention is between 10 and 100 % by weight, and preferably between 50 and 100 % by weight.

[0010] An appropriate manufacturing helper substance will be used in the above in accordance with the administration method, for example, internal use substances (oral medicine), injection use substances (injected medicine), adhesive administration substances (buccal, troche, and suppositories).

[0011] For example, in oral and adhesive administration excipients (example: starch, milk sugar, crystal cellulose, milk calcium, metakei acid aluminum acid magnesium, waterless silicic acid), collapse agents (example: carboxymethylcellulose, carboxymethyl cellulose calcium), lubricants (example: sterin acid magnesium, talc), coatings (example: hydroxyl methyl

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cellulose, sugar, hydroxyl propyl cellulose), and taste making agents, and other production substances may be used. [0012] In order to manufacture granules, wet or dry droplets are formed, and in order to produce pills, it is permissible to form the tablets with the powder and granules either left as they are or with additional stearic acid magnesium, talc, or other lubricant. These granules or tablets are coated with a stomach settling agent such as hydroxypropyl - methyl cellulose phthalate or methacrylic acid or methacrylic acid methyl copolymer, among others, and coating is made using stomach setting agent or ethyl cellulose, carnauba wax, hardened oil, or other substance. By doing so a durable pharmaceutical product may be produced. Also, in order to produce the medicine in capsule form, the powder or granules are filled into a hard capsule or the active ingredients are coated with a gelatin film either as is or after being melted into gelatin, polyethelyn glycol, sesame oil, olive oil, or other oil. In this way it is possible to generate a soft capsule.

[0013] In order to produce liquid medicine for oral administration, the active ingredient and a sweetener such as refined sugar, sorbitol, glycerol are dissolved in water, a clear syrup, essential oil, and ethanol are added making an elixir-like medicine, or alternatively gum arabic, tragacanth gum, polysorbate 80, carboxymethyl cellulose (CMC), or another such substance is added and an emulsion or a suspension is produced. This is also acceptable. Flavor agents, color changing agents, and/or preservatives may be added to the liquid solutions discussed herein, according to taste.

[0014] Also, stable production medicine components are used for injectable medicine, such as solution from water soluble injectable medicine and melted helper substances (example injection use distilled water, biological salt water, or propylene glycol), suspension substances (example: polysorbate 80 or other surfactant), pH regulation substances (example: organic acid or its metal salt).

[0015] In order to produce injection-use medication, the active ingredients are mixed with salts, sodium hydroxide, emulsion, emulsion natrium, dibasic sodium phosphate, sodium dihydrogen-phosphate, and other pH adjusting agents, sodium chloride, grape sugars, and other tonicity adjusting agents in injection use distilled water. The solution is sterilized and poured into an ampoule. Alternatively, mannitol, dextrin, cyclodextrin, gelatin, and other substances are added, fired into crystals under vacuum conditions, and placed into a form to be melted at the time of injection. To the active ingredients are added lecithin, polysolvent 80, polyoxyethylene hydrogenated castor oil, and other substances, melted into water and made into an injectable solution. [0016] Additionally, water or oil soluble medicines or soluble helper substances (example: alcohol, fatty acid esters), adhesives (example: carboxy vinyl polymer multi-sugars), emulsifiers (example: surfactants), and other substances are used as ingredients in externally administrable medicine. In producing rectally administered medicine the active ingredients and cocoa butter, fatty acid salts, monoglycerides and other suppository use substances are humidified, melted, poured into a mold, hardened, and frozen. Alternatively, the active ingredient could be melted in polyethylene glycol, soybean oil, or other oil, and thereafter coated in a gelatin film. [0017] Additionally, the medicine for the improvement of dementia symptoms of the present invention with the above listed characteristics may be produced using publicly known manufacturing methods, for example as stipulated in version 10 of the Pharmacy Act of Japan, noted in the manufacturing addendum, or a method that has appropriately modified the aforementioned method.

[0018] In particular, the medicine for the improvement of dementia of the present invention administration of a high purity concentration of DHA (for example, 90% or above) via a soft capsule is desirable because of the ease of administration. [0019] The amount of DHA administered in the medicine to prevent the symptoms of dementia of the present invention will vary based on the body weight and health conditions of the patient, but in general, the dose will range from 100 to 2000 mg / person with between one and several administrations per day. [0020] Below we explain the present invention in detail by following an embodiment of the present invention. [0021]

[Embodiments]

Embodiment 1. Test to measure level of psychological improvement

The targets of this test were 13 cranial blood vessel related dementia patients and 5 Alzheimer's related dementia patients. In addition to traditional treatments, 10 - 20 capsules including 70 mg of DHA each were administered (hereinafter referred to as the "DHA Administration Group"), and the results of the test were compared before administration and 6 months after administration. Also, a group that continued traditional pharmaceutical treatments (hereinafter referred to as the "Unchanging Administration Group"; 24 individuals) were targeted for the same test and the variance from the DHA Administration Group was observed. The results appear in Table 1.

[0022]

[Table 1]

solution.				
Table 1. Level of Improvement in Psychological State				
	Recovered	Somewhat	No Change	Worsened
		Recovered		
Cranial Blood Vessel	9	1	2	1
Dementia				
Alzheimer's Dementia	0	5	0	0

[0023] To further break down the content of the "improvements" seen in the cranial blood vessel related dementia patients, 2 cases of improvement in delirium were seen, 3 cases of greatly improved ambition were observed, 3 cases of improved loitering were observed. Also, among the Alzheimer's dementia patients we observed 1 case each of improved ambition, human relationships, and manic states, respectively, for a total of 3 observed improvements. The Unchanging Administration Group did not show any change in symptoms in this same period, and all cases were evaluated to have no change.

[0024] Embodiment 2. Test to measure improvement of loss of intellectual capacity.

The calculation skills, judgment, and higher functions of the same test group as test 1 were evaluated. This test was a simple evaluation of intellectual abilities. Also, a course correction and pathfinding test was administered as a simple measure of motor control. The test was administered twice, once before administration of DHA and once 6 months after the administration of DHA. The results were statistically aggregated. The results are shown in Table 2. [0025]

[Table 2]

1000	Table 2. Level of improvement of interfectual Abilities					
	DHA Admin G	roup	Unchanged Group			
Intellectual Ability	Preadmin (at 6 months pos		Pre-admin	6 months		
	test start time)	– admin		post - admin		
Calculation Abilities	6.2 +- 3.3	6.9 +- 3.0	3.4 +- 2.4	3.1 +- 3.3		
Total						
Judgment Total	4.6 +- 3.0	5.5 +- 3.3	4.1 +- 2.9	2.6 +- 2.4		
Higher Function Total	5.1 +- 2.9	6.0 +- 2.9	3.6 +- 3.3	2.4 +- 2.8		

Table 2. Level of Improvement of Intellectual Abilities

[0026] Note that in both the motor and IQ tests,

improvements were seen after the administration of DHA.

Continued from the front page

(72) Inventor Miyakawa Fumio Kanagawa-ken, Sumohara-shi, Nandai 1 – 2 – 12 (72) Inventor Muramatsu Ei Tokyo-to, Hino-shi, Nishi Hirayama 5 – 27 – 10

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Bruheim et al.Group No.:1651Serial No.:12/057,775Examiner:D. K. WareFiled:28 March 2008BIOEFFECTIVE KRILL OIL COMPOSITIONSExaminer:

INFORMATION DISCLOSURE STATEMENT LETTER

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Sir or Madam:

The citations listed in the attached IDS Form PTO-SB08 may be material to the examination of the above-identified application, and are therefore submitted in compliance with the duty of disclosure defined in 37 C.F.R. §§ 1.56 and 1.97. The Examiner is requested to make these citations of official record in this application.

Applicants wish to bring to the Examiner's attention that we are not providing copies of US Patents or published US patent applications as instructed under 37 CFR 1.98(a)(2).

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The Commissioner is hereby authorized to charge any required fees or credit any overpayments to Attorney Deposit Account No.: 50-4302, referencing Attorney Docket No.: AKBM-14409/US-5/ORD.

Respectfully submitted,

Dated: 14 January 2014

/J. Mitchell Jones/ J. Mitchell Jones Registration No. 44,174 Casimir Jones S.C. 2275 Deming Way Suite 310 Middleton, WI 53562 Phone: (608) 662-1277 Fax: (608) 662-1276

Electronic A	Electronic Acknowledgement Receipt					
EFS ID:	17912253					
Application Number:	12057775					
International Application Number:						
Confirmation Number:	1945					
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS					
First Named Inventor/Applicant Name:	Inge Bruheim					
Customer Number:	72960					
Filer:	John Mitchell Jones/Thomas Vita					
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Application Number		12057775	
Filing Date		2008-03-28	
First Named Inventor Inge		Bruheim	
Art Unit		1651	
Examiner Name Ware			
Attorney Docket Number		NATNUT-14409/US-5/ORD	

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INFORMATION DISCLOSURE	Application Number		12057775	
	Filing Date		2008-03-28	
	First Named Inventor Inge		Bruheim	
(Not for submission under 37 CFR 1.99)	Art Unit		1651	
	Examiner Name	Ware		
	Attorney Docket Numb	er	NATNUT-14409/US-5/ORD	

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INFORMATION DISCLOSURE	Application Number		12057775	
	Filing Date		2008-03-28	
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STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit		1651	
	Examiner Name	Ware		
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Name/Print	J. Mitchell Jones	Registration Number	44174

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(54) Title: IMPROVED ABSORPTION OF FAT-SOLUBLE NUTRIENTS

(57) Abstract: Carotenoids are provided as dietary supplements to animal feed. These supplements improve the bioavailability of carotenoids by providing them in combination with phospholipids. The invention provides animal feeds for aquatic and terrestrial animals, and methods for making the feeds.

TITLE OF THE INVENTION

[0001] Improved Absorption of Fat-Soluble Nutrients

BACKGROUND OF THE INVENTION

[0002] This application is related to improving the bioavailability of carotenoids as provided in formulated mixtures to animals. The invention provides both a specific composition and a method of manufacture for improved delivery of carotenoids.
[0003] This invention relates to a carotenoid composition and methods for its manufacture and use. In one aspect, the invention relates to carotenoids, synthetic or naturally produced by a single-celled organism, and phospholipids containing highly unsaturated fatty acids. In another aspect, the invention relates to methods of increasing carotenoid stability during feed processing and improving bioavailability in the gastrointestinal (GI) tract of coldwater species. In yet another aspect, the invention relates to using products made from these carotenoid compositions as a dietary supplement in various animal feeds.

[0004] The carotenoids, as a class of compounds, are classified into two main groups: carotenes and xanthophylls. In contrast to carotenes, which are pure polyene hydrocarbons, such as beta-carotene or lycopene, xanthophylls contain oxygen functional groups, such as hydroxyls, epoxy and/or oxo groups. Typical representatives of the xanthophyll group are astaxanthin, canthaxanthin and zeaxanthin.

[0005] A distinct red color is of prime importance to customer acceptance of a subset of food products, particularly aquatic food animals such as salmon, trout, shrimp, lobster and many other marine animals (Hinostroza, Huberman et al. 1997; Bjerkeng and Berge 2000). The oxygenated carotenoids (xanthophylls) are responsible for the red color of these aquatic animals. These xanthophylls are also useful for adding pigmentation to the flesh and products of other animals, and to other foodstuffs, for example poultry and eggs, various dairy products, snack foods, and the like.

[0006] Astaxanthin is the most abundant carotenoid present in the aquatic world (Shahidi, Metusalach et al. 1998). Aquatic animals, like terrestrial animals, generally cannot synthesize astaxanthin or any other carotenoid, although many of these animals

accumulate carotenoid compounds that are present in their diets. Some of these animals, such as crustaceans, can interconvert some carotenes to xanthophylls, of which astaxanthin is the predominant compound formed. However, aquatic fish accumulate dietary astaxanthin even though these fish cannot convert any other carotenoid compound to astaxanthin. Therefore, the astaxanthin present in aquatic fish, and in products produced from these fish, must be derived directly from dietary sources.

[0007] Currently, synthetic astaxanthin is added to feeds of aquacultured salmonids to provide a source of this carotenoid (Bell, McEvoy et al. 1998). In some cases, synthetic canthaxanthin (another xanthophyll that is very closely related to astaxanthin) is used in place of astaxanthin in feeds for salmonids, but this compound does not function as well in these fishes as the naturally predominant astaxanthin (Bell, McEvoy et al. 1998).

[0008] Natural sources of dietary astaxanthin, including krill, crawfish, crustacean processing by-products, bacteria, yeast, algae, and higher plants are in great demand by aquacultural industries. However, these natural sources tend to be too expensive and of limited availability and reliability to be commercially viable. Lycopene is an alternative natural carotenoid that might meet the cost criterion for inclusion in feeds (Clark, Yao et al. 2000). It is in a class of carotenoids that characteristically gives color to many vegetables.

[0009] Carotenoids are easily isomerized by heat, acid or light. Once isomerized, they lose their biological antioxidant properties (Fennema 1996). The high demands placed on xanthophyll-containing formulations with respect to coloring action and bioavailability can thus not always be met because of these problems (Yeum and Russell 2002). Indeed, various processes and a number of combined emulsifying/spray-drying processes (see patents DE-A-12 11 911 or in EP-A-0 410 236) have been proposed to improve the color yields and to increase the absorbability or bioavailability carotenoids.

[0010] One specific problem which has not yet been addressed is related to the low body temperature of salmonid fishes, which is equal to the temperature of the water in which they inhabit, generally 0 to 14°C. Natural astaxanthin, especially those in *Phaffia* yeasts, are concentrated in oil droplets that contain about 13% palmitic acid (16:0) with a melting point of 64°C, and about 32% oleic acid (18:1n9) with a melting point of 16°C (Deuel 1951). Because of these high melting point fatty acids, the astaxanthin containing oil droplets solidify near 10°C. This makes it difficult for the fish to incorporate the

astaxanthin from the solidified oil droplet at water temperatures below 10°C. This is especially problematic for coldwater fish.

BRIEF SUMMARY OF THE INVENTION

[0011] The invention alleviates these problems by providing a process for preparing a mixture of carotenoids and phospholipids rich in highly unsaturated fatty acids (PUFA). The process comprises the following steps:

[0012] a) Preparing a molecularly-associated composition of carotenoids and a phospholipid with an edible oil or a mixture of water and a water-miscible organic solvent. If appropriate, a water-dispersible dry powder could also be prepared. To achieve dispersion, e.g., in the form of a suspension or an emulsion, it is advantageous to use an edible oil (such as, but not limited to, sesame oil, corn oil, cottonseed oil, soybean oil, or peanut oil) plus esters of medium chain-lengths vegetable fatty acids or fish oils (such as, but not limited to, mackerel, capelin, menhaden or cod liver oil).

[0013] b) Further increasing the stability of the carotenoids to oxidative decay by adding stabilizers such as, but not limited to, alpha-tocopherol, *t*-butylated hydroxytoluene, *t*-butylated hydroxyanisole, ascorbic acid or ethoxyquin.

[0014] c) Providing the carotenoids used to produce the composition from natural sources and/or synthetic sources.

[0015] d) The phospholipids used to produce the composition are rich in polyunsaturated fatty acids (PUFA) having two or more double bonds in at least 20% of total fatty acids.

[0016] e) The carotenoid composition according to the invention can also contain at least one other active substance in concentrations of 0.01 to 40% by weight.

[0017] Possible examples of these active substances are the following:

[0018] Other carotenoids such as for example bixin, zeaxanthin, cryptoxanthin, citranaxanthin, canthaxanthin, astaxanthin, beta-apo-4-carotenal, beta-apo-8-carotenal, beta-apo-8-carotenoic esters, lycopene, or lutein, singly or as a mixture.

[0019] Vitamins, such as vitamin A, vitamin A acetate, vitamin A palmitate, riboflavin, vitamin B₁₂, ascorbic acid, ascorbyl palmitate, nicotinic acid, nicotinamide, pyridoxine hydrochloride, vitamin D₃, tocopherol, tocopherol acetate, tocopherol palmitate, tocotrienol, vitamin K, thiamine, calcium pantothenate, biotin, lipoic acid, folic acid, and folic acid

derivatives (such as tetraBASF hydrofolic acid, 5-methyltetrahydrofolic acid, 10formyltetrahydrofolic acid) and 5-formyltetrahydrofolic acid).

[0020] Compounds with vitamin or coenzyme characteristics, such as choline chloride, carnitine, taurine, creatine, ubiquinones, S-methylmethionine, and S-adenosylmethionine.

[0021] Polyunsaturated fatty acids, such as linoleic acid, linolenic acid, arachidonic acid (ARA), eicosapentaenoic acid (EPA), and docosahexaenoic acid (DHA) and esters thereof including but not limited to triglycerides.

[0022] Glutathione and its esters such as, for example GSH monomethyl ester, GSH dimethyl ester, GSH monoethyl ester, and GSH diethyl ester.

[0023] Depending on the nature of the formulation, it may contain, besides the carotenoids, at least one other additive such as, for example, oils, protective colloids, alkaloids (such as peperine (Badmaev, Majeed et al. 1999)), and antioxidants.

[0024] Examples of protective colloids that can be used are gelatin, fish gelatin, starch, dextrin, plant proteins, pectin, gum arabic, casein, caseinate, or mixtures thereof. It is also possible to employ polyvinyl alcohol, polyvinylpyrrolidone, methylcellulose, carboxymethylcellulose, hydroxypropylcellulose, and alginates.

[0025] To increase the mechanical stability of the dry powder, it is also possible to add to the colloid a plasticizer such as sugars or sugar alcohols, such as sucrose, glucose, lactose, invert sugar, sorbitol, mannitol, or glycerol.

[0026] The use of the PUFA-rich phospholipids as part of this formulation also provides additional benefit to the survival and health of the animal consuming the invention's formulation (Bracco and Decekbaum 1992; Furuita, Takeuchi et al. 1998; Place and Harel 2002).

[0027] The present invention provides a mixture comprising a carotenoid and PUFArich phospholipid.

[0028] The present invention provides a composition comprising a mixture including a carotenoid either in synthetic or natural form and a phospholipid having at least 20% PUFA, where the phospholipid is in an amount sufficient to improve carotenoid stability and bioavailability and prevent solidification when the composition is fed to coldwater species, and the carotenoid is in an amount sufficient to produce acceptable coloring in edible tissues.

[0029] The present invention also provides a molecularly-associated complex comprising a carotenoids and a phospholipid.

[0030] The present invention provides a composition comprising a molecularlyassociated complex including an amount of a carotenoid and an amount of a phospholipid, wherein the amount of the phospholipid is sufficient to improve carotenoids stability and bioavailability and prevent solidification when the composition is fed to coldwater species and the amount of the carotenoid is sufficient to produce acceptable coloring of edible tissues.

[0031] The present invention also provides a mixture comprising a carotenoid, a phospholipid, and a bioactive compound, or a bioactive complex (comprising a carotenoid/phospholipid/bioactive compound), and/or mixtures or combinations thereof.

[0032] The present invention provides a composition comprising a mixture including a carotenoid, a phospholipid and a bioactive compound, a bioactive complex, or mixtures or combinations thereof, wherein the phospholipid is present in an amount sufficient to improve the carotenoids' stability and bioavailability and prevent solidification when the composition is fed to coldwater species, and wherein the amount of the total carotenoid is sufficient to produce acceptable coloring of edible tissues.

[0033] The present invention provides a composition comprising a cellular material and a phospholipids wherein the phospholipid to cellular material is in the ratio of from about 1:1 to about 1:100 and the cellular material comprises long chain polyunsaturated fatty acids and/or carotenoids.

[0034] The present invention also provides a method for making a carotenoidcontaining composition with increased carotenoid stability and bioavailability with low melting temperature when fed to cold-water species, including the step of mixing carotenoids and a PUFA-rich phospholipid. The method can further include the step of mixing the carotenoid/phospholipid composition with another bioactive compound forming an alternative and useful composition.

[0035] The present invention also provides a method for making a carotenoidcontaining composition with increased stability and bioavailability including the step of contacting a carotenoid and a phospholipid under conditions sufficient to maintain the carotenoid and the phospholipid in a molecularly-associated form. The method can further

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include the step of admixing the carotenoid/phospholipid molecular association with a bioactive compound.

[0036] The present invention also provides for making a long chain polyunsaturated fatty acid (LC-PUFA) composition with increased stability and bioavailability including the step of contacting a cellular material containing said LC-PUFA and a phospholipid under conditions sufficient to maintain the LC-PUFA and the phospholipid in a molecular association form. The method can further include the step of admixing the LC-PUFA/phospholipid molecular association with a bioactive compound.

[0037] The present invention also provides a method for enhancing the pigmentation of coldwater animals by providing such animals with a feed enriched with a composition that consists of a cellular source of carotenoid such as, but not limited to *Phaffia* yeast, *Haematococcus* algae, marigold flowers, mixed with a PUFA-enriched phospholipid such as, but not limited to, plant lecithins, egg yolk lecithin, phospholipid-rich extracts from animals or animal byproducts, and phospholipid-rich extracts from microbial sources. The cellular or synthetic carotenoid material and phospholipid material are premixed and homogenized prior to the addition to a feed in order to stabilize and solubilize the carotenoid and such a process surprisingly results in the enhanced bioavailability of the carotenoids by the coldwater animal.

BRIEF SUMMARY OF THE SEVERAL VIEWS OF THE DRAWINGS

[0038] Figure 1. Improved total carotenoid content of rainbow trout using conditions as described in Example 5 (for the Astaxanthin compared to Astaxanthin + DHA-phospholipid) and Example 4 for Astaxanthin compared to Astaxanthin + soy lecithin. The control had no added astaxanthin in the diet (some residual carotenoids were in the original diet). The soy lecithin gave a 34% higher incorporation of astaxanthin (AX) than AX alone. The DHA-rich phospholipid gave 56% higher incorporation of AX than AX alone.

DETAILED DESCRIPTION OF THE INVENTION

[0039] Definitions

[0040] Unless otherwise stated, the following terms shall have the following meanings:
[0041] The term "solution" means a liquid and any mixture of a liquid and a solid that has fluid attributes, e.g., flowable or having appreciable fluidity at standard temperature and

pressure, including, without limitation, a dispersion of a solid(s) in a liquid, an emulsion, a slurry, a micro-emulsion, colloidal suspension, a suspension, or the like.

[0042] An "emulsion" is suspension of one liquid in another with which the first will not mix. The first liquid can be suspended as small globules in the second liquid. An oil or an aqueous form of the compositions of this invention can be emulsified into an aqueous solution.

[0043] An "active substance" is any material that functions or is capable of functioning in a manner characteristic of that substance.

[0044] The term "molecular association" or "molecularly-associated" means a combination of two or more molecular species associated via any known stabilizing atomic or molecular level interaction or any combination thereof, where the interactions include, without limitation, bonding interactions such as covalent bonding, ionic bonding, hydrogen bonding, coordinate bonding, or any other molecular bonding interaction, electrostatic interactions, a polar or hydrophobic interactions, or any other classical or quantum mechanical stabilizing atomic or molecular interaction.

[0045] The term "species" is defined as any species in the animal kingdom, including mammals, fish, crustaceans and mollusks.

[0046] An "aquatic animal" is an animal that lives primarily in an aquatic environment, and includes fish, crustaceans, and mollusks. Aquaculture methods and/or commercial production practices have been developed to cultivate aquatic animals.

[0047] A "fish" and the plural "fish" are defined in this invention as any Ostiechthyean or Chondrichthyean fish, such as, but not limited to, sharks, rays, sturgeon, eels, anchovy, herring, carp, smelt, salmon, trout, hakes, cod, rockfish, bass, drum, mackerel, tuna, butterfish, catfish, flounder, and seabream.

[0048] A "crustacean" and the plural "crustaceans" are defined in this invention as any member of the Class *Crustacea*, such as, but not limited to, shrimp, lobsters, red claws, and crabs.

[0049] A "terrestrial animal" is one that lives primarily on land in a non-aquatic environment, such as, but not limited to cows, pigs, and chickens.

[0050] The term "phospholipid" refers to any lipid or fatty acid having a covalently attached a phosphate group in the molecular structure. These phospholipids are preferably sourced from vegetable material such as, but not limited to, soy, corn, palm, canola, rice,

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flax, coconut, combinations thereof, and are usually obtained as byproduct of the process of refining the vegetable oil. These phospholipids may be comprised of any of phosphatidyl choline (PC), phosphatidyl serine (PS), phosphatidyl ethanolamine (PE) and/or phosphatidyl inositol (PI), or a combination thereof.

[0051] The term "PUFA-rich phospholipid" means a phospholipid containing at least 20% fatty acids with 2 or more double bonds.

[0052] The term "carotenoid" encompasses any molecule in a class of yellow to red pigments, including carotenes and xanthophylls. "Carotenes" are orange-yellow to red pigments that are found in some animal tissues and plants, and may be converted to Vitamin A in the liver. "Xanthophylls" are yellow pigments, some of which may be found with chlorophyll in green plants.

[0053] Description

[0054] The inventors have found that a unique mix, including carotenoid compounds and PUFA-rich phospholipid (such as soy lecithin, DHA-, EPA- or ARA-rich phospholipid extracts) improves the bioavailability of carotenoids when consumed by coldwater fish. Additionally, the phospholipids increase oxidation stability of the carotenoids compared to other types of standard preparations. It is well documented that carotenoids are sensitive to photo- and thermal-oxidation, which results in major carotenoid losses during feed preparation and storage. Moreover, natural sources of carotenoids include a high level of saturated oils. Saturated oils become solidified at low water temperature and thereby reduce bioavailability of the carotenoid in the animal GI tract. The present invention overcomes the problems associated with standard carotenoid formulations by combining carotenoids with PUFA-rich phospholipid, where the phospholipid increases the efficacy of the carotenoid absorption at low temperatures.

[0055] The present invention relates broadly to formulations including carotenoids and PUFA-rich phospholipid compositions. Additionally, methods for producing such compositions and their use in formulation of novel feeds are disclosed.

[0056] Examples of phospholipid include, without limitation, phosphatidyl cholines (such as phosphatidyl choline (PC), dipalmitoylphosphatidylcholine (DPPC), other disaturated phosphatidyl cholines), phosphatidyl ethanolamines, phosphatidylinositol, phosphatidyl serines (sphingomyelin or other ceramides), various other phospholipids,

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phospholipid-containing oils (such as lecithin oils derived from soy beans), or mixtures and combinations thereof. The phospholipids of the present formulation can also be found in PUFA-rich extracts of single cell organisms such as, but not limited to, *Crypthecodinium* sp., *Schizochytrium* sp., *Mortierella* sp. and *Paracoccus* sp. Phospholipids of the present invention can also be derived from animal sources including, but not limited to, animal organ extracts (*e.g.*, brain, liver, other animal process wastes), egg yolk, egg yolk extracts, fish byproducts and fish byproduct extracts (*i.e.*, processed waste products from preparation of fish meal or purified fish oil). Preferred phospholipids are from *Crypthecodinium* sp., *Schizochytrium* sp. and *Mortierella* sp., and plant lecithins. Phospholipids useful for this invention would be those wherein at least 20% of the fatty acid residues have 2 or more double bonds. Preferred phospholipids would be those containing at least 20% of the fatty acid residues with 3 or more double bonds. Particularly preferred phospholipids would be those with 4 or more double bonds. Most particularly preferred phospholipids would be those containing at least 20% of the fatty acid residues with 4 or more double bonds.

[0057] Generally, the weight ratio of carotenoids to PUFA-rich phospholipid is between about 2:1 and about 1:100, with ratios between about 2:1 and 1:50 being preferred and ratios between about 1:1 and 1:10 being particularly preferred and ratios between about 1:1 and about 1:5 being especially particularly preferred.

[0058] The effective amount of the carotenoids for use in the composition of this invention ranges from about 0.1 mg per kg feed to about 1000 mg per kg feed depending on the carotenoids and the phospholipid used in the composition. Amounts between about 1 mg per kg feed to about 500 mg per feed being preferred, with amounts between about 2 mg per kg feed and 50 mg per feed being particularly preferred. A sufficient amount of phospholipid is generally an amount of phospholipid between about 0.01 mg per mg carotenoids and about 5000 mg per mg carotenoids, with amounts between about 0.5 mg per mg carotenoids and 2500 mg per mg carotenoids being preferred, and amounts between 2 mg per mg carotenoids and about 250 mg per mg carotenoids being particularly preferred, and amounts between about 2 mg per mg carotenoids being particularly preferred, and amounts between about 2 mg per mg carotenoids and about 250 mg per mg carotenoids being particularly preferred, and amounts between about 2 mg per mg carotenoids being particularly preferred, and amounts between about 2 mg per mg carotenoids and about 2 mg per mg carotenoids and about 100 mg per mg carotenoids being per mg carotenoids b

[0059] The compositions of the present invention can be in any desirable form, including, without limitation, a solid (such as a powder, granules, a semi-solid such as a

paste or the like), an emulsion, or a solution. An emulsion means that an oil or aqueous form of the compositions of this invention is emulsified in an aqueous solution. In addition, the emulsion can be a standard emulsion or a micro-emulsion where the mixture is forced through a nozzle or in other methods that generate micro-emulsions. Solutions of this invention employ a suitable solvent in which the composition is soluble or highly soluble. [0060] Generally, the compositions of this invention are formulated to be directly mixed with other feed ingredients prior to processing. However, the formulations can also be emulsified or blended with a carrier oil to top-coat the feed after processing.

[0061] In formulations of this invention that combine a phospholipid, such as lecithin, and a carotenoid, such as astaxanthin, the phospholipid acts to prevent oxidation of the carotenoids as well as to improve its solubility. Thus, the formulations of this invention, which supplement carotenoids with phospholipids, show significantly more stability, thus removing a major impediment that severely limits the utility of natural carotenoids in feed preparation. The carotenoid/phospholipid formulations of this invention not only have increased stability, but the formulations also increase the bioavailability of the carotenoids when taken by coldwater animals. Current carotenoid formulations contain large quantities of high melting temperature oils. These preparations therefore lose a major part of their effectiveness when taken by coldwater species due to the phase of the oil (i.e., solid). The carotenoids of the invention associate with PUFA-rich phospholipids in such a way as to preserve their liquidity and become more available for uptake in the small intestines, especially at low temperatures. Additionally, it is thought that the PUFA-rich phospholipidcarotenoid formulations of this invention improve carotenoid bioavailability by interfering with the interaction of carotenoids with other feed components during digestion in the fish stomach, permitting carotenoids to exit the stomach in a bioavailable form.

[0062] For example, the carotenoids (naturally produced by a single celled-organism or synthetic) can be combined with different concentrations of either purified phospholipids or crude phospholipids. For example, PC is available in a purified form comprising > 90% PC or in crude extracts from soybeans in de-oiled and oiled states (American Lecithin Company). Crude phospholipid extracts containing over 40% DHA or ARA of total fatty acids are also available (Advanced BioNutrition Corp., Columbia, MD). The presence of PUFA-rich phospholipid, such as lecithin, in the formulations of this invention prevents carotenoid solidification, thereby increasing bioavailability of carotenoids in the GI tract of

coldwater species. Thus, the presence of a PUFA-rich phospholipid in the compositions of this invention allows a reduction in carotenoid dosages in feed and the shortening of the administration period prior to harvesting without loosing the desired coloring.
[0063] Further improvement in bioavailability may be achieved by the addition of an alkaloid, such as piperine, to the carotenoid/phospholipid composition.

[0064] The addition of PUFA-rich phospholipids can also significantly increase the bioavailability of the carotenoids. This is an improvement, since in certain instances carotenoids have bioavailabilities of about 50% or less necessitating relatively large doses of the carotenoids for a longer period of time. The PUFA-rich phospholipids result in improved bioavailability of the carotenoids especially by coldwater species. The improved bioavailability can range from about a 20% increase to as much as about a 60% or greater increase by carefully choosing the type of PUFA-rich phospholipid and the ratio of the carotenoids and PUFA-rich phospholipids.

[0065] It should be noted that a number of substances that are used as additives to enhance carotenoid absorption are known irritants or damaging agents of the GI mucosa. Therefore, these would be contraindicated for use with carotenoids. Such substances would include: short chain fatty acids (such as citric acid, decanoic acid, caprylic acid or the like), long-chain unsaturated free fatty acids (such as oleic acid or the like), detergents (such as BRIJ, TWEEN-80, sodium deoxycholate, or the like), and chelators of polyvalent metal cations (such as EDTA, EGTA, or the like).

[0066] Because of their degree of unsaturation, carotenoids are inherently prone to oxidative degradation. Preserving the integrity of the double bonds of the carotenoids through processing and storage is a critical problem in the preparation of feeds, food and supplements therefore containing such materials. At the same time the preservation of the double bonds of the carotenoids is critical for the efficacy of the carotenoid itself. Kyle and Becker (WO 00/54575) have described a process whereby a DHA-containing oil is stabilized by lecithin at levels up to 8% of the oil. AN additional aspect of this invention involves the combination of lecithin with the carotenoid containing material is in the stabilization of the carotenoid against oxidation.

[0067] Another aspect of the present invention is the combination of the lecithin with other cellular materials comprising long chain polyunsaturated fatty acids (LC-PUFAs). Microorganisms such as, but not limited to, Crypthecodinium, Schizochytrium,

Theraustochytrium, Ulkenia, Mortierella, etc. are prone to oxidation as a result of their high content of LC-PUFA. Schzochytrium, Thraustochytrium and Ulkenia, in particular, are very fragile and can release oil during the process of harvesting and drying. The use of high concentration of phospholipids (especially lecithin) during the drying process can impart a high degree of stability to the resulting dry biomass of these microorganisms and increase the bioavailability of the LC-PUFAs themselves. Lecithin to biomass ratios from about 1:100 to about 1:1 are effective in increasing stability and bioavailability of the oils.

[0068] Methods for Making Carotenoid/Phospholipid Compositions

[0069] One preferred class of compositions of this invention are compositions that include a carotenoid or carotenoids and PUFA-rich phospholipid or PUFA-rich phospholipids generally prepared by contacting carotenoid and phospholipid under conditions to promote molecular association of the carotenoid and phospholipid. Such conditions typically will include the use of mixing procedures that promote molecular interactions and associations, use of a solvent and/or buffer, and controlled physical parameters (such as temperature, pressure and time) to permit an optimal degree of interaction and association.

[0070] The chemical interaction is preferably performed by aggressive or vigorous mixing. Such mixing procedures include vortex mixing, other high shear mixing procedures, sonication, other molecular level mixing procedures, or the like. The time and temperature of mixing should be designed to maximize interactions between the carotenoids and the phospholipids without causing thermal or shear damage to the molecules themselves. Generally, the mixing time will range from about 5 minutes to several hours, with times ranging between 10 minutes and 1 hour being preferred.

[0071] Generally, the mixing temperature will range from ambient to a temperature of at least 10% below the lowest breakdown temperature for the carotenoids or phospholipids being mixed. Preferably, the temperature will be between ambient temperature to about 60°C.

[0072] In preparing the formulations of this invention, the carotenoids can be mixed with synthetic, purified naturally derived, or crude phospholipids or can be mixed with various grades of lecithin or other PUFA-rich oils obtained from single-celled organisms. Carotenoids may be in the form of pure carotenoid (synthetic or otherwise) or as cellular

material from high carotenoid microorganisms such as but not limited to Pfaffia or Heamatococcus and the mixture of phospholipids to microbial cell biomass may be in the range from 1 part phospholipid to from 1 to 100 parts cellular biomass. Especially useful phospholipid concentrations range from about 15 to about 93% PC by weight. Moreover, the formulations can use either de-oiled or oil-based phospholipid preparations.

[0073] Regardless of the form of the phospholipid, generally the ratio of carotenoids to phospholipids ranges from about 1:100 to about 10:1, preferably, from about 1:25 to about 2:1, and particularly from about 1.0:10.0 to about 1.0:1.0.

[0074] In formulations using de-oiled phospholipids, the de-oiled phospholipids are initially dissolved in an organic solvent such as ethanol, and then mixed with carotenoids. This is followed by mixing, such as vortexing and/or sonication mixing. In formulations using oiled phospholipids, the oil-based phospholipids are simply combined with a carotenoid compound and mixed by vortexing and/or sonication, if needed. Sonication or mixing temperatures are preferably between ambient and about 60°C.

[0075] Another preferred process for making the compositions of this invention includes the dissolving of phospholipids and carotenoids in a polar solvent. Suitable solvents include, without limitation, chlorocarbons (such as chloroform, or the like), lower alcohols (such as methanol, ethanol, isopropanol or the like), or any other solvent in which the phospholipids and the carotenoids have some solubility, and the solvent is removable, e.g., by evaporation, or the like.

[0076] Methods for making LC-PUFA phospholipids compositions. In preparing the formulations of this invention, the LC-PUFA-containing biomass such as, but not limited to Schyzochytrium, can be mixed with synthetic, purified naturally derived or crude phospholipids or can be mixed with various grades of lecithin or other PUFA-rich oils obtained from single cell organisms. Especially useful phospholipids concentrations ranging from about 15 to about 93% PC by weight. Moreover, the formulations can use either de-oiled and oiled-based phospholipids preparations. Mixtures of phospholipids_and cellular material containing LC-PUFAs can range from 1 part to from 1 to 100 parts cellular material.

[0077] Examples

[0078] The following examples are included for example only to illustrate the preparation of compositions of present invention containing a carotenoids and PUFA-rich phospholipid, and are in no way meant to limit the scope or teaching of this invention.

[0079] Example 1

[0080] Preparation of a composition of synthetic astaxanthin and soy lecithin.
[0081] A sample of 60 g of soy lecithin (American Lecithin Co) was dissolved in ethanol, 30 g synthetic astaxanthin (AHD International, Atlanta, GA) was added, the mixture sonicated at 60°C for 5 minutes, and the solvent evaporated under vacuum. The resulting powder can be incorporated with other feed ingredients or dissolved in oil and top-coated onto the feed particles.

[0082] Example 2

[0083] Preparation of a composition of *Haematococcus* (containing natural astaxanthin) and phospholipid extract from *Crypthecodinium* species.

[0084] A sample of 50 g of algal phospholipids (Advanced BioNutrition, Columbia, MD) and 100 g *Haematococcus* (Naturose, Cyanotech Corporation Kailua-Kona, HI) were mixed vigorously for 1 h at room temperature. The mixture was dissolved in 850 ml of Menhaden oil (Omega Protein, Houston, TX) and used to top-coat standard fish feed pellets. The feed pellets were top coated at a level of 20 g of the above mixture per kg feed. This produced a feed containing about 50 mg astaxanthin per kg feed. This feed was then used to color the flesh of aquatic animals that consumed the feed.

[0085] Example 3

[0086] Preparation of a composition of *Phaffia rhodozyma* yeast biomass and phospholipid extract from *Crypthecodinium* sp.

[0087] *Phaffia* yeast was grown under standard conditions in a fermentor and biomass was harvested by centrifugation and diluted to 30% solids with water. Then 13.3 g of algal phospholipids (8 g on a dry weight basis) (ABN, Columbia, MD) was mixed vigorously with 333 g of the *Phaffia* slurry (100 g on a dry weight basis) to facilitate molecular association between the carotenoid and the phospholipids. The material was then dried on a rotary drum dryer at low temperatures and the resulting flakes were milled under liquid

nitrogen to produce a coarse powder. The resulting powder was then mixed with a commercial trout feed and cold pressed into feed pellets (1.2 - 2.0 mm, Ziegler Bros Inc. Gardners, PA) using standard techniques.

[0088] Example 4

[0089] Preparation of a composition of *Phaffia rhodozyma* yeast biomass and soy lecithin.

[0090] One hundred grams of *Phaffia* yeast biomass (Archer-Daniels-Midland Company, Decatur, IL) was mixed with water to give a slurry with a 30% water content. Eight g of soy lecithin (American Lecithin Co) was added to the slurry and the resultant mixture was homogenized vigorously to facilitate molecular association between the carotenoid and the phospholipids. The slurry was then dried in a freeze dryer and collected as a powder. This material had the following composition: 1.5% astaxanthin, 8% phospholipid, 50% fatty acids with 2 or more double bonds, and 20% of the fatty acids with 4 or more double bonds. This mixture was then incorporated into 10 kg commercial fishmeal pellets using standard methods with cold pressing or cold extrusion (Ziegler Bros Inc. Gardners, PA).

[0091] Example 5

[0092] Feeding of trout fish with a feed containing natural astaxanthin from *Phaffia* and a PUFA-containing phospholipid.

[0093] Five diets were prepared by Ziegler Bros Inc. (Gardners, PA) according to the following compositions:

[0094] Diet 1 contained 12.5 g *Phaffia* biomass per kg feed (100 mg astaxanthin/kg feed).

[0095] Diet 2 contained 13.8 g of the composition described in Example 3 per kg feed (100 mg astaxanthin/kg feed).

[0096] Diet 3 contained 7.6 g of the composition described in Example 3 per kg feed (50 mg astaxanthin/kg feed).

[0097] Diet 4 contained no *Phaffia* (0 mg astaxanthin/kg feed).

[0098] Diet 5 contained 7.6 g of the composition described in Example 4 per kg feed (50 mg astaxanthin/kg feed).

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[0099] Five groups of 20 trout fish per group were fed 4.4% body weight/day for 21 days. White muscle tissues were sampled from 5 fish in each group on day 21 and freezedried for 48 h. Total carotenoids were extracted from the tissues by homogenizing in 5 ml of absolute ethanol and 5 ml ethyl acetate. The homogenates were centrifuged (1000 x g for 5 min) and the supernatants dried under a stream of nitrogen and dissolved in 2 ml of hexane. Total carotenoids were measured spectrophotometrically at 470 nm.

[0100] The effect of the diet on muscle pigmentation is presented in Table 1:

	Absorbance at 470 nm	
Diet 1	0.19	
Diet 2	0.30	
Diet 3	0.11	
Diet 4	0.05	
Diet 5	0.14	

[0101] As can be seen from Table 1, Diet 4, with no *Phaffia* and no astaxanthin, provided the least amount of muscle pigmentation indicative of carotenoid content ($A_{470} = 0.05$). Diet 3 and Diet 5, with no *Phaffia* and 50 mg astaxanthin provided by the compositions of Example 3 and Example 4, respectively, provided intermediate amounts of muscle pigmentation. Diet 1, with *Phaffia* biomass providing twice as much, i.e., 100 mg astaxanthin, provided only a slightly higher amount of coloration than Diets 3 and 5. Diet 2, with no *Phaffia* and 100 mg astaxanthin provided by the composition of Example 3, provided the highest amount of coloration. It improves the muscle coloring by 56%, compared to Diet 1.

[0102] Example 6

[0103] Preparation of Schyzochytrium biomass with a high degree of oxidative stability.
[0104] Schizochytrium biomass is produced using conventional fermentation
technology and harvested by centrifugal harvesting processes to a solid content of about
20%. To this 100g of slurry (20g dry weight Schizochytrium containing about 10g of LCPUFA enriched oil) 2 g of soy lecithin (American Lecithin Co.) is added. The resultant

mixture is thoroughly mixed and then dried using a rotary drum dryer, or any other drying process and collected as powder of flake. The resulting flake product has a high degree of oxidative stability and bioavailability relative to a similar product produced without the lecithin treatment.

[0105] While this invention has been described fully and completely, it should be understood that, within the scope of the appended claims, the invention may be practiced otherwise than as specifically described. Although the invention has been disclosed with reference to its preferred embodiments, from reading this description those of skill in the art may appreciate changes and modification that may be made which do not depart from the scope and spirit of the invention as described above and claimed hereafter.

[0106] All references cited herein are incorporated by reference, including the following.

- [0107] Patent References
- **[0108]** US 6,261,598
- **[0109]** US 6,476,010
- **[0110]** US 6,436,437
- **[0111]** US 6,403,056
- **[0112]** US 6,358,524
- [0113] US 6,296,877
- **[0114]** US 6,413,736
- **[0115]** US 6,022,701
- [0116] US 5,972,642
- [**0117**] US 5,935,808
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- [0119] EP-A-0 410 236
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CLAIMS

We Claim:

1. A composition comprising at least one carotenoid and at least one phospholipid, wherein the carotenoid comprises at least 1% of the total mass and the phospholipid comprises at least 5% of the total mass, and the ratio of carotenoid to phospholipid is from about 1:100 to about 1:0.01.

2. The composition of Claim 1, wherein the carotenoid has a microbial source.

3. The composition of Claim 2, wherein the microbial source is chosen from *Phaffia*, *Haematococcus*, Schizochytrium and *Paracoccus*.

4. The composition of Claim 1, wherein the carotenoid is chosen from astaxanthin, zeaxanthin, canthaxanthin, lutein, beta-carotene, and lycopene.

5. The composition of Claim 1, wherein the carotenoid is synthetic.

6. The composition of Claim 1, wherein the phospholipid comprises more than about 20% polyunsaturated fatty acids having two or more double bonds.

7. The composition of Claim 1, wherein the phospholipid comprises more than about 10% polyunsaturated fatty acids having three or more double bonds.

8. The composition of Claim 1, wherein the phospholipid comprises more than about 10% polyunsaturated fatty acids having four or more double bonds.

9. The composition of Claim 1, wherein the phospholipid comprises more than about 20% polyunsaturated fatty acids having four or more double bonds.

10. The composition of Claim 1, wherein the phospholipid is of microbial origin.

11. The composition of Claim 1, wherein the phospholipid is an egg lecithin.

12. The composition of Claim 1, wherein the phospholipid has an origin chosen from fish, crustacean, and shellfish.

13. The composition of Claim 1, wherein the phospholipid is of mammalian origin.

14. The composition of Claim 13, wherein the phospholipid is of mammalian brain origin.

15. The composition of Claim 1, wherein the ratio of carotenoids to phospholipids is from about 1:50 to about 2:1.

16. The composition of Claim 1, wherein the ratio of carotenoids to phospholipids is from about 1:10 to about 1:1.

17. The composition of Claim 1, wherein the ratio of carotenoids to phospholipids is from about 1:5 to about 1:1.

18. An animal feed comprising the composition of any of Claims 1 to 17, wherein the carotenoid content is between about 0.1 mg and about 1000 mg per kg of feed.

19. The composition of Claim 18, wherein the animal is an aquatic animal.

20. The composition of Claim 19, wherein the aquatic animal is typically cultured below a temperature of 20°C.

21. The composition of Claim 19, wherein the aquatic animal is a fish.

22. The composition of Claim 19, wherein the aquatic animal is a crustacean.

23. The composition of Claim 18, wherein the animal is a terrestrial animal.

24. The composition of Claim 23, wherein the terrestrial animal is a bird.

25. A method of pigmenting an animal or animal-derived product, comprising providing a feed as described in Claims 18-24.

26. The method of Claim 25, wherein the animal-derived product is chosen from an egg and a processed egg product.

27. The method of Claim 25, wherein the animal or animal-derived product is chosen from a whole animal, the processed flesh of an animal, and a processed animal product.

28. A method of preparing an animal feed, feed supplement, or feed ingredient by (a) first mixing at least one carotenoid and at least one phospholipid, wherein the carotenoid comprises at least 1% of the total mass and the phospholipid comprises at least 5% of the total mass, wherein the ratio of the carotenoid to phospholipid is from about 1:100 to about 1:0.01; and

(b) then combining the carotenoid/phospholipid mixture with animal feed, feed supplement, or feed ingredient to provide a final carotenoid content between about 0.1 mg and about 1000 mg per kg; and

(c) then processing the feed, feed supplement, or feed ingredient into a deliverable form.

29. A method as in Claim 28, wherein the deliverable form comprises a pelleted feed.

30. A method as in Claim 28, wherein more than one carotenoid is mixed with the phospholipid.

32. A method of preparing an animal feed by:

(a) first mixing at least one carotenoid and at least one phospholipid, wherein the carotenoids comprise at least 1% of the total mass and the total phospholipids comprise at least 5% of the total mass, and the ratio of the carotenoid to phospholipid is from about 1: 00 to about 1: 0.01;

(b) then combining the mixture with a carrier in a ratio of about 1 to about 100 parts carrier:mixture; and

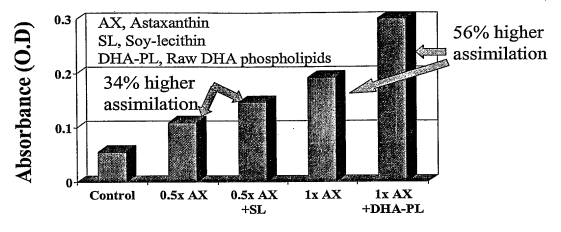
(c) then coating standard feed pellets with a composition comprising the mixture and the carrier.

33. The method of Claim 32, wherein the carrier comprises an oil.

34. The method of Claim 32, wherein the coating comprises a top coating.

Figure 1.

Improved Total Carotenes (TC) Assimilation With DHA-PL (Trout)





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INTERNATIONAL SEARCH REPORT

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International application No.

PCT/US04/19972

A. CLASSIFICATION OF SUBJECT MATTER IPC(7) : A61K 31/065					
USCL	: 514/726	tand designation and IDC			
	<u>International Patent Classification (IPC) or to both 1</u> DS SEARCHED	national classification and IPC			
	Minimum documentation searched (classification system followed by classification symbols)				
U.S. : 5					
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C. DOC	UMENTS CONSIDERED TO BE RELEVANT				
Category *	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.		
A	US 6,261,598 B1 (RUNGE et al.) 17 July 2001 (17	.07.2001), see the netire document.	1-32		
Furthe	r documents are listed in the continuation of Box C.	See patent family annex.			
	Special categories of cited documents:	"T" later document published after the inte date and not in conflict with the applic			
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EFS ID:	16476339					
Application Number:	12057775					
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Confirmation Number:	1945					
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS					
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Customer Number:	72960					
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File Listing:						
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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Inge Bruheim, et alConfirmation:1945Serial No.:12/057,775Group No.:1651Filed:03-28-2008Examiner:Ware, Deborah K.Entitled:BIOEFFECTIVE KRILL OIL COMPOSITIONS

INFORMATION DISCLOSURE STATEMENT LETTER

EFS Web Filed Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir or Madam:

The citations listed in the attached **IDS Form SB08A** may be material to the examination of the aboveidentified application, and are therefore submitted in compliance with the duty of disclosure defined in 37 C.F.R. §§ 1.56 and 1.97.

Applicants wish to bring to the Examiner's attention that we are not providing copies of US Patents as instructed under 37 CFR 1.98(a)(2). The Examiner is requested to make these citations of official record in this application.

CERTIFICATION STATEMENT

Applicants wish to bring to the Examiner's attention that the references supplied in this IDS are from a June 10, 2013 EP Search Report (copy attached). The present IDS is filed within three months of the mailing of the Search Report; therefore, no fees are due.

This Information Disclosure Statement under 37 C.F.R. §§ 1.56 and 1.97 is not to be construed as a representation that a search has been made, that additional information material to the examination of this application does not exist, or that any one or more of these citations constitutes prior art.

The Commissioner is hereby authorized to charge any required fees or credit any overpayments to Attorney Deposit Account No.: **50-4302**, referencing Attorney Docket No.: **NATNUT-14409/US-5/ORD**.

Dated: <u>August 1, 2013</u>

<u>/J. Mitchell Jones/</u> J. Mitchell Jones Registration No. 44,174 CASIMIR JONES, S.C. 2275 Deming Way, Suite 310 Middleton, WI 53562 608.662.1277

PTO/SB/81 (01-09) Approved for use through 11/30/2011, OMB 0651-0035 Trademark Office; U.S. DEPARTMENT OF COMMERCE LLC Detect and

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POWER OF ATTORNEY	Application Number	12/057,775
OR	Filing Date	28-Mar-2008
REVOCATION OF POWER OF ATTORNEY	First Named Inventor	Inge Bruheim
WITH A NEW POWER OF ATTORNEY	Title	BIOEFFECTIVE KRILL OIL COMPOSITION
	Art Unit	1651
	Examiner Name	Ware, Deborah
HANGE OF CORRESPONDENCE ADDRESS	Attorney Docket Number	AKBM-14409/US-5/ORD
hereby revoke all previous powers of attorney given	in the above-identified a	application
A Power of Attorney is submitted herewith.		
 OR I hereby appoint Practitioner(s) associated with the followin Number as my/our attorney(s) or agent(s) to prosecute the identified above, and to transact all business in the United and Trademark Office connected therewith: 	application	72960
OR I hereby appoint Practitioner(s) named below as my/our att to transact all business in the United States Patent and Tra	orney(s) or agent(s) to prosed idemark Office connected the	cute the application identified above, and rewith:
Practitioner(s) Name	R	egistration Number
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USPT0 to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is estimated to take 3 minutes to complete including gathering, preparing, and submitting the completed application form to the USPT0. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

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STATEMENT	[UNDER 37 CFR 3.73(b)
Applicant/Patent Owner: Inge Bruheim	
Application No./Patent No.: 12/057,775	Filed/Issue Date: 28-Mar-2008
Titled: BIOEFFECTIVE KRILL OIL COMPOSITIONS	
AKER BIOMARINE AS, a	corporation
(Name of Assignee)	(Type of Assignee, e.g., corporation, partnership, university, government agency, etc.
states that it is:	
1. X the assignee of the entire right, title, and interest i	in;
2. an assignee of less than the entire right, title, and (The extent (by percentage) of its ownership inter	interest in rest is%); or
3 the assignee of an undivided interest in the entire	ty of (a complete assignment from one of the joint inventors was made)
the patent application/patent identified above, by virtue of eith	her:
A. An assignment from the inventor(s) of the patent the United States Patent and Trademark Office a copy therefore is attached.	application/patent identified above. The assignment was recorded in t Reel, or for which a
OR	
	application/patent identified above, to the current assignee as follows:
1. From: INVENTORS	To: AKER BIOMARINE ASA
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Additional documents in the chain of title are list	ed on a supplemental sheet(s).
As required by 37 CFR 3.73(b)(1)(i), the documentar or concurrently is being, submitted for recordation pur	ry evidence of the chain of title from the original owner to the assignee was rsuant to 37 CFR 3.11.
[NOTE: A separate copy (<i>i.e.</i> , a true copy of the origination accordance with 37 CFR Part 3, to record the assignment of the sestimation of the second seco	inal assignment document(s)) must be submitted to Assignment Division ir nent in the records of the USPTO. <u>See</u> MPEP 302.08]
The undersigned (whose title is/supplied below) is authorized	d to act on behalf of the assignee. March $1,2013$
Signature	Date
HALLVARD MURI	CEO
Printed or Typed Name	Title
This collection of information is required by 37 CFR 3.73(b). The information	is required to obtain or retain a benefit by the public which is to file (and by the USPTO to

This collection of information is required by 37 CFR 3.73(b). The information is required to obtain or retain a benefit by the public which is to file (and b) the USP10 to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USP10. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sant to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

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Privacy Act Statement

The **Privacy Act of 1974 (P.L. 93-579)** requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b)(2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

The information provided by you in this form will be subject to the following routine uses:

- 1. The information on this form will be treated confidentially to the extent allowed under the Freedom of Information Act (5 U.S.C. 552) and the Privacy Act (5 U.S.C 552a). Records from this system of records may be disclosed to the Department of Justice to determine whether disclosure of these records is required by the Freedom of Information Act.
- A record from this system of records may be disclosed, as a routine use, in the course of
 presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to
 opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
- 4. A record in this system of records may be disclosed, as a routine use, to a contractor of the Agency having need for the information in order to perform a contract. Recipients of information shall be required to comply with the requirements of the Privacy Act of 1974, as amended, pursuant to 5 U.S.C. 552a(m).
- A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
 A record in this system of records may be disclosed, as a routine use, to another federal
- A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (*i.e.*, GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
- 8. A record from this system of records may be disclosed, as a routine use, to the public after either publication of the application pursuant to 35 U.S.C. 122(b) or issuance of a patent pursuant to 35 U.S.C. 151. Further, a record may be disclosed, subject to the limitations of 37 CFR 1.14, as a routine use, to the public if the record was filed in an application which became abandoned or in which the proceedings were terminated and which application is referenced by either a published application, an application open to public inspection or an issued patent.
- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.

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EFS ID:	15093234					
Application Number:	12057775					
International Application Number:						
Confirmation Number:	1945					
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS					
First Named Inventor/Applicant Name:	Inge Bruheim					
Customer Number:	72960					
Filer:	John Mitchell Jones/Vickie Hoeft					
Filer Authorized By:	John Mitchell Jones					
Attorney Docket Number:	AKBM-14409/US-5/ORD					
Receipt Date:	01-MAR-2013					
Filing Date:	28-MAR-2008					
Time Stamp:	14:10:55					
Application Type:	Utility under 35 USC 111(a)					

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New Applications Under 35 U.S.C. 111

If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application.

National Stage of an International Application under 35 U.S.C. 371

If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course.

New International Application Filed with the USPTO as a Receiving Office

If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application. Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

Application Number		12057775	
Filing Date		2008-03-28	
First Named Inventor	Inge Bruheim		
Art Unit		1651	
Examiner Name	Ware, Deborah K.		
Attorney Docket Numb	er	AKBM-14409/US-5/ORD	

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	Application Number		12057775	
	Filing Date		2008-03-28	
INFORMATION DISCLOSURE	First Named Inventor Inge Bruheim		Bruheim	
(Not for submission under 37 CFR 1.99)	Art Unit		1651	
	Examiner Name	Ware,	, Deborah K.	
	Attorney Docket Numb	er	AKBM-14409/US-5/ORD	

Examiner Initials*	Cite No	ude name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item ok, magazine, journal, serial, symposium, catalog, etc), date, pages(s), volume-issue number(s), isher, city and/or country where published.					
	1	CN Office Action mailed April 27, 2012, JP Patent Application No. 200880112125.6 (and English translation)	X				
	2	FRICKE, et al., Lipid, Sterol and Fatty Acid Composition of Antarctic Krill (Euphausia superba Dana), Lipids (1984) 19 (11): 821-827.					
	3	FRICKE, et al., 1-O-Alkylglycerolipids in Antarctic Krill (Euphausia Superba Dana), Comp. Biochem. Physiol. (1986) 85B(1): 131-134					
	4	GORDEEV, K.Y., et al. "Fatty Acid Composition of the Main Phospholipids of the Antarctic Krill, Euphausia superba," Chem. Nat. Cmpds. (1990) 26(2), pp. 143-147					
	5	GRANTHAM (1977) Southern Ocean Fisheries Survey Programme, FAO Rome, GLO/SO/77/3: 1-61.					
	6 RAVENTOS et al., Application and Posssibilities of Supercritical CO2 Extraction in Food Processing Industry: An Overview, Food Science and Technology International (2002) 8: 269-284						
	7 TANAKA, T., et al., Platelet-activating Factor (PAF)-like Phospholoipds Formed during Peroxidation of Phosphatidylcholines from Different Foodstuffs, Biosci. Biotech. Biochem. (1995) 59 (8), pp. 1389-93						
	8 WINTHER, et al., Elucidation of Phosphatidylcholine Composition in Krill Oil Extracted from Euphausia superba, Lipids (2011) 46: 25-36						
If you wish to add additional non-patent literature document citation information please click the Add button Add							
EXAMINER SIGNATURE							
Examiner	Signa	ure Date Considered					
*EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through a citation if not in conformance and not considered. Include copy of this form with next communication to applicant.							

	Application Number		12057775
	Filing Date		2008-03-28
INFORMATION DISCLOSURE	First Named Inventor	Inge E	Bruheim
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit		1651
	Examiner Name	Ware,	, Deborah K.
	Attorney Docket Number		AKBM-14409/US-5/ORD

¹ See Kind Codes of USPTO Patent Documents at <u>www.USPTO.GOV</u> or MPEP 901.04. ² Enter office that issued the document, by the two-letter code (WIPO Standard ST.3). ³ For Japanese patent documents, the indication of the year of the reign of the Emperor must precede the serial number of the patent document. ⁴ Kind of document by the appropriate symbols as indicated on the document under WIPO Standard ST.16 if possible. ⁵ Applicant is to place a check mark here if English language translation is attached.

	Application Number		12057775	
	Filing Date		2008-03-28	
INFORMATION DISCLOSURE	First Named Inventor Inge B		Bruheim	
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit		1651	
	Examiner Name	Ware,	, Deborah K.	
	Attorney Docket Number		AKBM-14409/US-5/ORD	

CERTIFICATION	STATEMENT
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Please see 37 CFR 1.97 and 1.98 to make the appropriate selection(s):

That each item of information contained in the information disclosure statement was first cited in any communication from a foreign patent office in a counterpart foreign application not more than three months prior to the filing of the information disclosure statement. See 37 CFR 1.97(e)(1).

OR

That no item of information contained in the information disclosure statement was cited in a communication from a foreign patent office in a counterpart foreign application, and, to the knowledge of the person signing the certification after making reasonable inquiry, no item of information contained in the information disclosure statement was known to any individual designated in 37 CFR 1.56(c) more than three months prior to the filing of the information disclosure statement. See 37 CFR 1.97(e)(2).

See attached certification statement.

The fee set forth in 37 CFR 1.17 (p) has been submitted herewith.

X A certification statement is not submitted herewith.

SIGNATURE

A signature of the applicant or representative is required in accordance with CFR 1.33, 10.18. Please see CFR 1.4(d) for the form of the signature.

Signature	/J. Mitchell Jones/	Date (YYYY-MM-DD)	2012-11-15
Name/Print	J. Mitchell Jones	Registration Number	44174

This collection of information is required by 37 CFR 1.97 and 1.98. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 1 hour to complete, including gathering, preparing and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. **SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450**.

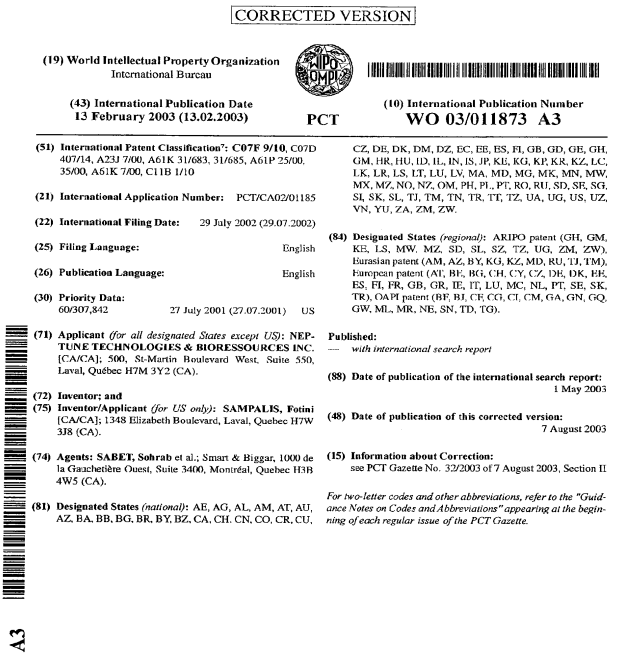
The Privacy Act of 1974 (P.L. 93-579) requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b)(2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

The information provided by you in this form will be subject to the following routine uses:

- The information on this form will be treated confidentially to the extent allowed under the Freedom of Information Act (5 U.S.C. 552) and the Privacy Act (5 U.S.C. 552a). Records from this system of records may be disclosed to the Department of Justice to determine whether the Freedom of Information Act requires disclosure of these record s.
- 2. A record from this system of records may be disclosed, as a routine use, in the course of presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
- 4. A record in this system of records may be disclosed, as a routine use, to a contractor of the Agency having need for the information in order to perform a contract. Recipients of information shall be required to comply with the requirements of the Privacy Act of 1974, as amended, pursuant to 5 U.S.C. 552a(m).
- 5. A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
- 6. A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
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- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.

(12) STANDARD PATENT (11) Application No. AU 2002322233 B2 (19) AUSTRALIAN PATENT OFFICE			
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. 56)	 56) Related Art HOSOKAWA et al, Journal of Agricultural and Food Chemistry 2000, Vol. 48, No. 10, Pages 4550-4554. HENDERSON et al, Lipids, 1994, Vol. 29, No. 5, Pages 311-317 WO 2000/023546 A1 (UNIVERSITE DE SHERBROOKE) 27 April 2000 EP 0209037 B1 (INVERNI DELLA BEFFA S.P.A) 28 February 1990 EP 0275005 B1 (INDENA S.p.A) 11 August 19 BELL et al "Molecular Species Composition of the Major Diacyl Glycerophospholipids from Muscle, Liver, Retina and Brain of Cod" Lipids, 1991, Vol. 26, No. 8, Pages 565-573 US 4963527 A (BOMBARDELLI et al) 16 October 1990 		

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)



VO 03/011873 A3

(54) Title: NATURAL MARINE SOURCE PHOSPHOLIPIDS COMPRISING FLAVONOIDS, POLYUNSATURATED FATTY ACIDS AND THEIR APPLICATIONS

(57) Abstract: A phospholipid extract from a marine or aquatic biomass possesses therapeutic properties. The phospholipid extract comprises a variety of phospholipids, fatty acid, metals and a novel flavonoid.

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NATURAL MARINE SOURCE PHOSPHOLIPIDS COMPRISING FLAVONOIDS, POLYUNSATURATED FATTY ACIDS AND THEIR APPLICATIONS

Cross-Reference to Related Application

This application claims the benefit of United States Provisional Patent Application Serial No. 60/307,842, filed July 27, 2001, which is incorporated herein by reference in its entirety.

Field of the Invention

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The present invention is directed to nutraceutical, pharmaceutical or cosmetic compositions, particularly to phospholipid compositions derived from natural marine or aquatic sources.

Background of the Invention

WO 92/21335 published on December 10, 1992 and corresponding United States Patent No. 5,434,183 issued on July 18, 1995 describes a phospholipid emulsion derived from marine and/or synthetic origin comprising polyunsaturated fatty acids and having anti-inflammatory and immunosuppressive effects and which promotes normal brain or retinal development and function. U.S. 5,434,183 does not disclose the presence of flavonoids or nervonic acid (a mono-unsaturated fatty acid) in the composition.

JP 2215351, published on August 28, 1990, discloses a 25 method for extracting and purifying phospholipids from fresh krill. Krill is lyophilized and then extracted with ethanol to produce an extract which is fractionated by absorption column chromatography to produce high purity phosphatidyl choline and phosphatidyl ethanolamine. There is no disclosure of a 30 composition comprising a flavonoid or nervonic acid.

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WO 00/23546, published on April 27, 2000, discloses methods for extracting lipid fractions from marine and aquatic animal material by acetone extractions. The resulting nonsoluble and particulate fraction is further solvent extracted with ethanol or ethylacetate to achieve further lipid extractions.

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Hosokawa et al. (35), published in 2000, discloses the conversion of docosahexanoic acid containing phosphatidylcholines (DHA-PC) from squid skin lecithin to docosahexanoic acid containing phosphadylserines (DHA-PS) via transphosphatidylation with phospholipase D (PLD). According to Table 2 of this reference, the fatty acid composition of the phospholipid includes important portions of eicosapentanoic acid. There is no disclosure concerning any pharmaceutical, nutraceutical, or cosmetic use of a composition comprising a flavonoid.

Henderson et al. (36), published in 1994, discloses lipid compositions of the pineal organ from rainbow trout comprising phospholipids. According to Table 4 of this reference, said phospholipids contain fatty acids corresponding 20 to eicosapentanoic and docosahexanoic acid. Similarly, Bell et al. (37), published in 1991, discloses phospholipid compositions derived from different organs of cod. Moreover, Wiegand et al. (38), published in 1983, discloses polyene derivatives of phosphatidylcholine as phospholipid molecular 25 species of frog receptor membranes. However, there is no disclosure in any of these references concerning any pharmaceutical, nutraceutical, or cosmetic use of a composition comprising a flavonoid.

WO 97/39759, published on October 30, 1997, discloses ω -3 fatty acids and ω -3 phosphatidylcholine in the treatment of bipolar disorder. The preferred ω -3 phosphatidylcholine

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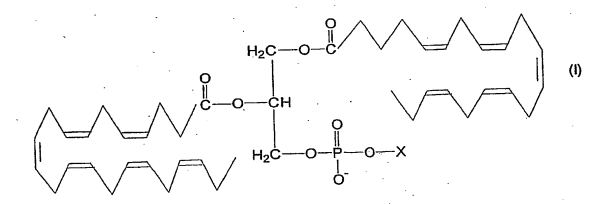
derivatives comprise eicosapentanoic and/or docosahexanoic acid. However, there is no disclosure concerning any pharmaceutical, nutraceutical, or cosmetic use of phospholipids beyond the treatment of bipolar disorder or the use of a composition comprising a flavonoid.

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EP 0609078 A1, published on March 8, 1994, discloses a phospolipid comprising two different unsaturated fatty acids, wherein a preferred phospholipid contains both eicosapentanoic and docosahexanoic acid. Furthermore, the phospholipid can be used in the preparation of foods, skin care preparations, or pharmaceutical agent. However, there is no disclosure concerning any pharmaceutical, nutraceutical, or cosmetic use of a composition comprising a flavonoid.

15 Summary of the Invention

In one aspect, the invention provides novel phospholipids, wherein the two fatty acids chains of the phospholipid are occupied by eicosapentanoic acid (EPA) and docosahexanoic acid (DHA) simultaneously, within the same molecule, i.e.: a phospholipid of the general formula (I):



wherein X represents a moiety normally found in a phospholipid.

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(I)

and

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According to a further aspect of the present invention there is provided a composition, comprising: (a) a phospholipid of the general formula (I),

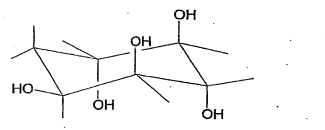
3

 H_2C-O-C $H_2C-O-CH$ $H_2C-O-CH$ $H_2C-O-P-O-X$

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wherein X is $-CH_2CH_2NH_3$, $-CH_2CH_2N(CH_3)_3$ or



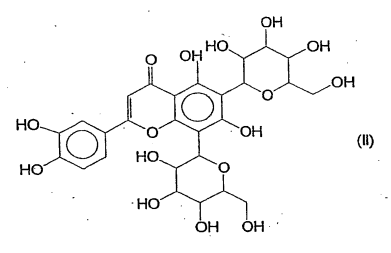
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(b) a flavonoid of the general formula (II),

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In a further aspect, the invention provides a novel flavonoid compound (II):



There is also provided a composition comprising the above noted phospholipid and flavonoid derived from a marine or aquatic biomass. The composition and the components are useful in the prevention or treatment of a variety of disease states and for the aesthetic enhancement of an animal, including human, body. Commercial packages containing the composition are also within the invention.

The novel phospholipids and the novel flavonoid compound are derived from an extract from a marine or aquatic biomass.

20 There is also provided a phospholipid extract comprising the above noted phospholipids and flavonoid compound derived from a marine or aquatic biomass. The extract and the components are useful in the prevention or treatment of a variety of disease states and for the aesthetic enhancement of 25 an animal, including human, body. Pharmaceutical, nutraceutical and cosmetic compositions containing the extract and uses thereof are also within the invention, as are commercial packages contain the compositions of the invention.

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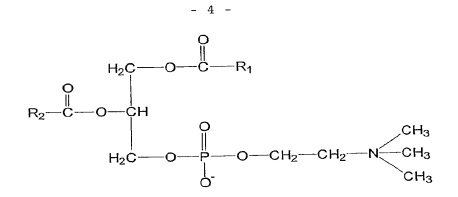
Detailed Description of the Invention

1. Phospholipids

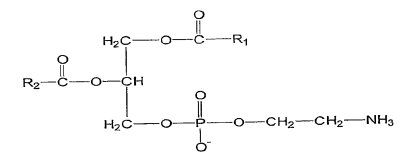
Phospholipids are complex lipids containing The phosphatides, known as phospholipids, are phosphorus. 5 usually divided into groups on the basis of compounds from which they are derived. In addition to two chains of fatty acids they contain phosphoric acid, glycerol and nitrogenous bases such as choline. Important phospholipids are phosphatidylcholine (PC), phosphatidylethanolamine (PE) and 10 phosphatidylinositol (PI). Their nature as amphophilic molecules provides them with unique physicochemical properties. Their function as the principle components of cell membranes makes phospholipids essential for all vital cell processes. They are wide spread as secretory and structural components of 15 the body and can mimic or enhance natural physiological process.

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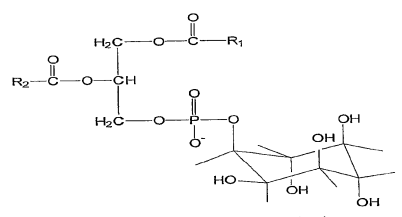


Phosphatidylcholine — common structure R_1 and R_2 are fatty acid residues, different for each molecular species



Phosphatidylethanolamine- common structure

 R_1 and R_2 are fatty acid residues, different for each molecular species



Phosphatidylinositol— common structure R_1 and R_2 are fatty acid residues, different for each molecular species

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Phospholipid production may be either synthetic or through extraction from natural tissues. The chief source of commercial natural phospholipids are soybean, egg yolk and cows (brain and liver). Since an individual phospholipid may

- 5 contain a variety of fatty acid residues, it may be described as pure only with this limitation in mind. Naturally occurring essential polyunsaturated fatty acids can contribute to the activation of cellular metabolism. The main fatty acid found in phospholipid products is linoleic acid (C18:2n6), present in
- 10 soybean at more than 65%. The longest chain polyunsaturated fatty acids found in commercially available phospholipids either as preparations or individually are 20:4 among the eicosanoids, known as arachidonic acid, and 22:6 known as docosahexanoic acid.

15 Arachidonic acid is a fatty acid that is found as part of phospholipid membranes, generally as part of phosphatidylcholine and phosphatidylinositol. Adverse cellular stimuli will activate enzymes (phospholipase) that cleave arachidonic acid from the phospholipid backbone in the cell

- 20 membrane. Arachidonic acid, which serves as the precursor for prostaglandins and prostacyclin (PGs, PGI₂) and thromboxane (TXs), can then be metabolized by one of two major pathways: the cyclooxygenase (COX) pathway or the lipoxygenase pathway. The COX pathway products, PGG₂ and PGH₂, can then be acted upon 25 by thromboxane synthase (in platelets) or prostacyclin synthase (in endothelium) to form TXs or PGI₂, respectively. Arachidonic acid can also be acted upon by 5-lipoxygenase, primarily in leukocytes, to form leukotrienes (LTs). One or more of these
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with arachidonic acid, i.e. inflammatory disease and pain.

Platelets, leukocytes, smooth muscle, and endothelium can produce vasoactive substances, products of arachidonic acid metabolsim such as prostaglandins (PGs), prostacyclin (PGI₂),

metabolites can mediate all the signs and symptoms associated

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leukotrienes (LTs), and thromboxanes (TXs). These substances can either act as vasodilators or as vasoconstrictors. PGI_2 is essential in vascular function since it inhibits platelet adhesion to the vascular endothelium and has significant

- 5 vasodilatation qualities. Damaged endothelial cells cannot produce PGI₂, making the vessel more susceptible to thrombosis and vasospasm. Thromboxanes and leukotrienes serve a vascular function during inflammation, generally producing vasoconstriction. Prostaglandins have a vascular role during 10 inflammation, and also play a more subtle role in normal flow
- regulation, most notably as modulators of other control mechanisms. Prostaglandins have both vasoconstrictor and vasodilator activities. Leukotrienes and prostaglandins can also increase the endothelial membrane permeability thus 15 promoting edema during inflammation. Arachidonic acid is naturally present in most phospholipid mixtures or emulsions
 - naturally present in most phospholipid mixtures or emulsions available today.
- Nervonic acid (C24:1) is also called selacholeic acid or tertracosenic acid. Nervonic acid is the predominant nutrient of white matter in glucoside, which is quantitatively 20 contained in nerve tissue and white matter. The absence of nervonic acid may result in cerebral lesion, fatigue, hypodynamia, amentia, and senile dementia. Nervonic acid, tertracosenic acid in another name, is monounsaturated, nonoxidable/decomposed and absorptive. It is called a rare tonic 25 as it is rare existent in nature. It may be obtained in small quantities by extracting from cerebral chrondriosome. Therefore, the substantance is far below the demand of human body. In foreign countries, nervonic acid mainly comes from shark brain and oil. 30

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1.1 Phosphatidylinositol Clinical Applications

Recent advances in nutritional and biochemical research have documented inositol as an important dietary and cellular constituent. Functions of phosphatidylinositol in 5 biological membranes include the regulation of cellular responses to external stimuli and/or nerve transmission as well as the mediation of enzyme activity through interactions with

Inositol has been identified as an important dietary 10 and cellular constituent. Biochemical functions:

a. Regulation of cellular responses to external stimuli

b. mediation of enzyme activity.

various specific proteins (1).

Phosphoinositide composition of the central nervous system cell membranes are fatty-acid enriched and consist 15 primarily of phosphatidylinositol (PI), phosphatidylinositol-4phosphate (PIP), and phosphatidylinositol-4,5-biphosphate (PIP2). Once the membrane is stimulated, phospholipase C is activated and consequently inositol triphosphate along with diacylglycerol is produced. PI is used as a precursor for 20 phosphatidylinositol-3-phosphate and 3,4,5-triphosphate (2).

Active transport carriers, calcium pumps in the cell membrane itself, and in the endoplasmic reticulum, keep cytoplasmic calcium concentration very low. Usually the calcium concentration inside the cytoplasm is 5,000-10,000

25 times less than the concentration in the extracellular fluid. This endoplasmic store of calcium can be accessed upon stimulation by inositol. Inositol triphosphate is released from the cell membrane and travels through the cytoplasm until it reaches the endoplasmic reticulum. This inositol then 30 releases the sequestered calcium, which can go on to mediate - 8 -

the release of neurotransmitters in response to depolarization (3).

In addition to releasing endoplasmic reticulum calcium, inositol functions as the major central nervous system non-nitrogenous osmoregulator. Modulation of this inositol pool is regulated in response to states of high or low osmolalities. The inositol pool is supplied via a sodium/inositol transporter, a sodium dependent active transport system, and a passive low affinity transporter (4,5).

10 Numerous non-inositol receptors have been identified in the central nervous system that can potentially interact with the inositol signaling system. Most of these receptors are linked to the G proteins and produce inositol-1,4,5triphosphate as second messengers. These receptors can be 15 found in nearly every human organ system. The potential interactions between these receptors and their agonists are responsible for regulation of the body on a day-to-day basis. In view of the complexity of these systems and their actions, a perfect balance is required for regulation of the signaling

20 systems.

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Theoretically, an imbalance of inositol concentration could potentially affect the development and function of one or all of these receptors. Cholinergic receptors are located in the liver, heart, stomach, and lungs. Serotonin and glutamine receptors are found mostly in the central nervous system (CNS) tissues. Adrenergic receptors are present in various tissues including CNS, vascular tissues, and heart. Histaminergic receptors are predominantly found in the lungs and stomach.

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Clinical Applications

A change in CNS availability of inositol may produce altered brain signaling and eventually lead to the development of neurological disorders.

5 a. Depression:

The pathophysiology of depression is believed to be linked to a deficiency of neurotransmitters at post-synaptic receptor sites. According to the catecholamine theory, the deficiency is in the amount of norepinephrine; in the

10 indolamine theory the deficiency is in the amount of serotonin. Receptors linked to the inositol signalling system include serotonin (5HT2a and 5HT2b) and norepinephrine (alpha 1a, 1b, and 1d).

In 1978, Barkai et al demonstrated depressed patients had significantly decreased cerebospinal fluid (CSF) levels of inositol as compared to healthy patients (6). In 1993 this theory was expanded to conclude that administration of highdose inositol could increase CSF levels by as much as 70 percent (7). This led to the study of inositol for treatment

- 20 of depression (8,9). In 1995 Levine et al completed a doubleblind study for treatment of depression using inositol at a dose of 12 grams daily compared to placebo. Patients receiving inositol showed significant improvement in depression as ranked by the Hamilton Depression Rating Scale (33.4 +/- 6 versus .6
- 25 +/- 10). Another important observation was the absence of manic episodes in the bipolar patients treated with inositol. This lack of manic episodes may suggest that when the signalling system is not overactive, addition of inositol will not increase the signalling system's activity (10,11). It can
- 30 be concluded that inositol is effective in managing the clinical manifestations of depression.

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b. Panic Disorder:

Benjamin et al expanded the clinical use of inositol by evaluating its effectiveness in panic disorder (12). This was an eight week double-blind, crossover study whereby patients were treated with inositol daily for four weeks and then crossed over to the other study arm. Improvement was assessed using patient diaries, the Marks-Matthews Phobia Scale, the Hamilton Anxiety Rating Scale, and the Hamilton Depression Scale. The frequency and severity of panic attacks and the severity of agoraphobia declined significantly more after inositol than after placebo (a decrease from 10 attacks per week to 3 per week in the treated group compared to a decrease from 10 to 6 in the placebo group). The authors conclude inositol's efficacy and safety, and the fact that inositol is a natural component of the human diet, make it a

potentially attractive therapeutic agent for panic disorder.

c. Obsessive Compulsive Disorder (OCD):

Since the phosphatidylinositol cycle, as a second messenger is known to affect several neurotransmitters, 20 including serotonin receptors, inositol was studied for treatment in OCD in a double-blind, placebo controlled, crossover trial. Thirteen patients were treated for six weeks. There was a significant improvement at week six during the inositol period when compared to placebo period. There were no 25 side-effects reported during the study period (1).

d. Alzheimer's Disease (AD):

Although the role of aluminum in AD is still speculative at best, the presence of aluminosilicates at the core of senile plaques in diseased neurons is a consistent feature found in the CNS of AD patients during autopsy. It is known that aluminum inhibits the incorporation of inositol into 5

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phospholipids and the hydrolysis of the phosphoinositides by binding to one of two specific phosphate groups. This binding of phosphate and aluminum affects the calcium releasing effects of the cell. The resulting profound disturbance of the phosphatidylinositol second messenger system may account for

neuronal malfunction and eventual cell death (13).

Since the potential role of aluminum as a causative agent for cell death may be affected by the deregulation of calcium concentration, possibly due to inositol depletion,

10 supplementation with inositol may produce positive CNS effects. Recent data suggests the loss of PI second messenger system target sites and IP3 receptors may add to cognitive impairment and the failure of conventional therapies in AD. Therefore, supplementation of inositol to replenish the diminished PI 15 system may be beneficial in the treatment of AD (13-20).

In 1996 Barak et al completed a double-blind, controlled, crossover study of six grams inositol daily compared to placebo for 30 days in 11 Alzheimer's patients. Patients in the study were diagnosed with dementia of the AD type as classified by DSM - IIIR and aged 65 years or older. The Cambridge Mental Disorder of the Elderly Examination (CAMDEX) was used as the basic assessment parameter and was administered upon admission into the study. Included in CAMDEX is part A: patient's present physical and mental state, part B:

- 25 Cognitive.Subscale of CAMDEX (CAMCOG), part C: interviewers observations, and part D: physical examination. CAMCOG was repeated at two, four, six, and eight weeks. Participants scored 80 or less on the CAMCOG examination and their symptoms of depression were not severe (21).
- 30 Patients were excluded from the study if they had a history of psychiatric, alcohol, and/or drug addiction disorders, or abnormalities in baseline laboratory values

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(blood count, electrolytes, liver or kidney functions, VDRL, or CT scan) not consistent with AD. Patients with additional neurologic, metabolic, endocrinologic disorders, or presence of internal disease that grossly impaired brain functioning were also excluded.

Subjects were given either three grams inositol or placebo in the morning and again in the evening. After four weeks patients were crossed over into the other arm (inositol or placebo) for an additional four weeks. Only benzodiazepines were allowed during the study period (15 mg of oxazepam or equivalent), provided the patient was receiving it on study entry.

Analysis of the improvement scores of all patients who completed the study showed inositol increased the total 15 CAMCOG score from a baseline of 31.36 +/- 20.90 to 40.09 +/-24.54, while the placebo group increased from baseline of 35.9 +/- 25.96 to 39.27 +/- 25. The authors concluded only two of the eight subscales (language and orientation) showed significant improvement with inositol.

20 Inositol's proposed mechanism of action in the CNS does not include direct manipulation with either pre- or postreceptors. However, it may indirectly affect the relationship between receptor and agonist. By mediating the physiochemical characteristics of the M1 pre-synaptic receptor (solubility, 25 osmolality, etc.), inositol may alter the binding site and influence the signaling that occurs as a result.

1.2 Aging

Phosphatidylcholine rich in polyunsaturated fatty acids is indispensable for cellular differentiation, 30 proliferation and regeneration. The physiologic functions of these phospholipids are related to the morphology of the

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biological membranes, the incorporation of these molecules into membranes and thus the maintenance of intact cell membranes.

The current study was designed to investigate the effects of Polyunsaturated phosphatidylcholine on age-related hearing loss by evaluating its ability to preserve mitochondrial function, protect mitochondrial DNA from oxidative damage and preserve auditory sensitivity (22).

Harlan-Fischer 344 rats, 18-20 months of age, were used as the experimental subjects.

10 The subjects were caged individually and maintained at 21 to 22° C in a 12:12 light-dark cycle b.

A dose of 300mg/kg/day of Polyunsaturated phosphatidylcholine was supplemented to each subject, by adding it to the oral diet.

15 The animals were divided randomly into two groups (n = 7 for each group). Group-1 served as the control, and group-2 as the experimental group.

At the onset of the study, Auditory Brainstem Responses were obtained to measure baseline hearing thresholds 20 in all subjects.

Age-associated changes in hearing sensitivities were then recorded at two-month intervals for six months.

In order to assess age-related changes in mitochondrial function, mitochondrial membrane potentials were 25 studied using flow cytometry. For this purpose, peripheral blood was obtained from each subject at the beginning and at the end of the protocol.

At the conclusion, the subjects were euthanized (according to NIH protocol), and tissue samples were obtained

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from brain and cochlea (stria vascularis and auditory nerve) to study mitochondrial DNA deletion associated with aging. This was achieved by amplifying the specific common aging mitochondrial deletion by Polymerase Chain Reaction. DNA

5 quantification was performed. The data obtained for each protocol was compared between the two groups and analyzed using ANOVA.

The effects of Polyunsaturated phosphatidylcholine on age-related hearing loss demonstrate a gradual age-associated decline in hearing sensitivities at all the frequencies tested (3, 6, 9, 12 and 18 kHz).

There was a statistically significant preservation of hearing noted in the treated subjects at all frequencies, which was observed at four and six months of treatment.

15 Overall, there was a continued decline in hearing in the control subjects and a statistically significant protective effect of Polyunsaturated phosphatidylcholine on the experimental subjects (p<.005).

Mitochondrial membrane potentials were recorded by 20 flow cytometry as a measure of the uptake of Rhodamine 123 by mitochondria.

The mean fluorescence intensity (MFI) in group-1 subjects measured 3190 and 2100 at the beginning and end of the study, respectively.

25 This, approximately, 30% decline in membrane potential with time was statistically significant (p=0.003).

Conversely, the MFI in the experimental group remained essentially unchanged at 2990 from 3165 at the beginning of the study.

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This difference between the control and treated groups was statistically significant (p<0.05), demonstrating the protective effect of polyunsaturated phosphatidylcholine supplementation on mitochondrial membrane potential.

Phospholipids are integral structural components of all biological membranes with polyunsaturated phosphatidylcholine and phosphatidylethanolamine being the predominant types, quantitatively. They constitute the phospholipid bilayer structure of cellular membranes, which is

- 10 responsible for membrane stability and cellular function. Polyunsaturated phosphatidylcholine maintains and promotes the activity of several membrane bound proteins and enzymes, including Na-K ATPase, adenylate cyclase and glutathione reductase. They are also known to be precursors of
- 15 cytoprotective agents such as eicosanoids, prostaglandins and antioxidants.

The results of these studies suggest that polyunsaturated phosphatidylcholine and phosphatidylethanolamine may protect mitochondrial function by

20 preserving the age-related decline in mitochondrial membrane potentials and hence their activity. The observation that there was less mitochondrial DNA damage in the treated group may explain the effect of preservation of hearing loss associated with aging, by the ability of polyunsaturated

25 phosphatidylcholine and phosphatidylethanolamine to specifically up-regulate cochlear mitochondrial function. There are many studies demonstrating the effects of mitochondrial metabolites on cognition and aging (22-33). Additionally, recent work has shown that acetyl-L-carnitine and -lipoic acid

30 delay the progression of age-related hearing loss by protecting cochlear mitochondrial DNA from oxidative damage (34). These results support the membrane hypothesis of aging and provide further evidence to support this theory as a possible

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explanation for age-related hearing loss. Thus, PPC may be one of many rational approaches to consider for the purpose of membrane preservation, enhanced mitochondrial function, reduction of age-associated mitochondrial DNA damage and

5 slowing of some of the aging processes.

1.3 Effect of phosphoglycolipid exract (NT factor) on normal and cancerous cells

Reduced levels of phospholipids in normal cells can limit metabolic activity and limit available energy. 10 Phospholipids, as part of the membrane structure:

i. maintain membrane integrity

ii. regulate enzyme activities and membrane transport processes through changes in membrane fluidity (Spector 1981, 1985)

- 15 iii. Signal transduction utilizes phospatidylcholine and phosphatidylinositol for the production of diacyl-glycerol (DAG) by phospholipase C (Berridge 1989) and for the production of inositol triphosphate (IP3) (Ranan 1990, Michell 1988, Margolis 1990).
- 20 iv. One of the choline phospholipids (1-alkyl-2 acetyl-SNglycerol-3-phosphocholine) is the substrate for the synthesis of platelet activating factor (Synder 1989).

v. The arachidonic acid found as part of the structure of choline or inositol phospholipid is utilized for the production of prostaglandin and leukotriene (Nordoy 1990).

vi. The choline of phosphatidylcholine may be used in neural tissue for the synthesis of acetylcholine (Blusztain 1987)

vii. Phosphoglycolipid improves cell maintenance and metabolic activity of normal cells.

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viii. Phosphatidylcholine derivatives disrupt cancer cells at concentrations that do not affect normal cells.

ix. Phosphatidylcholine is selectively cytotoxic to cancer cells in vitro (Hoffman 1986, Harmann 1986, Berger 1984).

5 a. Such compounds inhibit HL60 leukemic cells at a dosage that has no effect on normal human marrow cells, the tissue from which the leukemic cells are derived.

b. Normal cells were able to tolerate 4 times higherdosage than the leukemic cells during 24 hours incubation with10 the phospholipid preparation (Berdel 1986).

c. There was up to a 5-fold difference in sensitivity between the normal and tumor cells with breast, ovarian, and lung cancer cells, as well as with mesothelioma cells (Namba 1993).

15 1.4 Imaging

Polyunsaturated phospholipids are known to be important with regard to the biological functions of essential fatty acids, for example, involving neural tissues such as the brain and retina. The NMR spectra of polyunsaturated bilayers

- 20 are dramatically different from those of less unsaturated phospholipid bilayers. MD simulations can aid in interpreting the complex NMR spectra of polyunsaturated bilayers, in conjunction with electron density profiles determined from small-angle X-ray diffraction studies. This work clearly
- 25 demonstrates preferred helical and angle-iron conformations of the polyunsaturated chains in liquid-crystalline bilayers, which favor chain extension while maintaining bilayer flexibility. The presence of relatively long, extended fatty acyl chains may be important for solvating the hydrophobic
 30 surfaces of integral membrane proteins, such as rhodopsin. In

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addition, the polyallylic DHA chains have a tendency to adopt back-bended (hairpin-like) structures, which increase the interfacial area per lipid. Finally, the material properties have been analyzed in terms of the response of the bilayer to

5 mechanical stress. Simulated bilayers of phospholipids containing docosahexaenoic acid were less sensitive to the applied surface tension than were saturated phospholipids, possibly implying a decrease in membrane elasticity (area elastic modulus, bending rigidity). The above features
10 distinguish DHA-containing lipids from saturated or nonunsaturated lipids and may be important for their biological

1.5 In Summary

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modes of action.

The functions of the phospholipids are multiple and 15 different for each phospholipid:

a. Sphingosine and carbohydrate containing lipids are mainly concentrated in nervous tissues.

b. The hydrophilic and hydrophobic parts of the phospholipid molecule allow them to function as emulsifying agents in order to maintain the proper colloidal state of protoplasm.

c. Phospholipids aid the transport of triglycerides through the liver, especially during mobilization from adipose tissue.

d. Phospholipids and their metabolites play an important role in intracellular signalling, for example via

25 phosphatidylinositol specific phospholipase C, phospholipase D or phosphatidylinositol-kinases.

e. Through their concentration in cell membranes they may somehow be involved in the transport of hydrophobic constituents into and out of cells.

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f. Phospholipids affect brain function in two substantial ways: (Cohen B.M., Babb S.M., Yurgelun-Todd D., et al. Brain choline uptake and cognitive function in middle age. Biol. Psych. 1997;41:905.)

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- a. The membranes of brain cells depend on phospholipids as part of their structure.
 Phosphatidylserine (PS) is concentrated in the cell membranes of the brain.
- b. Phospholipids are required for the production of neurotransmitters.
- c. Choline is a component of the neurotransmitter acetylcholine. Without adequate levels of acetylcholine, the brain can't store or retrieve information efficiently.
- d. Lower choline levels in the brain are an underlying factor for age-related cognitive disorders.
- e. Patients submitted to increased choline uptake show significant improvement in their ability to recall information and perform on memory retention tests, suggesting a causal relationship between poor choline status and cognition.
- g. Phosphatidylserine (PS) in Dementia-Related Diseases: 25
 - a. Dementia is the deterioration of mental function, particularly affecting memory, concentration, and judgment.
 - b. A frequent cause of dementia is Alzheimer's disease.
 - c. The first double-blind trial of PS for Alzheimer's disease was published about a decade ago. (Delwaide P.J., et al. Double-blind randomized controlled study of phosphatidylserine in demented subjects. Acta Neur. Scand. 1986;73:136-140.) In this study, 35 Alzheimer's patients were either given a placebo or 300 mg. per day of PS for six weeks. The PS group showed significant improvement after this short-term supplementation period.
 - d. More recently, a large double-blind study of 494 elderly patients with symptoms of cognitive decline compared a placebo to 300 mg. per day of PS for six months. (Cenacchi T., Bertoldin T., Farina C., et al. Cognitive decline in the elderly: A double-blind, placebo-controlled multicenter study on efficacy of phosphatidylserine administration. Aging Clin.

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Exp. Res. 1993;5:123-133.) Memory and learning of the PS-treated group was significantly improved over the placebo group, as well as certain emotional and behavior components of Alzheimer's disease.

e. Supplements of PS have also shown impressive results in older populations with memory impairment unrelated to Alzheimer's disease. (Crook T.H., et al. Effects of phosphatidylserine in age-associated memory impairment. Neurology 1991;41:644-649.) Three months of taking 300 mg. of PS daily, in one study, reversed the decline of memory function in a group of 149 patients. The memory function of these men and women initially averaged that of a typical 64 years old. After taking PS supplements, the average memory function was 52 years old -- a mental gain of 12 years.

20 h. Restoring and Preserving Liver Function:

- a. While the phospholipid PS dominates in the mental function arena, the phospholipid phosphatidylcholine (PC) is the major player for liver health.
- b. PC protects the liver against damage from alcoholism, pharmaceuticals, pollutant substances, viruses, and other toxic influences, most of which operate by damaging cell membranes.
- Many of the studies using PC supplements to aid c. recovery of the liver are based on 800 mg. per day (taken with meals). (Kidd P.M. Phosphatidylcholine: A superior protectant against liver damage. Alt. Med. Rev. 1996; 1:258-274.) Although PC is a source of choline, studies reviewed by Dr. Kidd suggest that PC is superior to choline; in fact choline in its pure form may be detrimental to the liver's recovery from toxic overload (such as in alcoholism). As a lipotropic, choline transports fats within the body, while inadequate choline intake might result in an unhealthy accumulation of fat in the liver. (Newberne P.M., Nauss K.M., and de Camargo J.L. Lipotropes, immunocompetence, and cancer. Cancer Res. 1983;43:24265-24345.)

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2. Flavonoids

Flavonoids are polyphenolic compounds ubigitous in nature. They are categorized into isoflavones, anthocyanidins, flavans, flavonols, flavones, citrus flavonoids, hesperidin,

- 5 chalcones, catechins, rutin, and flavanones. Essential flavonoids, such as quercetin in onions and genistein in soy are actually considered subcategories rather than independent categories. Over 4,000 flavonoids have been identified in fruits, vegetables and beverages (tea, coffee, beer, wine and
- 10 fruit drinks). Even though they have a similar molecular structure between them, their functions are different from each other. Flavonoids have been shown to have antibacterial, antiinflammatory, antiallergic, antimutagenic, antiviral, antineoplastic, anti-thrombotic, and vasodilatory activity.
- 15 Quercetin has been proven to block the "sorbitol pathway" which is directly associated with diabetes as well as to prevent LDLcholesterol oxidative damage, which is essential for the maintenance of a healthy cardiovascular system.
- Flavonoids are found in a wide range of fruits and vegetables. For example, Quercetin (a flavonol in vegetables, fruit and onions), Xanthohumol (a prenylated chalcone in beer), Isoxanthohumol (a prenylated flavanone in beer), Genistein (an isoflavone in soy), Chalconaringenin (a non-prenylated chalcone in citrus fruits) and Naringenin (a non-prenylated flavanone in 25 citrus fruits).

In plants flavonoids have very well defined functions. First, the accumulation of pigment in flower petals, seeds and leafs. Flowers, as pollinators, must attract pollen carriers. Second, they protect plants from UV damage, by absorbing UV at the epidermal layer. Third, they protect the plants against insects and pathogens.

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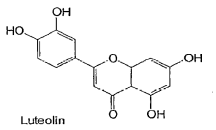
The flavonoid biosynthetic pathway is one of the best understood plant secondary metabolism pathways (1992, Gerats). The key enzymes are phenylalanine-ammonia lyase and chalcone synthase. Phenylalanine-ammonia lyase converts phenylalanine into cinnamic acid as it controls the total flow of carbons 5 into phenolics which is shown to be the limiting step in this pathway (1974, Creasy). Another key enzyme of the flavonoid pathway is the chalcone synthase. It condenses three molecules of malonyl-CoA with one molecule p-courmaroyl-CoA to form a C_{15} intermediate, naringenin chalcone, with a R stereochemistry at 10 the 2nd carbon. Chalcone isomerase, transforms the intermediate into the first flavonoid of the pathway, 2S-flavonone (naringenin). This reaction is part of all major flavonoid biosynthesis pathways. Chalcone synthase and chalcone isomerase form a complex ensuring the right stereochemistry 15 (1996, Lyster).

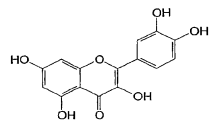
The structural components of flavonoids include two benzene rings on either side of a 3-carbon ring. Different combinations of hydroxyl groups, sugars, oxygens, and methyl groups attached to these structures create the various

- 20 groups attached to these structures create the various categories of flavonoids mentioned above. The capacity of flavonoids to act as an antioxidant depends upon their biochemical structure, and more specifically, the position of the hydroxyl groups. Epicatechin gallate, epigallocatechin 25 gallate, luteolin and quercetin exhibit the highest antioxidant
- activity, followed by epigallocatechin, gallic acid, epicatechin, catechin, rutin, and dihydroquercetin. It is worth noting at this point that the only difference between quercetin or luteolin (the most potent) and dihydroquercetin
- 30 (the least potent) is the double bond between the second (2nd) and third (3rd) carbons on the center (C) ring. The presence of this double bond significantly increases the antioxidant activity of the flavonoid. Antioxidant activity can be

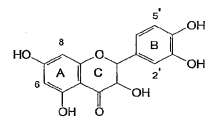
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increased with the addition of another hydroxyl group on the B or C ring.





Quecertin



Dihydroquercetin

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The potent antioxidant activity of flavonoids seems to be the most important function of flavonoids, responsible for many of the above mentioned health benefits.

The flavonoids most recognised by scientists until today are:

20 Quercetin and quercetin chalcone

Quercetin chalcone, is quercetin with an opened C ring and the oxygen found in the C-ring of quercetin converted into a hydroxyl group. Quercetin is mainly found in tea and even more in green tea.

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Oligomeric Proanthocyanidins

Oligomeric proanthocyanindins are oligomeric flavonoids, usually dimers and trimers, based on the flavan- 3ol, or catechin, molecule, sometimes attached to gallic acid. They are found in the bark of pine trees, in grape seeds and skins, in peanut skins, cranberries, tea, and other sources.

Ginkgo Biloba Extract

Ginkgo biloba extracts contain 24% ginkgo flavone glycosides and 6% terpenes. They are extracted from the eldest 10 living tree species, Gingo Biloba. Scientific research suggests that the beneficial constituents of gingo biloba extracts are quercetin and myricetin.

Luteolin

Luteolin is a flavonoid found in the same foods as 15 apigenin (vegetables and fruits). Scientific research has shown that luteolin and quercetin can inhibit platelet activating factor and suppress the inflammatory response induced by allergens.

Flavonoids have been studied for the last 60 years. Their antioxidant activity is accepted as a scientific fact. Epidemiological, clinical, and laboratory research on flavonoids demonstrates the use of flavonoids in the prevention and/or treatment of cardiovascular disease, cancer, inflammatory conditions, asthma, peridontal disease, liver

25 disease, cataracts and macular degeneration. Until today there has never been a flavonoid extracted from anything other than a plant, vegetable, fruit or algae.

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3. Preparation of Extracts

The phospholipid extract of the present invention may be extracted from a variety of marine or aquatic biomass sources. Preferred sources of the phospholipid composition are crustaceans, in particular, zooplankton. A particularly preferred zooplankton is Krill. Krill can be found in any

- marine environment around the world. For example, the Antarctic Ocean (where the krill is <u>Euphasia</u> superba), the Pacific Ocean (where the krill is <u>Euphasia</u> pacifica), the
- 10 Atlantic Ocean and the Indian Ocean all contain krill habitats. In particular, the coastal regions of Mauritius Island and/or Reunion Island off Madagascar, the Canadian West Coast, the Japanese Coast, the Gulf of St. Lawrence and the Bay of Fundy are krill habitats.

The phospholipid extract of the present invention is 15 preferably a product of initial processing of the biomass. As such, the phospholipids are extracted from the biomass grease as opposed to the oil, the oil being a product of subsequent processing steps of a biomass. Since the phospholipid extract is derived from the biomass grease, the viscosity of the 20 phospholipid extract tends to be higher than extracts from biomass oils. The extract has a very high natural stability with a peroxide value of zero or approaching zero and a good Oil Stability Index of less than about 0.2 Meg/kg after 20 or more hours. Table 1 below details the stability of the 25 extract.

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TABLE 1

Stability indexes of the extract after 50 hours at 97.8°C<0.1</th>Peroxide value (mEq/kg)<0.1</td>Oil Stability Index (after 50 hours) at 97.8°C (mEq/kg)<0.1</td>Saponification Index70-180Iodine Index (%)60-130

Phospholipids are generally present in the extract in an amount of at least 40% w/w, preferably at least 45% w/w. More preferably, the amount of phospholipid is from about 45-60% w/w. A variety of types of phospholipids may be present in the extract. These include phosphatidylethanolamine, phosphatidylinositol, phosphatidylserine, phosphatidylcholine and sphingomyelin.

The phospholipid extract preferably further comprises a number of other components. The extract may also comprise fatty acids, antioxidants and/or metals.

Fatty acids found in the phospholipid extract may be saturated, monounsaturated or polyunsaturated fatty acids. Polyunsaturated fatty acids are particularly preferred, the omega-3 and omega-6 fatty acids being most preferred. In particular, docosahexaenoic acid (DHA), eicosapentaenoic acid (EPA), myristic acid, myristoleic acid, lignoceric acid, linolenic acid, alpha linolenic acid, nervonic acid, linoleic acid, oleic acid, stearic acid, palmitic acid and palmitoleic acid are present in significant quantities. Arachidonic acid content of the extract is generally very low to non-existent despite the presence of phosphatidyl inositol and phosphatidyl serine. Other lipid components that may be present in the

25 extract include monoglycerides, triglycerides and/or cholesterol.

Table 2 below details the fatty acid compositions of the phospholipids of the extract.

TABLE 2

5 The fatty acid composition of the extract of the phospholipids

	Total PL	PC	PE
Fatty Acids	FA%	FA%	FA%
C14:0 MYRISTIC	2.04	1.70	0.7
C14:1 MYRISTOLEIC	1.22		
C15:0 PENTADECANOIC	0.2	0.30	0.3
C16:0 PALMITIC	24.08	26.50	23.9
C16:1 PALMITOLEIC	2.24	2.30	0.7
C18:0 STEARIC	1.02	1.30	2.9
C18:1 OLEIC	9.18	11.90	24.1
C18:2n6 LINOLEIC	1.63	2.30	0.8
C18:3n6 GLA	1.02	0.30	
C18:3n3 ALA	1.02	1.30	
C18:4n3 OTA	1.84	2.00	0.3
C20:0 ARACHIDIC			
C20:1 cis-11-EICOSENOIC	0.41	.0.60	0.7
C20:2n6 EICOSADIENOIC			
C20:3n6 METHYL ETA		0.20	
C20:4n6 ARACHIDONIC	0.61	0.70	0.6
C20:3n3 Homo-y-LINOLENIC			
C20:4n3			
C20:5n3 EPA	27.35	31.90	12.9
C22:0 BEHENIC			
C22:1 ERUCIC	1.22	1.50	
C22:2n6			
C22:4n6			
C22:5n6 METHYL DPA			
C22:5n3 DPA		1.00	
C22:6n3 DHA	24.9	14.20	32.1
C24:0 LIGNOCERIC			
C24:1 NERVONIC			
Total	100.0	100	100

Compared to phospholipids existing in the market today, the extract phospholipids:

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- a. achieve a superior profile;
- b. have the highest quantities of polyunsaturated fatty acids;
- c. have the highest quantities of DHA;
- d. are the only phospholipids that contain EPA; and
- e. are the only phospholipids that contain a

combination of EPA and DHA on the same molecule.

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PL = phospholipid FA = fatty acid PC = phosphatidylcholine PE = phosphatidylethanolamine

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Free fatty acids are present in the extract in an amount of at least 4% w/w and preferably at least 5% w/w. Polyunsaturated fatty acids, in particular omega-3 fatty acids, preferably make up at least 15% w/w, more preferably at least 40% w/w, and even more preferably at least 45% w/w, of the total lipids in the extract. DHA and EPA are generally the largest component of the fatty acids and preferably account for at least 32% w/w, more preferably at least 35% or 37%, of the total lipid content of the extract.

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Table 3 below details the fatty acid composition of the total lipids of the extract.

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TABLE 3

Fatty acid composition of total lipids of the extract

Sample	%
Fatty Acid Composition	
C14:0	<u>></u> 3.00
C14:1	<u>></u> 0.01
C15:0	<u>></u> 0.3
C16:0	>20.00
C16:1	>3.25
C18:0	<u>></u> 1.00
C18:1	>10.00
C18:2n6	>2.00
C18:3n6 GLA	>0.04
C18:3n3 ALA	<u>></u> 0.01
C18:4n3	<u>></u> 1.50
C20:0	<u>></u> 0.05
C20:1	<u>></u> 1.00
C20:2n6	>0.05
C20:3n6	<u>></u> 0.05
C20:4n6	<u><</u> 0.50
C20:3n3	<u>></u> 0.01
C20:4n3	<u>></u> 0.20
C20:5n3 EPA	<u>></u> 25.00
C22:0	<u>></u> 0.01
C22:1	<u>></u> 1.50
C22:2n6	<u>></u> 0.03
C22:4n6	<u>></u> 0.01
C22:5n6	<u>></u> 0.01
C22:5n3 DPA	<u>></u> 0.50
C22:6n3 DHA	<u>></u> 10.00
C24:0	<u>></u> 0.01
C24:1	>0.05

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Table 4 below also details the fatty acid composition of the total lipids of the extract.

TABLE 4

Fatty acid composition of total lipids of the extract

Saturated (g/10	0g lipid)	>22.00
Monounsaturated	(g/100g lipid)	>11.00
Polyunsaturated	(g/100g lipid)	<u>></u> 35.00
Omega-3 (g/100g	lipid)	>30.00
Omega-6 (g/100g	lipid)	≥1.00

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Antioxidants present in the extract may include vitamin A (for example, all-trans retinol), vitamin E (for example, alpha-tocopherol), beta-carotene, astaxanthin (mainly esterified but non-esterified may be present), canthaxanthin

5 and/or flavonoids. Antioxidants are preferably present in the extract in an amount of at least 20 and preferably at least 200 mg/100 ml.

Table 5 below details the lipids and other compounds (non-metal) of the extract.

TABLE 5

Lipid composition, vitamins A and E, pigments and flavonoids of the extract

Monoglycerides (MG) (g/100g sample)	<u>>0.7</u>
Triglycerides (TG) (g/100g sample)	<u>></u> 3.00
Free Fatty Acids (FFA) (g/100g sample)	<u>></u> 5.00
Cholesterol (g/100g sample)	<u><</u> 2.00
Total Phospholipids (PL) (g/100g sample)	<u>></u> 40.00
Phosphatidyl Ethanolamine (PE) (g/100g sample)	<u>></u> 2.50
Phosphatidyl Inositol (PI) (g/100g sample)	<u>></u> 0.20
Phosphatidyl Serine (PS) (g/100g sample)	<u>></u> 0.20
Phosphatidyl Choline (PC) (g/100g sample)	<u>></u> 35.00
Sphingomyelin (g/100g sample)	<u>></u> 0.50
	>1,400
Vitamin A (μ g/100 ml)	>15
Vitamin E (μ g/100 ml)	>1,600
Beta-Carotene (µg/100 ml)	<u> </u>
Astaxanthin (mg/100 ml)	≥10 >10
Canthaxanthin (mg/100ml)	>7.0
Flavonoid (mg/100ml)	

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The metals present in the extract are preferably zinc and selenium. Zinc is preferably present in an amount of at least 0.05 mg/100g of extract while selenium is generally present in an amount of less than 3 mg/100g of extract.

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Table 6 below details the metals content of the extract.

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TABLE 6

Metal composition and solvent residue of the extract mixture

>0.1
<2
<25 ppm

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Table 7 below details the physiochemical characteristics of the extract.

TABLE 7

10 Physiochemical characteristics of the extract

Color	Red	
Viscosity (cPs)	<1300	
Odor	Fish	

Extraction of the phospholipid composition from the biomass is generally carried out by a method similar to the one 15 described in commonly owned PCT publication number WO 00/23546, published on April 27, 2000, the disclosure of which is incorporated herein by reference. The extraction is generally carried out by successive acetone and alcohol treatments. For the extraction of the instant application, the preferred

- 20 treatment involves the use of >60% acetone in the first extraction followed by extraction with a mixture of organic solvents at 65-95%/45-50% preferably acetone, ethyl acetate/ethanol mixture. The most preferred extraction solvent system is 100% acetone in the first extraction followed with a
- 25 95%/5% ethyl acetate/ethanol mixture. However, other ketones can also be used in combination with or in place of acetone. The alcohol can be other than ethanol, e.g., isopropanol or tbutanol. The acetate may also vary. Further, the ratio of alcohol to acetate may vary widely from 100:0 to 0:100. The
- 30 procedure produces two successive lipid fractions and a dry residue enriched in protein, including active enzymes.

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Preferably, freshly harvested and finely divided marine and aquatic animal material is subjected to acetone extraction, for at least about two hours and preferably overnight. However, extraction time is not critical to the yield of lipid extracted. Particle sizes of comminuted crustacean less than 5 mm are preferred. The extraction is preferably conducted under an inert atmosphere and at a temperature of about 5 degrees Celsius or less. The mixture may be agitated during extraction and a volume ratio of about 6:1 of acetone to biomass is generally most preferred.

The solubilized lipid fraction is separated from the solid starting material by known techniques, for example, by filtration, centrifugation or sedimentation. Filtration is preferred. The residue is optionally washed with acetone to 15 recover more lipid and the acetone removed by flash evaporation or spray drying. Water residue is allowed to separate from the lipid extract at low temperature.

The solid residue left on the filter from the initial extraction is suspended and extracted with 95/5 ethyl 20 acetate/ethanol, preferably two volumes (original volume of material). The filtrate is evaporated yielding a second fraction of lipids. Extraction period is not critical although it is preferred to extract for about 30 minutes at a temperature below about 5 degrees Celsius.

Each phospholipid is subdivided into multiple categories depending on the fatty acids that are attached to the molecule. The biological activity, bioavailability as well as the value of phospholipids is determined by the purity and the source:

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a. Purity:

i. Optimal purity of the phospholipid or flavonoid of the invention is at least 99% by weight. The purity of the phospholipid or flavonoid after extraction from

- 5 the krill may vary, but will normally be in the range of at least 90% to 100% of the/or mixture of phospholipid compound(s). Usually, the purity will be at least 95%. Preferably, the purity will be at least 96%, 97% or 98%. More preferably, the purity will be at least 99.5%. Most
- 10 preferably, the purity will be at least 99.9%. By "purity" is meant that the phospholipid or flavonoid of the invention is isolated from other phospholipids, flavonoids, or components of the extract, to the weight percent specified. Isolation may be performed by e.g. HPLC. For example, a phospholipid that is
- 15 99% pure, contains less than 1% by weight of any material other than the specified phospholipid.

ii. Higher bioavailibility and efficacy is achieved with higher purity.

iii. Phospholipid market value is directlyanalogous to the purity achieved for the final product.

b. Source and fatty acid content:

i. The types of fatty acids attached to the phospholipid is widely dependent upon the source.

ii. Plant source phospholipids contain mainly 25 palmitic acid (16:0), stearic acid (18:0), vaccenic acid (18:1), linoleic acid (18:2) or alpha-linoleic acid (18:3).

iii. Animal source phospholipids contain a higher percentage of longer-chain fatty acids with higher degree of unsaturation like homo-gamma-linoleic acid (20:3),

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arachidonic acid (20:4), behenic acid (22:0) and docosahexanoic acid -DHA (22:6).

iv. Neptune Krill Oil[™] (the present invention) phospholipids contain high quantities of eicosapentanoic acid -EPA (20:5) and docosahexanoic acid -DHA (22:6). Their fatty acid profile closely resembles that of human brain phospholipids.

v. The efficacy in human health and the value of phospholipids increases directly analogous to the length of the fatty acid chain and the degree of unsaturation. Therefore, phospholipids with more polyunsaturated fatty acids attached to them are more efficacious and of higher value.

vi. Arachidonic acid, although polyunsaturated, has been proven to predispose to inflammatory disease. Hence, 15 moderate quantities are preferred.

vii. DHA and EPA are the two most active polyunsaturated fatty acids in the human body, contributing to all health benefits associated with omega-3 fatty acids.

viii. The highest quantities of polyunsaturated 20 fatty acids contained in the phospholipids in the market today are:

	а	Arachidonic acid		:30.1%
		Homo-gamma-linolenic a	acid	:9.0%
		nono ganana renoremente		:8.4%
c.	DHA			

25 4. Pharmaceutical, nutraceutical and cosmetic compositions

The phospholipid extract of the present invention may be used with or without other additives. Preferably, no other additives are used. However, if other additives are used, pharmaceutical or nutraceutical formulations may be made by

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methods known in the art. For example, the compositions of the present invention may be formulated in a conventional manner using one or more pharmaceutically or nutraceutically acceptable carriers. Thus, the extract may be formulated for

- 5 oral administration. For oral administration, the pharmaceutical or nutraceutical compositions may take the form of, for example, tablets or capsules prepared by conventional means with pharmaceutically or nutraceutically acceptable excipients such as binding agents (<u>e.g.</u>, pregelatinised maize
- 10 starch, polyvinylpyrrolidone or hydroxypropyl methylcellulose); filters (e.g., lactose, microcrystalline cellulose or calcium phosphate); lubricants (e.g., magnesium stearate, talc or silica); disintegrants (e.g., potato starch or sodium starch glycollate); or wetting agents (e.g., sodium lauryl sulphate).
- 15 The tablets may be coated by methods well known in the art. Liquid preparations for oral administration may take the form of, for example, solutions, syrups or suspensions, or they may be presented as a dry product for constitution with water or other suitable vehicle before use. Such liquid preparations
- 20 may be prepared by conventional means with pharmaceutically or nutraceutically acceptable additives such as suspending agents (<u>e.g.</u>, sorbitol syrup, methyl cellulose or hydrogenated edible fats); emulsifying agents (<u>e.g.</u>, lecithin or acacia); nonaqueous vehicles (<u>e.g.</u>, almond oil, oily esters or ethyl
- 25 alcohol); and preservatives (e.g., methyl or propyl phydroxybenzoates or sorbic acid).

When the phospholipid extract of the inventions is used as a nutraceutical, it can be in the form of foods, beverages, energy bars, sports drinks, supplements or other forms all as are known in the art.

As noted above, the phospholipid extract of the invention is also useful in cosmetic preparations, e.g.,

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moisturizing creams, sun-block products and other topical cosmetic products as known in the art.

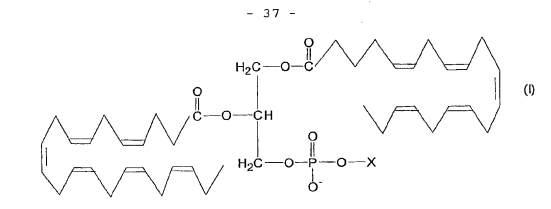
The phospholipid extract of the present invention may be used in the treatment or prevention of a variety of disease states including: liver disease; chronic hepatitis; steatosis; liver fibrosis; alcoholism; malnutrition; chronic parenteral nutrition; phospholipid deficiency; lipid peroxidation; disarrhythmia of cell regeneration; destabilization of cell membranes; coronary artery disease caused by

10 hypercholesterolemia; high blood pressure; menopausal or postmenopausal conditions; cancer, e.g., skin cancer; hypertension; aging; benign prostatic hyperplasia; kidney disease; edema; skin diseases; gastrointestinal diseases; peripheral vascular system diseases (e.g. leg ulcers); pregnancy toxemia; and

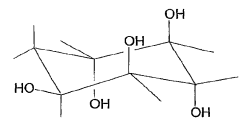
15 neurodegenerative and psychiatric diseases (e.g. Parkinson's, Alzheimer's, autism, attention deficit disorder, learning disorders, mood disorders, bipolar depression, multiple sclerosis, muscular dystrophy).

The extracts are also useful for targeting tumors and 20 can be used in conjunction with radioisotopes for diagnosing central nervous system tumors. The extract can also be used to reduce local fat deposits and reducing visible cellulite. The extract can also be used in aesthetics such as breast enlargement by acting on the lobular tissue of the breast and 25 by increasing hydration of the breast.

As noted above, the present invention provides novel phospholipids derived from a marine or aquatic biomass. The novel phospholipids have the general formula (I):



wherein X represents a moiety normally found in phospholipids, e.g., $-CH_2CH_2N(CH_3)_3$, $CH_2CH_2NH_3$ or



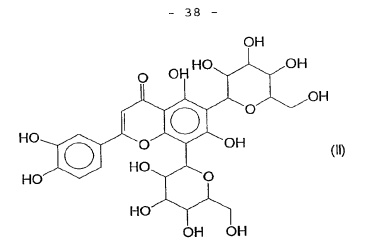
for phophatidylcholine, phosphatidylethanolamine or phosphatidylinositol, respectively.

The left hand acid residue is derived from 15 docosahexanoic acid (DHA) [C22:6n3]. The right hand acid residue is derived from eicosapentaenoic acid (EPA) [C20:5n3].

These novel phospholipids have all of the uses noted above for phospholipids in pharmaceutical, nutraceutical and cosmetic compositions.

20 As noted above, the present invention also provides a novel flavonoid compound derived from a marine or aquatic biomass. The novel flavonoid compound has the formula (II):

10



The novel flavonoid is an antioxidant and thus is useful in the pharmaceutical, nutraceutical and cosmetic compositions of the invention.

As used herein and in the claims, where the term 10 "about" is used with a numerical value, the numerical value may vary by at least ±50%. Preferably, the variation will be ±40% or ±30% and more preferably ±20% or ±10%. Even more preferred variations are in the range ±5%, ±4%, ±3% or ±2%. Most preferably, the variation is in the range of ±1%.

15 Brief Description of the Drawings

Figs. 1 to 3 are chromatograms of the product of Example 1.

Fig. 4 is a mass spectrograph for characterizing the novel flavonoid compound (II).

20 The invention is further illustrated by the following non-limiting examples.

The extraction of the phospholipids for Example 1 was as described above for krill extractions.

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Examples

Materials and Methods

For analysis of lipids, samples were dissolved in solvent and standards were added. Lipid classes were isolated 5 using silica gel and quantified. Fatty acid composition of total lipids and individual phospholipids was determined by gas chromatography. Pigments were measured by reversed phase high performance liquid chromatography.

Example 1

10 This example illustrates the isolation and molecular characterization of the phospholipids from the extract.

Sample #804 molecular species determination

The sample contains large amounts of phospholipids,

- mainly:
- 15 PC (438.48 mg/g lipid) PE (183.15 mg/g lipid)

Preliminary results were obtained only for these two phospholipid fractions.

METHODS

20 Separation of main phospholipid fractions

To obtain large quantities of PC and PE, separation was done by Thin Layer Chromatography (TLC) and bands identity was confirmed by HPLC.

Diacylglycerol formation

25 Both fractions (PC and PE) were incubated with phospholipase C, the enzyme which removes choline phosphate

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from PC and ethanolamine phosphate from PE. The remaining diacylglycerols were extracted with ethyl ether.

Benzoate derivatization

Each mixture of diacylglycerols needed to be derivatized (using benzoic anhydride and 4-dimethylaminopyridine) to make further separation possible. In a parallel experiment, derivatization was done for three standard authentic diacylglycerols, dilinolein, diolein and dipalmitin.

Subclass separation

10 A preliminary separation of diacylglycerols derivatives into subclasses was done by TLC. Diacylglycerol derivatives obtained from PC and from PE separated into two major bands (#3 and #4). Additional bands #2 were also visible very close to the start. Only bands #3 and #4 were processed 15 further because their localization corresponded to the localization of main band #2 obtained for a mixture of standards (benzoate derivatives of dilinolein, diolein and dipalmitin).

and a second se	#4(R£#⊨70.*37%	∛ #4.≥(RE =: 0.37)
#2		//#3: (R£ '=*0.25)
Start Std mix	#2 Start PC	Start PE

Example TLC plate separation

20

HPLC fractionation

10

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Bands #3 and #4 obtained for PC and PE were eluted and further separated into individual diacylglyercol species by HPLC. To confirm a number of peaks for the subsequent GC analysis, each peak was collected and separately re-run on HPLC.

Number of confirmed peaks:

For PC band #3, nine peaks were identified and confirmed. For PC band #4, nine peaks were identified and confirmed. For PE band #3, eight peaks were identified and confirmed. For PE band #4, eight peaks were identified and confirmed.

See Figure 1.

Hydrolysis, methyl ester derivatization and GC analysis

For both PC and PE, all confirmed peaks obtained from HPLC separation of band #3 were hydrolized and fatty acid 15 profiles were determined by GC after conversion into methyl esters. Peak identity was assessed by mass spectrometry. Fatty acid profiles were compared to those obtained for intact PC and PE fractions subjected to hydrolysis and methylation.

Results

20 The peak surface areas calculated for fatty acid molecular species in selected fractions are summarized in Table 8. The peak fatty acid areas for intact PC and PE fraction are in Table 9. The representative Gas Chromatography profiles for an individual fraction and for intact phospholipid (PC) are 25 presented in Table 10.

The Gas Chromatography profiles obtained for individual peaks were only partly consistent with profiles obtained for intact PC. They contained only 5-6 major peaks while Gas Chromatography profiles of intact phospholipids

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consist of much higher number of peaks. Among the 5-6 peaks consistently found in molecular species profiles, only two had identity confirmed by mass spectrometry (Cl6:0 and Cl8:0). Among the remaining three peaks, one did not correspond to any fatty acid and two had retention times identical to those of

authentic omega-3 fatty acids, EPA and DHA.

The Cl6:0 peak was prominent in all individual molecular species profiles and was also prominent in the intact phospholipid fractions. For the Cl8:0 peak, its proportions found in individual peaks were relatively high. Oleic acid (Cl8:1) was found at high levels in both PC and PE fatty acid profile.

TABLE 8

Molecular species peak areas obtained for selected fractions.

15

	<u> </u>	C18:0	EPA	RT 48.33	DHA
Fraction	C16:0		42.76	103.83	62.07
PC band #3 F1	205.27	57.79		71.96	7.11
PC band #3 F2	21.39	8.87	0		14.75
PC band #3 F3	58.74	17.70	0	45.64	
PC band #3 F4	93.41	9.72	0	44.31	9.19
	19.87	9.67	4.56	46.89	3.96
PC band #3 F5		10.34	12.45	59.86	14.29
PC band #3 F6	15.26		30.70	56.83	25.12
PC band #3 F7	28.32	10.93		84.24	11.89
PC band #3 F8	6.39	4.49	0	58.95	28.22
PC band #3 F9	14.65	8.21	8.60		9.19
PB band #3 F2	4.50	10.79	0	77.68	
PE band #3 F3	26.85	22.14	14.45	49.62	21.76
		22.45	28.70	62.11	29.43
PE band #3 F4	13.08	20.34	11.06	100.79	30.61
PE band $#3 F5$	22.42			54,88	7.28
PE band #3 F6	3.05	6.13	4.93		

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TABLE 9

Selected fatty acid peak areas of intact PC and PE

	C16:0	C18:0	C18:1	EPA	Un- identified	DHA
Retention time	15.80	21.66	22.36+22.63	39.68	48.34	53.59
PC	1141.36	35.75	257.99	642.50	68.61	192.22
PE	166.43	20.45	87.75	59.77	110.27	109.63

5

See Fig. 2

TABLE 10

The representative GC profiles for an individual fraction and

10 for intact phospholipid (PE)

CH	PKNO	TIME	AREA	HEIGHT	MK	IDNO	CONC
1	5	0.826	17654310	1368301	E		21.8397
	13	2.637	11027760	1352920	Е		13.6422
	14	2.916	2167386	203115	Е		2.6812
	15	3.15	597812	87264	v		0.7395
	22	4.408	667991	60799	v		0.8264
	29	7.063	7293939	290768			9.0231
	30	8.397	144489	13997			0.1787
	32	9.933	32467398	1384059	Е		40.1646
	33	10.252	8166303	661493	v		10.1023
	43	14.451	348072	20030			0.4306
	44	14.813	102126	9975			0.1263
	45	15.12	198366	21561	·		0.2454
TOTAL			80835952	5474282			100

See Fig. 3

Example 2

15 UVB-Induced Skin Cancer

Objectives

To evaluate the photoprotective potential of krill extract against UVB-induced skin cancer.

Study Design

Randomized control trial

Statistical significance p<0.05

Study Phase

5 Pre-clinical

Experimental Animals

Type: Nude Mice

Strain: C57BL6 Nude Congenic Mice - B6NU -T (heterozygotes) (Preference of specific type because of proven susceptibility to skin cancer).

Study Protocol

Number of nude mice = 96

Randomization groups: 48 placebo: 16 per os

16 local application

16 per os and local application

15

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48 krill extract: 16 per os

16 local application

16 per os and local application

In order to establish efficacy of krill extract for

20 the prevention of skin cancer, the test was conducted as a randomized double blind controlled trial (both the pathologist and the research assistant were blind). Half of the mice were treated orally or topically or both with extract containing 100% by weight of krill extract and the other half underwent

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the same method of treatment with a placebo. The groups were divided as follows:

Nutrition: Week 1: fat-free chow

Week 2-20: according to group

5 Experimental Design:

20

The mice were divided in six groups as follows: Group A: fat-free chow with supplementation of soy extract (20% of total calories)

Group B: fat-free chow (100% of calories) + local application 10 of soy extract 2 times per day

Group C: fat-free chow with supplementation of soy extract (20% of total calories) + local application of soy extract 2 times per day

Group D: fat-free chow with supplementation of krill extract 15 (20% of total calories)

Group E: fat-free chow (100% of calories) + local application of krill extract 2 times per day

Group F: fat-free chow with supplementation of krill extract (20% of total calories) + local application of krill extract 2 times per day

Week 2-20: UVB radiation using a fluorescent test lamp, emission spectrum 270 - 400 nm.

Week 3-20: liquid from blisters formed is examined for PGE2 levels

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Week 3-20: mice are anaesthetized with ether and sacrificed when malignant tumours have formed or at the end of the 20 weeks.

Skin is examined by pathologist for signs of carcinogenesis.

The results are shown in the following Table 11.

TABLE	11

	Frequency of cancer	
	Krill Oil	Placebo
"mulication	Frequency %	Frequency %
Application	13	69.3
Oral	0	63.8
Topical	0	37.5
Oral & Topical		

10

5

In conclusion, the results of the present study demonstrate that both oral and topical krill extract maybe effectively used for the protection of skin against the harmful effects of UVB radiation including skin cancer.

15 Example 3

This example illustrates the use of the present krill extract in improving dyslexia and abnormal motor function in a 7 year old girl.

2g per day of the krill extract were given to a 7 20 year old girl suffering from dyslexia and abnormal motor function. After 1.5 months, she showed:

- Increased learning ability (blind observation by psychologist)
- Improved motor function (moderate ice skating)
- 25

(

Improved speech

Improved social skills

Accordingly, the krill extract has beneficial neurological properties.

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All publications cited in this specification are herein incorporated by reference as if each individual publication were specifically and individually indicated to be incorporated by reference. The citation of any publication is for its disclosure prior to the filing date and should not be construed as an admission that the present invention is not

entitled to antedate such publication by virtue of prior

Although the foregoing invention has been described 10 in some detail by way of illustration and example for purposes of clarity of understanding, it is readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the

15 appended claims.

invention.

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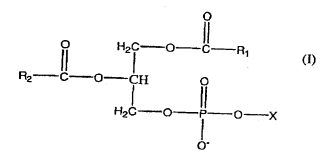
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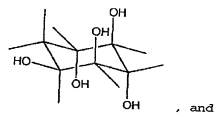
CLAIMS:

1. A composition, comprising :

(a) a phospholipid of the general formula (I),



wherein R_1 and R_2 is docosahexaenoic acid (DHA) or eicosapentaenoic acid (EPA), and X is $-CH_2CH_2NH_3$, $-CH_2CH_2N(CH_3)_3$ or



(b) an antioxidant,

wherein DHA and EPA are attached simultaneously on the phospholipid and wherein the carboxy moiety attached to R1 and R2 is the carbonyl group of said EPA and DHA.

2. The composition according to claim 1, further comprising a flavonoid of the general formula (II)

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3. The composition according to claim 1 or 2, wherein the composition is derived from at least one marine or aquatic biomass.

4. The composition according to claim 3, wherein the biomass is crustaceans.

5. The composition as according to claim 3 or 4, wherein the biomass is zooplankton.

6. The composition according to claim 5, wherein the zooplankton is krill.

7. The composition according to any one of claims 3 to 6, wherein the composition is derived from initial processing of the biomass.

8. The composition according to claim 7, wherein the processing is conducted at a temperature of about 5° C or less.

9. The composition according to claim 7 or 8, wherein the composition is derived from the biomass grease.

10. The composition according to any one of claims 1 to 9, comprising at least about 40% w/w phospholipid.

11. The composition according to claim 10, comprising at least about 45% w/w phospholipids.

12. The composition according to claim 11, comprising from about 45 to 60% w/w phospholipid.

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13. The composition according to any one of claims 1 to 12, wherein the composition comprises at least one phospholipid selected from the group consisting of phosphatidylcholine, phosphatidylethanolamine, phosphatidylinositol, phosphatidylserine and sphingomyelin.

14. The composition according to any one of claims 1 to 13, further comprising a phospholipid comprising saturated, monounsaturated or polyunsaturated fatty acids.

15. The composition according to claim 14, wherein the polyunsaturated fatty acids are selected from the group consisting of omega-3 and omega-6 fatty acids.

16. The composition according to claim 14, wherein the fatty acids are selected from the group consisting of docosahexaenoic acid (DHA), eicosapentaenoic acid (EPA), myristic acid, myristoleic acid, lignoceric acid, linolenic acid, alpha linolenic acid, nervonic acid, linoleic acid, oleic acid, stearic acid, palmitic acid and palmitoleic acid.

17. The composition according to any one of claims 1 to 16, further comprising lipids.

18. The composition according to claim 17, wherein the additional lipids are selected from at least one lipids from the group consisting of monoglycerides, triglycerides, and cholesterols or a mixture thereof.

19. The composition according to any one of claims 1 to 18, further comprising at least about 4% w/w of free fatty acids.

20. The composition according to claim 19, comprising at least about 5% w/w of free fatty acids.

21. The composition according to any one of claims 17 to 20, wherein polyunsaturated fatty acids comprise at least about 15% w/w of the total lipids in the composition.

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22. The composition according to claim 21, wherein polyunsaturated fatty acids comprise at least about 40% w/w of the total lipids in the composition.

23. The composition according to claim 22, wherein polyunsaturated fatty acids comprise at least about 45% w/w of the total lipids in the composition.

24. The composition according to any one of claims 21 to 23, wherein the polyunsaturated fatty acids are omega-3 fatty acids.

25. The composition according to any one of claims 17 to 24, wherein DHA and EPA comprise at least about 32% w/w of the total lipids in the composition.

26. The composition according to claim 25, wherein DHA and EPA comprise at least about 35% w/w of the total lipids in the composition.

27. The composition according to any one of claims 1 to 26, wherein the antioxidant is selected from at least one of the group consisting of vitamin A, vitamin E, beta-carotene, astaxanthin, canthaxanthin, flavonoid and mixture thereof.

28. The composition according to claim 27, wherein the vitamin A is all-trans retinol, the vitamin E is alpha-tocopherol, and the astaxanthin is mainly esterified.

29. The composition according to any one of claims 1 to 28, further comprising a metal.

30. The composition according to claim 29, wherein the metal is zinc, selenium or a mixture thereof.

31. The composition according to any one of claims 17 to 30, wherein the fatty acid composition of the lipids in the composition is about:

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	Total	Phosphatidy1-	
	Phospholipid	choline	ethanolamin
Fatty Acids	Fatty Acids	Fatty Acid %	Fatty Acid
C14:0 MYRISTIC	2	2	0.7
C14:1 MYRISTOLEIC	1		
C15:0 PENTADECANOIC	0.2	0.3	0.3
C16:0 PALMITIC	24	27	24
C16:1 PALMITOLEIC	2	2	0.7
C18:0 STEARIC	1	1	3
C18:1 OLEIC	9	12	24
C18:2n6 LINOLEIC	2	2	0.8
C18:3n6 GLA	1	0.3	· · · · · · · · · · · · · · · · · · ·
C18:3n3 ALA	1	1	
C18:4n3 OTA	2	2	0.3
C20:0 ARACHIDIC			
20:1 cis-11- COSENOIC	0.5	0.6	0.7
20:2n6 EICOSADIENOIC			
20:3n6 METHYL ETA		0.2	
20:4n6 ARACHIDONIC	0.6	0.7	0.6
20:3n3 Homo-γ- NOLENIC			

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C20:4n3			
C20:5n3 EPA	27	32	13
C22:0 BEHENIC			
C22:1 ERUCIC	1	1.5	
C22:2n6			
C22:4n6			
C22:5n6 METHYL DPA			
C22:5n3 DPA		1.0	
C22:6n3 DHA	25	14	32
C24:0 LIGNOCERIC			
C24:1 NERVONIC			
Total	100.0	100	100

32. The composition according to claim 31, wherein the total fatty acid composition of all the lipids in the composition is about:

Sample	ક
Fatty Acid Composition	
C14 : 0	≥3.00
C14 : 1	≥0.01
C15 : 0	≥0.3
C16 : 0	≥20.00

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C16 : 1	
C18 : 0	≥3.25
	≥1.00
C18 : 1	≥10.00
C18 : 2n6	≥2.00
C18 : 3n6 GLA	≥0.04
C18 : 3n3 ALA	≥0.01
C18 : 4n3	≥1.50
C20 : 0	≥0.05
C20 : 1	≥1.00
C20 : 2n6	≥0.05
C20 : 3n6	≥0.05
C20 : 4n6	≤0.50
C20 : 3n3	≥0.01
C20 : 4n3	≥0.20
C20 : 5n3 EPA	≥25.00
C22 : 0	≥0.01
C22 : 1	≥1.50
C22 : 2n6	≥0.03
C22 : 4n6	≥0.01
C22 : 5n6	≥0.01
C22 : 5n3 DPA	
C22 : 6n3 DHA	≥0.50
	≥10.00

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C24 : 0	≥0.01
C24 : 1	≥0.05

33. The composition according to claim 32, wherein the total fatty acid composition of all the lipids further is about:

Saturated (g/100g lipid)	≥22.00
Monounsaturated (g/100g lipid)	≥11.00
Polyunsaturated (g/100g lipid)	≥35.00
Omega-3 (g/100g lipid)	≥30.00
Omega-6 (g/100g lipid)	≥1.00

34. The composition according to any one of claims 1 to 32, further comprising about:

Monoglycerides (MG) (g/100g sample)	≥0.7
Triglycerides (TG) (g/100g sample)	≥3.00
Free Fatty Acids (FFA) (g/100g sample)	≥5.00
Cholesterol (g/100g sample)	≤2.00

Total Phospholipids (PL) (g/100g sample) ≥40.00

Phosphatidyl Ethanolamine (PE) (g/100g sample)	≥2.50
Phosphatidyl Inositol (PI) (g/100g sample)	≥0.20
Phosphatidyl Serine (PS) (g/100g sample)	≥0.20

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Phosphatidyl Choline (PC) (g/100g sample)	≥35.00
Sphingomyelin (g/100g sample)	≥0.50
Vitamin & (um (100	
Vitamin A (µg/100g ml)	≥1,400
Vitamin E (µg/100g sample)	≥15
Beta-Carotene (µg/100g ml)	≥1,600
Astaxanthin (g/100g ml)	≥10
Canthaxanthin (mg/100g ml)	
Flavonoid (mg/100 ml)	≥10
	≥7.0

35. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of liver disease.

36. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of liver disease.

37. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of chronic hepatitis.

38. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of chronic hepatitis.

39. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of steatosis.

40. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of steatosis.

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41. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of liver fibrosis.

42. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of liver fibrosis.

43. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of alcoholism.

44. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of alcoholism.

45. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of malnutrition.

46. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of malnutrition

47. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of chronic parenteral malnutrition.

48. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of chronic parenteral malnutrition.

49. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of phospholipids deficiency.

50. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of phospholipids deficiency.

51. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of lipid peroxidation.

52. Use of the composition according to any one of claims 1 to 33,

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for the prevention or treatment of lipid peroxidation.

53. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of disserrythmia of cell regeneration.

54. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of disserrythmia of cell regeneration.

55. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of destabilization of cell membranes.

56. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of destabilization of cell membranes.

57. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of coronary artery disease caused by hypercholesterolemia, high blood pressure, menopausal or post-menopausal conditions.

58. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of coronary artery disease caused by hypercholesterolemia, high blood pressure, menopausal or post-menopausal conditions.

59. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention of cancer.

60. Use of the composition according to any one of claims 1 to 33, for the prevention of cancer.

61. Use of claim 59 or 60, wherein cancer is skin cancer.

62. Use of claim 59 to 61, wherein the use of the composition is oral or topical.

63. Use of the composition according to any one of claims 1 to 33,

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for the manufacture of a medicament for the prevention or treatment of hypertension.

64. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of hypertension.

65. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of effects of ageing.

66. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of effects ageing.

67. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of benign prostatic hyperplasia.

68. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of benign prostatic hyperplasia.

69. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of kidney disease.

70. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of kidney disease.

71. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of edema.

72. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of edema.

73. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of skin diseases.

74. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of skin diseases.

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75. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of gastrointestinal diseases.

76. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of gastrointestinal diseases.

77. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of peripheral vascular system diseases.

78. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of peripheral vascular system disease.

79. Use of the composition according to claim 78, wherein the peripheral vascular system disease is ulcers.

80. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of pregnancy toxemia.

81. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of pregnancy toxemia.

82. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for the prevention or treatment of neurodegenerative or psychiatric disease.

83. Use of the composition according to any one of claims 1 to 33, for the prevention or treatment of neurodegenerative or psychiatric disease.

84. The use of claim 81 or 82, wherein the neurodegenerative or psychiatric disease is dementia, Parkinson's, Alzheimer's, autism, attention deficit disorder, learning disorders, mood disorders, behavioural disorders, multiple sclerosis or muscular dystrophy.

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85. Use of claim 84, wherein the learning disorder is dyslexia. 86. Use of the composition according to any one of claims 1 to 33,

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manufacture of a medicament for the prevention or treatment of abnormal motor function.

87. Use of the composition according to any one of claims 1 to 33. for the prevention or treatment of abnormal motor function.

88. Use of the composition according to any one of claims 1 to 33, in conjunction with a radioisotope for diagnosing or targeting tumors.

89. Use of the composition according to any one of claims 1 to 33, in conjunction with a radioisotope for the manufacture of a medicament for diagnosing or targeting tumors.

90. Use of claim 88 or 89, wherein the tumors are central nervous system tumors.

91. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for reducing local fat deposits and reducing visible cellulite.

92. Use of the composition according to any one of claims 1 to 33, for reducing local fat deposits and reducing visible cellulite.

93. Use of the composition according to any one of claims 1 to 33, for the manufacture of a medicament for aesthetic enhancement.

94. Use of the composition according to any one of claims 1 to 33, for aesthetic enhancement.

95. Use of the composition according to claim 94, wherein the aesthetic enhancement is breast enlargement.

96. A commercial package comprising the composition according to claim 1 to 33 together with instructions for use in treating cancer.

97. A commercial package comprising the composition according to claim 1 to 33 together with instructions for use in treating skin cancer.

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98. A commercial package comprising the composition according to claim 1 to 33 together with instructions for use in treating learning disorders.

99. A commercial package comprising the composition according to claim 1 to 33 together with instructions for use in treating dyslexia.

100. A commercial package comprising the composition according to claim 1 to 33 together with instructions for use in treating abnormal motor function.

101. Use of the composition according to any one of claims 2 to 33, for the manufacture of a medicament for the prevention or treatment of cardiovascular disease.

102. Use of the composition according to any one of claims 2 to 33, for the prevention or treatment of cardiovascular disease.

103. Use of the composition according to any one of claims 2 to 33, for the manufacture of a medicament for the prevention or treatment of inflammatory conditions.

104. Use of the composition according to any one of claims 2 to 33, for the prevention or treatment of inflammatory conditions.

105. Use of the composition according to any one of claims 2 to 33, for the manufacture of a medicament for the prevention or treatment of asthma.

106. Use of the composition according to any one of claims 2 to 33, for the prevention or treatment of asthma.

107. Use of the composition according to any one of claims 2 to 33, for the manufacture of a medicament for the prevention or treatment of peridontal disease.

108. Use of the composition according to any one of claims 2 to 33, for the prevention or treatment of peridontal disease.

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109. Use of the composition according to any one of claims 2 to 33, for the manufacture of a medicament for the prevention or treatment of cataracts.

110. Use of the composition according to any one of claims 2 to 33, for the prevention or treatment of cataracts.

111. Use of the composition according to any one of claims 2 to 33, for the manufacture of a medicament for the prevention or treatment of macular degeneration.

112. Use of the composition according to any one of claims 2 to 33, for the prevention or treatment of macular degeneration.

113. A composition substantially as described with reference to and as illustrated in the accompanying figures.

114. Use of a composition, said use being substantially as described with reference to and as illustrated in the accompanying figures.

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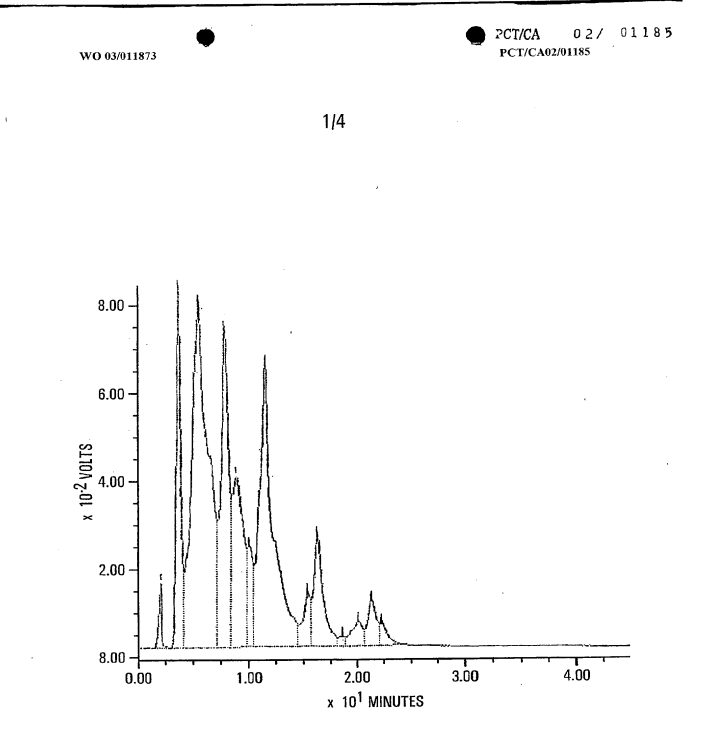
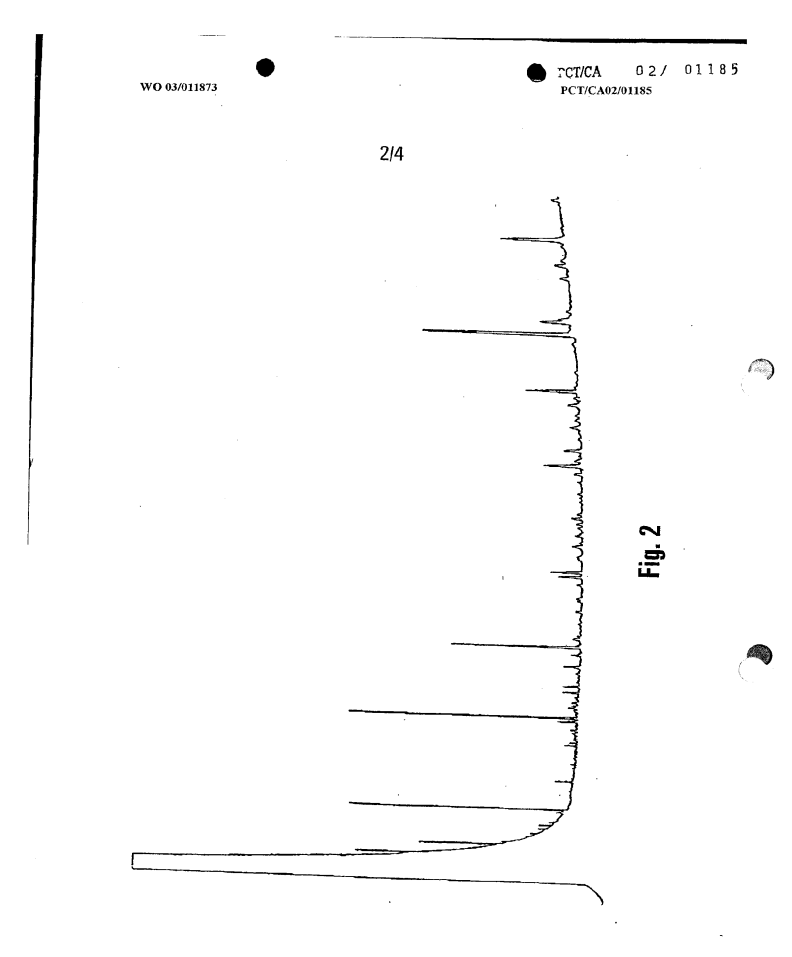
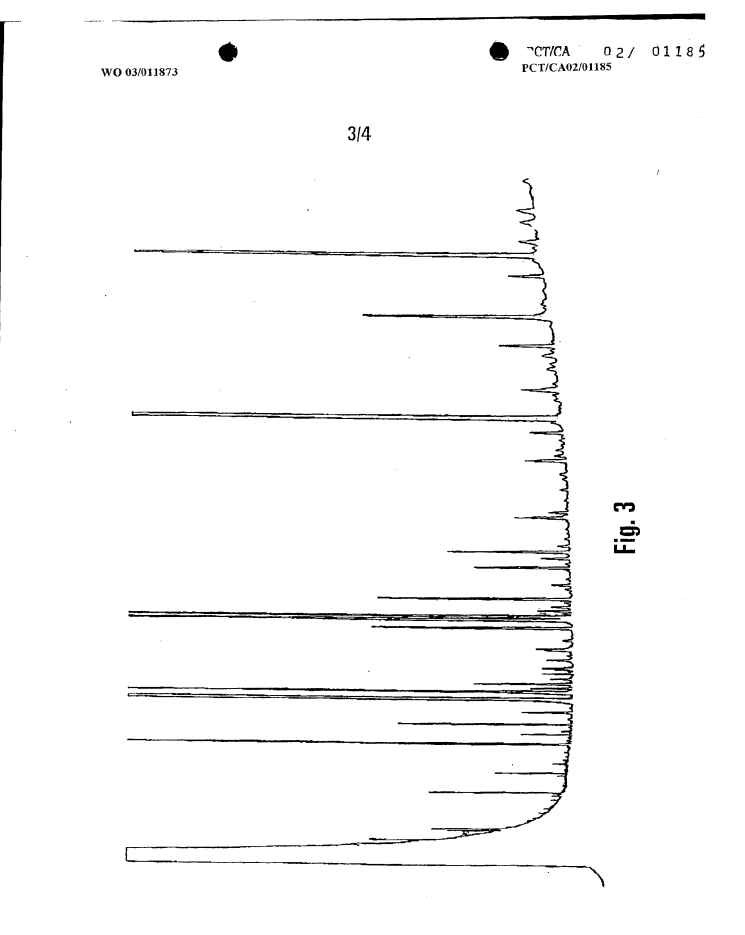


Fig. 1



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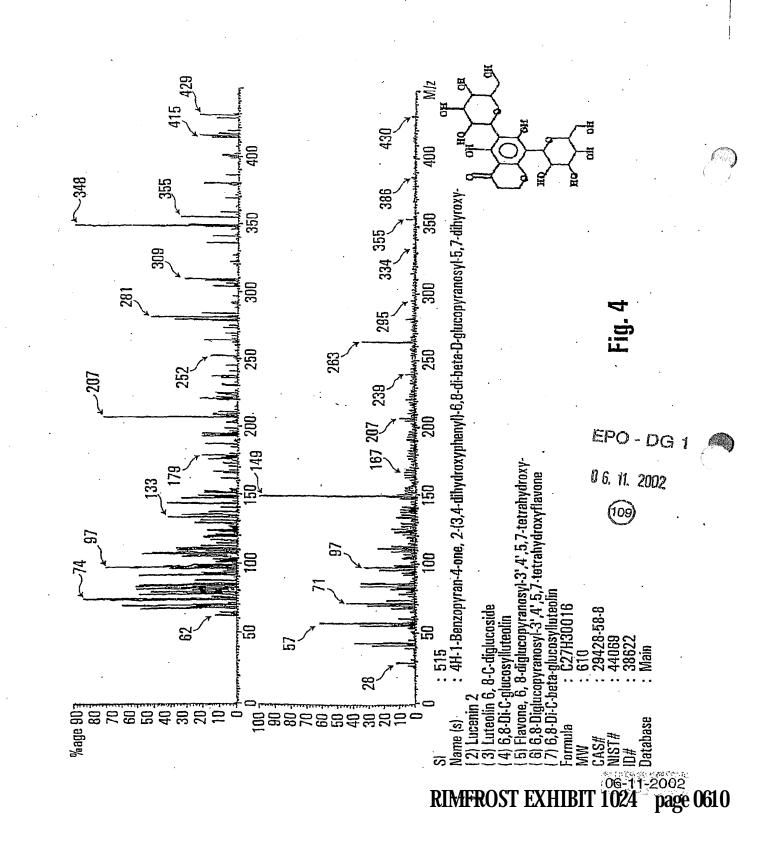
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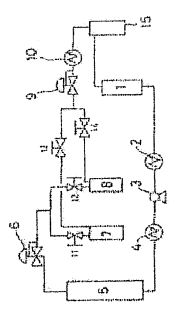
Bibliographic data: JP4057853 (A) - 1992-02-25

METHOD FOR EXTRACTING AND SEPARATING COLORING MATTER FROM KRILL

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Classification:	- international: <i>C09B61/00;</i> (IPC1-7): C09B61/00 - European:		
Application number:	JP19900170549 19900628		
Priority number (s):	JP19900170549 19900628		
Also published as:	<u>JP2963152 (B2)</u>		

Abstract of JP4057853 (A)

PURPOSE: To prepare a reddish orange coloring matter having a high safety in a high concn. by extracting, with CO2 in a supercritical state, krill shells of which the protein has been decomposed by a protease. CONSTITUTION:Krill shells are treated with a protease to decompose the protein in the shells and the treatment product is filtered. The residue of filtration is dried to give treated shells having a water content of 6-8% and a mean particle size of 200 mum or lower. The treated shells are put into an extraction vessel 5. An extractant comprising a liq.; CO2 in an amt. of 30-40 pts.wt. based on one pt.wt. treated shells having a coloring matter concn. of 30 mg/100 g is supplied through a supercooling apparatus 2 to a pump 3, pressurized at the pump 3 to 100-250 kg/cm<2>, heated with a heat exchanger 4 to 35-40 deg.C to bring it into a supercritical state, and transferred to the extraction vessel 5 to extract an oil in the treated shells. After the pressure of the oil-contg.



CO2 in the supercritical state is reduced to 40-60 kg/cm<2> with a pressure reducing valve 6, the CO2 is delivered through a selector valve 11 to the first separating vessel 7 to separate the oil, and recycled through a selector valve 13, a pressure reducing valve 9, a condense 10, a water separator 15, and a storage vessel 1 to the extraction vessel 5.; Then, selector valves 11 and 13 are closed while selector valves 12 and 14 are opened, and the CO2 contg. the coloring matter is transferred to the second separating

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vessel 8, where the CO2 is evaporated to give a coloring matter with a concn. of 2000-10000 mg/100g.

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			Examination Requeste	d: No	Number	of Claims: 5	(7 Pages Total)
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		(21)	Patent Application No. (22) Filing Date: June 2	: H2-17 28, 1990	70549 D		
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Specification

1. Title of Invention

Method for extracting and separating pigment from krill

2. Claims

(1) A method for extracting and separating pigment from krill, wherein, using as a starting material krill shells that are the residue after krill has been decomposed by a protease and the protein removed, pigment is extracted and separated with supercritical carbon dioxide as an extraction agent.

(2) The method for extracting and separating pigment from krill according to claim 1, wherein extraction and separation are characterized in that

the extract from the krill shells is fractionated while varying the pressure of supercritical carbon dioxide in two stages.

(3) The method for extracting and separating pigment from krill according to claim 1, wherein extraction and separation are characterized in that the extract from the krill shells is fractionated by separation over time without varying the pressure of supercritical carbon dioxide.

(4) The method for extracting and separating pigment from krill according to claim 1, wherein extraction and separation are characterized in that components extracted in an extraction tank are fractionated by a plurality of separation tanks of different conditions. (5) The method for extracting and separating pigment from krill according to any one of claims 1 through 4, characterized in that the moisture content ratio of krill shells is from 10% to 30%.

3. Detailed Description of the Invention (Industrial Field of Use)

The present invention relates to a method for obtaining high-concentration pigment by separating the reddish-orange pigment having a primary component of astaxanthin contained in krill, and in particular, it relates to a method of extraction and separation using supercritical carbon dioxide.

(Prior Art)

The reddish-orange pigment having a primary component of astaxanthin contained in krill has generally been extracted from krill organisms using an organic solvent. This extract contains various components, starting with the lipids that are contained in krill. In particular, concentration and separation of only the pigment contained in the pigment extract is necessary because oxidative decomposition products such as unsaturated fatty acids and glycerol esters thereof bonded to or coexisting with the pigment give off an unpleasant odor, or reaction products in the course of oxidative decomposition such as unsaturated fatty acids cause fading of the pigment.

As methods for concentrating and separating pigment from krill pigment extract liquid, Japanese Unexamined Patent Application Publication No. S60-4558 and Japanese Examined Patent Application Publication No. S61-52183 propose a method in which the pH of krill pigment extract liquid extracted by an organic solvent such as nhexane or acetone is neutralized and lipids are then decomposed by a lipase, and a method in which pigment liquid is separated from a liquid in which an alkali has been added to decompose lipids or other impurities, and then this pigment liquid is extracted and separated by molecular distillation or by a fluid in the supercritical state.

(Problems the Invention is to Solve)

In the proposed krill pigment concentration and separation methods of prior art, numerous steps are required, including a step of extracting krill pigment liquid from krill organisms by an organic solvent, a neutralization step, a step of decomposing lipids and impurities by lipase or alkali, a step of decomposing the decomposition products of impurities and krill pigment, and an extraction step by molecular distillation or a fluid in the supercritical state.

Furthermore, it has been reported that the reddishorange pigment contained in krill has astaxanthin as a primary component and has 100 to 1000 times the antioxidant action of vitamin E, and is anticipated to be used as a drug starting substance in the future. If used as a drug starting substance, however, steps such as solvent removal will be required in order to completely eliminate residue of the organic solvent used in the krill pigment liquid extraction step.

A method has also been considered wherein krill are extracted directly by supercritical carbon dioxide without going through treatment steps, but it is difficult to extract and separate only the pigment because the large amount of moisture and various useful components contained in krill are simultaneously extracted.

(Means for Solving Problems)

The present inventors arrived at the present invention as a result of diligent research to solve the above problems.

Since krill contains a large amount of useful components such as proteins, it has been used in applications such as starting materials for processed foods. Among these applications, the proteins contained in krill have been separated and used in the starting materials of amino acids, rather than krill being used as is. However, the krill shells from which such proteins used in starting materials of amino acids have been removed were discarded or only used as feed for cultivated fish in the past.

The present inventors discovered that krill pigment is produced without going through special pretreatment steps by employing a simple method wherein krill pigment liquid is extracted from krill using a supercritical fluid, by using the shells remaining after components such as proteins from krill have been removed as the starting material of pigment production.

That is to say, pigment is extracted by a supercritical fluid using as a starting material krill shells that are the residue obtained by methods such as filtration after the useful components such as proteins in the krill have been decomposed by enzymes.

The supercritical fluid in the present invention is a fluid in a state beyond the critical temperature and critical pressure. In the case of carbon dioxide, it is the state at 31°C or above, 75.3 Kg/cm² or above;

for propane, 96.7°C or above, 43.4 Kg/cm² or above; for ethane, 9.9°C or above, 52.2 Kg/cm² or above. These fluids are characterized by having density close to that of a liquid and a large expansion coefficient close to that of a gas, and can be used in extraction and separation of various organic matter. In the method of the present invention, carbon dioxide in particular is used as the supercritical fluid. When carbon dioxide gas is used, not only are the steps required in extraction and separation of pigment simplified, but there is absolutely no danger even if the carbon dioxide used as the extraction agent remains in the extracted pigment, for example, and the obtained pigment can be used without a problem in many fields starting

ith pharmaceuticals.

Additionally, supercritical carbon dioxide used as an extraction agent has no risk of explosion or combustion in air like hydrocarbons do. Furthermore, since the critical temperature and critical pressure of carbon dioxide are relatively low, dissolution characteristics can be easily varied by varying the temperature and pressure, and it is possible to perform extraction with an extraction agent having dissolution characteristics suited to pigment extraction and separation.

The method of the present invention is to extract pigment with supercritical carbon dioxide using krill shells as a starting material. The method of the present invention was achieved by studying the extraction conditions such as extraction pressure, temperature and fractionation method for

-reforming efficient extraction and separation of gment, and by studying the water content ratio of the starting material krill.

The present invention will be described below in reference to the drawings.

FIG. 1 is a flowchart of equipment having a means for switching among a plurality of separation tanks for implementing the method of the present invention.

The extraction agent carbon dioxide passes from a liquid carbon dioxide storage tank 1 to a supercooler 2, after which it is pressurized to a prescribed pressure by a pump 3, and then heated to a prescribed temperature by a heat exchanger 4, and supplied as supercritical carbon dioxide to an extraction tank 5 filled with krill shells.

The starting material krill shells primarily contain chitin, proteins, triglyceride esters, triglyceride

esters, monoglyceride esters (oil components) and pigment (astaxanthin). The residue obtained when frozen krill is thawed and then the extract portion in which the proteins decomposed by a protease have been filtered out is a powder with an average particle size of 200 μ m, and normally has a water content ratio of 6% to 8% after it is dried.

Since supercritical carbon dioxide has the characteristic that it decomposes the oil components and pigment of krill shells, only two components are extracted from the krill shells, but in order to separate these two components, the extraction operation is divided into two stages.

Namely, in the first extraction, the oil components contained in the krill shells are extracted by passing through 30 parts by weight to 40 parts by weight of supercritical carbon dioxide having a temperature of 30°C to 50°C and a relatively low pressure of 100 Kg/cm² to 250 Kg/cm² for every 1 part by weight of krill shells having a pigment concentration of 30 mg/100 g, which is equivalent to the concentration of contained astaxanthin.

The supercritical carbon dioxide that contains oil components is reduced in pressure to 40 Kg/cm^2 to 60 Kg/cm^2 by a pressure reducing value 6, and led into a first separation tank 7 via a switching value 11.

In the first separation tank 7, carbon dioxide in the gas state which has separated the oil components is further reduced in pressure and adiabatically expanded by a switching valve 13 and a pressure reducing valve 9, and after being liquefied by a condenser 10, it passes through a water separator 15 and returns to the liquid carbon dioxide storage tank 1 where it is recirculated.

Then, supercritical carbon dioxide is supplied to the extraction tank 5 at a pressure higher than the pressure during the first extraction stage. That is, 30 parts by weight to 40 parts by weight of supercritical carbon dioxide having a temperature of 30° C to 50° C and a pressure of 300 Kg/cm^2 to 500Kg/cm² is supplied to the extraction tank for every 1 part by weight of krill shells, and by closing the switching valves 11 and 13 and opening switching valves 12 and 14, carbon dioxide containing extract with a pressure of 40 Kg/cm² to 60 Kg/cm² is led to a second separation tank 8 by the pressure reducing valve 6.

In the second separation tank 8, carbon dioxide in the gas state is returned to the liquid carbon dioxide storage tank 1 in the same way as in the first extraction step. From the second separation tank, pigment with an extremely high concentration of 2000 mg/100 g to 10,000 mg/100 g can be obtained.

High-concentration pigment is obtained by successive two-stage high-pressure extraction as described above, but it is possible to efficiently collect it by providing a plurality of separation tanks and switching among them.

Furthermore, even if the two-stage extraction is not performed while varying the pressure as described above, it is possible to similarly perform extraction and separation at the same pressure.

That is, extraction and separation of highconcentration pigment is also possible by the extraction operation illustrated in FIG. 2. To describe the operation in reference to FIG. 2, carbon dioxide is fed from the liquid carbon dioxide storage tank 1 through the supercooler 2 to the pump 3, and pressured to a prescribed pressure. Then, it is heated to a prescribed temperature by the heat exchanger 4 to make a supercritical fluid, which is supplied to the extraction tank 5 filled with krill shells.

30 parts by weight to 50 parts by weight of supercritical carbon dioxide at a temperature of 35° C to 50°C and a pressure of 300 Kg/cm² to 500 Kg/cm² is passed through for every 1 part by weight of krill shells (pigment concentration 30 mg/100 g). In the extraction tank, oil components are extracted initially, and then high-concentration pigment is extracted, and the supercritical carbon dioxide gas containing the extract is reduced in pressure to 40 Kg/cm² to 60 Kg/cm² by the pressure reducing valve 6 and led to the first separation tank 7.

The carbon dioxide that comes out from the first separation tank is further reduced in pressure by the pressure reducing valve 9, and after being liquefied by the condenser 10, it passes through the water separator 15 and returns to the liquid carbon dioxide storage tank 1.

In this extraction method, because supercritical carbon dioxide of relatively high pressure is used from the start of extraction, pigment is also extracted together with the oil components, resulting in loss of pigment. Therefore, after extraction is performed by supplying 15 parts by weight to 25 parts by weight of supercritical carbon dioxide for every 1 part of krill shells, the extract, which is primarily made up of oil components, is separated from a feed out valve 16 provided on the bottom of the first separation tank 7. Then, by supplying 15 parts by weight to 25 parts by weight of supercritical carbon dioxide for every 1 part of krill shells, pigment concentrate is obtained in the first separation tank.

The pigment concentration in the oil components obtained by this method was from 10 mg/100 g to 30 mg/100 g, and the pigment concentration in the pigment concentrate was from 2000 mg/100 g to 10,000 mg/100 g.

In this method, because supercritical carbon dioxide of relatively high pressure is used from the start of extraction, a slight amount of pigment is contained in the oil components that constitute the initial extract, but this method has the advantage that extraction time can be shortened compared to the aforementioned method that uses supercritical carbon dioxide in two stages of low pressure and high pressure.

Furthermore, FIG. 3 illustrates a method in which a plurality of tanks are provided in succession, and pigment is efficiently recovered while varying the separation conditions by varying the set pressure and temperature of each separation tank.

The method will be described below in reference to FIG. 3.

Carbon dioxide is fed from the liquid carbon dioxide storage tank 1 through the supercooler 2 to the pump 3, and pressurized to a prescribed pressure. After that, it is heated to a prescribed temperature by the heat exchanger 4 and supplied as supercritical carbon dioxide to the extraction tank 5 filled with krill shells.

Here, the supercritical carbon dioxide supplied to the extraction tank 5 has a temperature of 35° C to 50° C and a pressure of 300 Kg/cm^2 to 500 Kg/cm^2 .

Oil components and pigment are extracted from the krill shells in the extraction tank, and the supercritical carbon dioxide that contains these oil components and pigment is reduced in pressure by the pressure reducing valve 6, and led to a highpressure separation tank 17.

The high-pressure separation tank 17 is held in the supercritical state at a pressure lower than inside the extraction tank at a temperature of 35° C to 50° C and a pressure of 100 Kg/cm² to 300 Kg/cm². Pigment concentrate is collected in the tank, and the supercritical carbon dioxide that contains oil components is reduced in pressure by a pressure

reducing valve 18 and led to a low-pressure separation tank 19.

While the low-pressure separation tank 17 is held in a gas state at a temperature of 20°C to 30°C and a pressure of 40 Kg/cm² to 60 Kg/cm², the carbon dioxide is again reduced in pressure by the pressure reducing valve 9, and after being liquefied by the condenser 10, the moisture it contains is removed by the water separator 15, and the carbon dioxide is returned to the liquid carbon dioxide storage tank 1.

When this method is used, by passing through 30 parts by weight to 40 parts by weight of supercritical carbon dioxide of a relatively high pressure of 300 Kg/cm² to 500 Kg/cm² for every 1 part by weight of krill shells (pigment concentration γ mg/100 g), pigment with an extremely high

Incentration of 2000 mg/100 g to 10,000 mg/100 g can be obtained in the high-pressure separation tank, and oil components having a low pigment concentration can be obtained as an extract in the low-pressure separation tank.

In the method in which the supercritical carbon dioxide supplied to the extraction tank is initially at a relatively low pressure below 300 Kg/cm² and then supercritical carbon dioxide at a relatively high pressure is supplied, the yielded quantity of pigment is high because almost no pigment is extracted in the initial extract, but extraction takes a long time. On the other hand, in the method in which the initial extraction step using supercritical carbon dioxide at a relatively high pressure of 300 Kg/cm²

10 500 Kg/cm² and then the extract of pigment incentrate is separated over time, the equipment configuration is simple and extraction time is short, but since some pigment is contained in the oil components obtained as the initial extract, there is the problem that the yielded quantity of pigment is reduced. However, the method illustrated in FIG. 3, in which a plurality of separation tanks having different set pressures and temperatures is provided and extracts of components are obtained in succession under different extraction conditions, is superior to the aforementioned two methods.

Furthermore, in the present invention, by performing extraction after increasing the moisture content of the krill shells (pigment concentration 30 mg/100 g) used as the starting material to 10 wt% to 30 wt%, it is possible to speed up extraction speed, particularly the initial extraction speed, and as a result, it is possible to reduce the amount of pigment contained in the extract of oil components initially obtained in extraction when supercritical carbon dioxide at a relatively high pressure of 300 Kg/cm² to 500 Kg/cm² is used, and therefore, a reduction of the amount of pigment contained in the oil components and lost can be prevented.

Because water has been added, water is extracted together with pigment, but since water and pigment can be easily separated into two layers, adding water does not hinder extraction and separation of pigment in any way.

However, if the water content ratio exceeds 30%, a reduction in extraction speed in the initial extraction is seen, and therefore it is undesirable if the amount of water exceeds 30%.

It is preferred that the water content ratio of the krill shells be adjusted by controlling the dry state in the krill treatment step, but in cases where krill shells of relatively low moisture content in the dry state are used, it is necessary to disperse water in the krill shells and sufficiently mix before the extraction step.

[Operation]

The present invention is a method for producing pigment made up of astaxanthin contained in krill in which it is extracted using as a production starting material krill shells that are the residue after krill has been decomposed by a protease and the protein and so forth removed, and using supercritical carbon dioxide as an extraction agent. The method of the present invention can produce krill pigment without going through a special pretreatment step using an organic solvent.

(Examples)

The present invention will be described in further detail below by giving examples of the present invention.

Example 1

Using a protease as a protein decomposition enzyme, frozen krill after thawing were made to undergo a proteolysis reaction for 2 hours at 47 °C to 48°C, and then the liquid was filtered and the separated residue was dried to obtain krill shells containing 6% water. An extraction tank having a volume of 25 liters was packed with 6 Kg of these krill shells (pigment concentration 30 mg/100 g). While holding the tank temperature at 40°C, supercritical carbon dioxide having a temperature of 40°C and pressure of 200 Kg/cm² was supplied for 4 hours at a supply rate of 60 Kg per hour. In a separation tank set to a pressure of 50 Kg/cm² and temperature of 30°C, carbon dioxide and liquid were separated, and 1398 g of extract with a pigment concentration of 7.1 mg/100 g was obtained from a feed out valve on the bottom of the separation tank.

Additionally, in the extraction tank, supercritical carbon dioxide at a temperature of 40°C and pressure of 400 Kg/cm² was supplied for 4 hours at a supply rate of 60 Kg per hour, and in the separation tank set to a pressure of 50 Kg/cm² and temperature of 30°C, 13.4 g of high-concentration extract having a pigment concentration of 8331 mg/100 g was obtained from the residue of the previous extract.

Example 2

An extraction tank having a volume of 25 liters was packed with 6 Kg of krill shells (pigment concentration 30 mg/100 g) having the same components as those used as a starting material in example 1. While holding the tank temperature at 40°C, supercritical carbon dioxide having a temperature of 40°C and pressure of 400 Kg/cm² was supplied for 2 hours at a supply rate of 60 Kg per hour. 1703 g of extract with a pigment concentration of 42.8 mg/100 g was obtained from a feed out valve on the bottom of a separation tank set to a pressure of 50 Kg/cm² and temperature of 30°C.

Additionally, in the extraction tank, supercritical carbon dioxide at a temperature of 40°C and pressure of 400 Kg/cm² was supplied for 5 hours at a supply rate of 60 Kg per hour, and in the separation tank set to a pressure of 50 Kg/cm² and temperature of 30°C, 10 g of high-concentration pigment having a pigment concentration of 5874 mg/100 g was obtained.

Example 3

An extraction tank having a volume of 1 liter was packed with 250 g of krill shells (pigment concentration 30 mg/100 g) having the same components as those used as a starting material in example 1. While holding the tank temperature at 40°C, supercritical carbon dioxide having a temperature of 40°C and pressure of 400 Kg/cm² was supplied for 2.5 hours at a supply rate of 2.5 Kg per hour. The supercritical carbon dioxide containing the extract obtained in the extraction tank was supplied to a high-pressure separation tank held in the supercritical state.

While holding the high-pressure separation tank at a temperature of 40°C and pressure of 250 Kg/cm², the extract in the low-pressure separation tank was supplied via a pressure reducing value to a low-pressure separation tank held at 20°C and 60 Kg/cm².

As a result, 0.44 g of pigment with a pigment concentration of 7072 mg/100 g was obtained from the high-pressure separation tank, and 70.57 g of pigment with a pigment concentration of 6.2 mg/100 g was obtained from the low-pressure separation tank.

Example 4

Water was dispersed and sufficiently mixed with 6 Kg of krill shells having the same components as those used as a starting material in example 1, and extraction and separation of pigment were performed while varying the water content ratio of the starting material.

An extraction tank having a volume of 25 liters was packed with the krill shells of different water content ratios, and while holding the extraction tank temperature at 40°C, supercritical carbon dioxide having a temperature of 40°C and pressure of 400 Kg/cm² was supplied, and it was separated in a separation tank set to a pressure of 50 Kg/cm² and temperature of 30°C. The amount of oil component fraction obtained from the start of extraction and the amount of pigment obtained after fractionation of the oil components ended are shown in Table 1 together with extraction time.

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Table 1

	Oil components		Pigment concentrate	
Water content ratio (%)	Extraction time (hours)	Extracted quantity (g) Pigment concentration (mg/100 g)	Extraction time (hours)	Extracted quantity (g) Pigment concentration (mg/100 g)
3	2	1649 12.3	3	18.7 1660
7	2	1703 42.8	3	6.8 7832
14	1	1546 28.3	2	11.9 6084

(Advantageous Effect of the Invention) The present invention is a method that extracts Idish-orange pigment containing astaxanthin from III shells using supercritical carbon dioxide, which makes effective use of krill shells as starting materials, which were treated as waste in the past, by extracting the useful components from krill. Moreover, since it does not use organic solvents and so forth, the process is simple and does not require organic solvent separation steps, and it can extract and separate pigment by a method that is highly safe even in the fields of foods and pharmaceuticals.

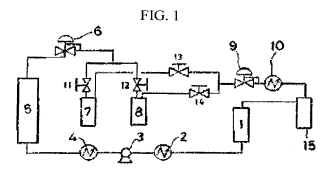
4. Brief Description of the Drawings

FIG. 1 is a flowchart of extraction equipment that switches among multiple separation tanks used for implementing the method of the present invention. FIG. 2 is a flowchart of equipment used when fractionating two components at the same pressure r implementing the method of the present

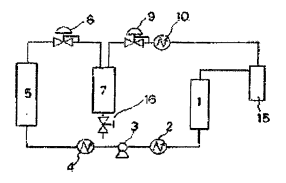
vention. FIG. 3 is a flowchart of equipment having separation tanks at different pressures used for implementing the method of the present invention. Liquid carbon dioxide storage tank ... 1 Supercooler ... 2 Pump ... 3 Heat exchanger ... 4 Extraction tank ... 5 Pressure reducing valve ... 6 First separation tank ... 7 Second separation tank ... 8 Pressure reducing valve ... 9 Condenser ... 10 Switching valves... 11, 12, 13, 14 Water separator ... 15 Feed out valve ... 16 High-pressure separation tank ... 17 Pressure reducing valve ... 18 Low-pressure separation tank ... 19

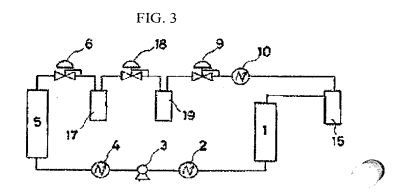
Patent Applicant: Chlorine Engineers Corp. Ltd. (and one other)

Agent: Akira YONEZAWA, Patent Agent (and seven others)









RIMFROST EXHIBIT 1024 page 0620

Electronic Acknowledgement Receipt			
EFS ID:	14233305		
Application Number:	12057775		
International Application Number:			
Confirmation Number:	1945		
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS		
First Named Inventor/Applicant Name:	Inge Bruheim		
Customer Number:	72960		
Filer:	John Mitchell Jones/Vickie Hoeft		
Filer Authorized By:	John Mitchell Jones		
Attorney Docket Number:	NATNUT-14409/US-5/ORD		
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Filing Date:	28-MAR-2008		
Time Stamp:	15:53:28		
Application Type:	Utility under 35 USC 111(a)		

Payment information:

Submitted with Payment			no			
File Listing:						
Document Number	Document Description		File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1	Transmittal Letter	14	409US5ORD_IDSletter11152 012.pdf	81767 f21a078b2154dd35095e098d8d96effd58d 9d672	no	1
Warnings:						
Information:			RIMF	ROST EXHIBIT	1024 p	age 0621

2	Information Disclosure Statement (IDS)	14409US5IDS11152012.pdf	613246	no	5
2	Form (SB08)	1440903510511152012.pdf	06b7557bb6422e7707d2e472de65fbecd78 925ca	10	
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3	Foreign Reference	AU2002322233.pdf	2639818	no	78
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5	5 Non Patent Literature	30102CN1PCTOfficeActionChin	1391615	no	21
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6	Non Patent Literature	Fricke1984.pdf	195109	no	9
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9	Non Patent Literature	Grantham1977.pdf	4188436	no	72
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lf a new appl 1.53(b)-(d) a	tions Under 35 U.S.C. 111 ication is being filed and the applica nd MPEP 506), a Filing Receipt (37 CF ement Receipt will establish the filin	R 1.54) will be issued in due	-	-		
<u>National Stage of an International Application under 35 U.S.C. 371</u> If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course.						
national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course. <u>New International Application Filed with the USPTO as a Receiving Office</u> If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.						

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Inge Bruheim, et alConfirmation:1945Serial No.:12/057,775Group No.:1651Filed:03-28-2008Examiner:Ware, Deborah K.Entitled:BIOEFFECTIVE KRILL OIL COMPOSITIONSFiled:

INFORMATION DISCLOSURE STATEMENT LETTER

EFS Web Filed Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir or Madam:

The citations listed in the attached **IDS Form SB08A** may be material to the examination of the aboveidentified application, and are therefore submitted in compliance with the duty of disclosure defined in 37 C.F.R. §§ 1.56 and 1.97.

Applicants wish to bring to the Examiner's attention that we are not providing copies of US Patents as instructed under 37 CFR 1.98(a)(2). The Examiner is requested to make these citations of official record in this application.

This Information Disclosure Statement under 37 C.F.R. §§ 1.56 and 1.97 is not to be construed as a representation that a search has been made, that additional information material to the examination of this application does not exist, or that any one or more of these citations constitutes prior art.

The Commissioner is hereby authorized to charge any required fees or credit any overpayments to Attorney Deposit Account No.: **50-4302**, referencing Attorney Docket No.: **NATNUT-14409/US-5/ORD**.

Dated: <u>November 15, 2012</u>

<u>/J. Mitchell Jones/</u> J. Mitchell Jones Registration No. 44,174 CASIMIR JONES, S.C. 2275 Deming Way, Suite 310 Middleton, WI 53562 608.662.1277 Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

REQUEST FOR CONTINUED EXAMINATION(RCE)TRANSMITTAL (Submitted Only via EFS-Web)							
Application Number	12057775	Filing Date	2008-03-28	Docket Number (if applicable)	NATNUT-14409/US-5/ORD	Art Unit	1651
First Named Inventor	Inge Bruheim	-		Examiner Name	Ware, Deborah K.		
Request for C	This is a Request for Continued Examination (RCE) under 37 CFR 1.114 of the above-identified application. Request for Continued Examination (RCE) practice under 37 CFR 1.114 does not apply to any utility or plant application filed prior to June 8 1995, or to any design application. The Instruction Sheet for this form is located at WWW.USPTO.GOV						
		S	UBMISSION REQ	UIRED UNDER 37	7 CFR 1.114		
in which they v	were filed unless	applicant in		applicant does not wi	nents enclosed with the RCE wi sh to have any previously filed u		
	v submitted. If a fi n even if this box			any amendments file	ed after the final Office action ma	ay be cor	sidered as a
	nsider the argum	ents in the A	ppeal Brief or Reply	Brief previously filed	l on		
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🗙 Am	endment/Reply						
🗌 🗌 Info	ormation Disclosu	ire Statemer	nt (IDS)				
Affi	Affidavit(s)/ Declaration(s)						
	Other						
MISCELLANEOUS							
	Suspension of action on the above-identified application is requested under 37 CFR 1.103(c) for a period of months (Period of suspension shall not exceed 3 months; Fee under 37 CFR 1.17(i) required)						
Other	Other						
FEES							
 The RCE fee under 37 CFR 1.17(e) is required by 37 CFR 1.114 when the RCE is filed. The Director is hereby authorized to charge any underpayment of fees, or credit any overpayments, to Deposit Account No 504302 							
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Signature of Registered U.S. Patent Practitioner				
Signature	/J. Mitchell Jones/	Date (YYYY-MM-DD)	2012-09-07	
Name	J. Mitchell Jones	Registration Number	44174	

This collection of information is required by 37 CFR 1.114. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450.

If you need assistance in completing the form, call 1-800-PTO-9199 and select option 2.

The Privacy Act of 1974 (P.L. 93-579) requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b)(2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

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- 5. A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
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- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (i.e., GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
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- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.

Electronic Patent Application Fee Transmittal					
Application Number:	12	057775			
Filing Date:	28-	Mar-2008			
Title of Invention:	BIC	DEFFECTIVE KRILL O	IL COMPOSITIC	NS	
First Named Inventor/Applicant Name:	Inge Bruheim				
Filer:	John Mitchell Jones/Vickie Hoeft				
Attorney Docket Number:	NA	TNUT-14409/US-5/0	ORD		
Filed as Large Entity					
Utility under 35 USC 111(a) Filing Fees					
Description		Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Basic Filing:					
Pages:					
Claims:					
Miscellaneous-Filing:					
Petition:					
Patent-Appeals-and-Interference:					
Post-Allowance-and-Post-Issuance:	Post-Allowance-and-Post-Issuance:				
Extension-of-Time:					

Description	Fee Code	Fee Code Quantity		Sub-Total in USD(\$)	
Miscellaneous:					
Request for continued examination	1801	1	930	930	
	Tot	al in USD) (\$)	930	

Electronic Ac	Electronic Acknowledgement Receipt				
EFS ID:	13684183				
Application Number:	12057775				
International Application Number:					
Confirmation Number:	1945				
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS				
First Named Inventor/Applicant Name:	Inge Bruheim				
Customer Number:	72960				
Filer:	John Mitchell Jones/Vickie Hoeft				
Filer Authorized By:	John Mitchell Jones				
Attorney Docket Number:	NATNUT-14409/US-5/ORD				
Receipt Date:	07-SEP-2012				
Filing Date:	28-MAR-2008				
Time Stamp:	13:45:19				
Application Type:	Utility under 35 USC 111(a)				

Payment information:

Submitted with Payment	yes			
Payment Type	Deposit Account			
Payment was successfully received in RAM	\$930			
RAM confirmation Number	11			
Deposit Account	504302			
Authorized User				
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Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1		14409US5Response09072012.	88799	yes	18
		pdf	f3a583da3d9837582e4692fa45f70835a4a1 a747	yes	10
	Multip	art Description/PDF files in a	zip description		
	Document Des	scription	Start	E	nd
	Amendment Al	fter Final	1		1
	Claims		2		3
	Applicant Arguments/Remarks	14	18		
Warnings:					
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2	Request for Continued Examination	14409US5RCE09072012.pdf	697798	no	3
	(RCE)		251edde30526ba30b1e5b4929b56cf13fb4 7194f		5
Warnings:			· · ·	-	
Information:					
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If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application.

National Stage of an International Application under 35 U.S.C. 371

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New International Application Filed with the USPTO as a Receiving Office

If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Bruheim et al.Serial No.:12/057,775Filed:March 28, 200Entitled:BIOEFFECTIV

Bruheim et al.Art12/057,775ExMarch 28, 2008CoBIOEFFECTIVE KRIL OIL COMPOSITIONS

Art Unit: 1651 Examiner: Ware Confirmation: 1945

REQUEST FOR CONTINUED EXAMINATION AND RESPONSE TO OFFICE ACTION MAILED JUNE 7, 2012

EFS WEB FILED

Commissioner for Patents P.O. Box 1450 Alexandria, Virginia 22313-1450

Examiner Ware:

This communication is responsive to the Office Action mailed June 7, 2012. The Commissioner is hereby authorized to charge any fees during the entire pendency of this application, including fees due under 37 C.F.R. §§ 1.16 and 1.17 that may be required, including any required extension of time fees, or credit any overpayment to Deposit Account 50-4302, referencing Attorney Docket No. **NATNUT-14409/US-5/ORD**. This paragraph is intended to be a CONSTRUCTIVE PETITION FOR EXTENSION OF TIME in accordance with 37 C.F.R. § 1.136(a)(3).

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CLAIM AMENDMENTS

1. (Withdrawn) A composition comprising:

from about 3% to 10% ether phospholipids on a w/w basis;

from about 35% to 50% non-ether phospholipids on w/w basis, so that the total amount of ether phospholipids and non-ether phospholipids in the composition is from about 48% to 60% on a w/w basis;

from about 20% to 45% triglycerides on a w/w basis; and from about 400 to about 2500 mg/kg astaxanthin.

2. (Withdrawn) The composition of Claim 1, wherein said ether phospholipids are selected from the group consisting of alkylacylphosphatidylcholine, lyso-alkylacylphosphatidylcholine, alkylacylphosphatidylethanolamine, and combinations thereof.

3. (Withdrawn) The composition of Claim 1, wherein said ether lipids are greater than 90% alkylacylphosphatidylcholine.

4. (Withdrawn) The composition of Claim 1, wherein said non-ether phospholipids are selected from the group consisting of phosphatidylcholine, phosphatidylserine, phosphatidylethanolamine and combinations thereof.

5. (Withdrawn) The composition of Claim 1, wherein said composition comprises a blend of lipid fractions obtained from *Euphausia superba*.

6. (Withdrawn) The composition of Claim 1, wherein said composition comprises from about 25% to 30% omega-3 fatty acids as a percentage of total fatty acids and wherein from about 80% to 90% of said omega-3 fatty acids are attached to said phospholipids.

7. (Withdrawn) A capsule containing the composition of Claim 1.

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8. (Withdrawn) A composition comprising:
from about 3% to 10% ether phospholipids on a w/w basis; and
from about 400 to about 2500 mg/kg astaxanthin.

9. (Withdrawn) The composition of Claim 8, further comprising from about 35% to 50% non-ether phospholipids on w/w basis, so that the total amount of ether phospholipids and non-ether phospholipids in the composition is from about 38% to 60% on a w/w basis.

10. (Withdrawn) The composition of Claim 8, further comprising from about 20% to 45% triglycerides on a w/w basis.

11. (Withdrawn) The composition of Claim 8, wherein said ether phospholipids are selected from the group consisting of alkylacylphosphatidylcholine, lyso-alkylacylphosphatidylcholine, alkylacylphosphatidylethanolamine, and combinations thereof.

12. (Withdrawn) The composition of Claim 11, wherein said ether lipids are greater than 90% alkylacylphosphatidylcholine.

13. (Withdrawn) The composition of Claim 8, wherein said non-ether phospholipids are selected from the group consisting of phosphatidylcholine, phosphatidylserine, phosphatidylethanolamine and combinations thereof.

14. (Withdrawn) The composition of Claim 8, wherein said composition comprises a blend of lipid fractions obtained from *Euphausia superba*.

15. (Withdrawn) The composition of Claim 10, wherein said composition comprises from about 25% to 30% omega-3 fatty acids as a percentage of total fatty acids and wherein from about 80% to 90% of said omega-3 fatty acids are attached to said phospholipids.

16. (Withdrawn) A capsule containing the composition of Claim 8.

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17. (Withdrawn) A blended krill oil composition comprising: from about 45% to 55% w/w phospholipids; from about 20% to 45% w/w triglycerides; and from about 400 to about 2500 mg/kg astaxanthin.

18. (Withdrawn) The composition of Claim 17, wherein said blended krill oil product comprises a blend of lipid fractions obtained from *Euphausia superba*.

19. (Withdrawn) The composition of Claim 17, wherein said composition comprises from about 25% to 30% omega-3 fatty acids as a percentage of total fatty acids and wherein from about 80% to 90% of said omega-3 fatty acids are attached to said phospholipids.

20. (Withdrawn) A *Euphausia superba* krill oil composition comprising:

from about 3% to about 10% w/w ether phospholipids;

from about 27% to 50% w/w non-ether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w;

from about 20% to 50% w/w triglycerides;

from about 400 to about 2500 mg/kg astaxanthin; and

from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

21. (Withdrawn) A dietary supplement comprising encapsulated *Euphausia superba* krill oil comprising from about 3% to about 10% w/w ether phospholipids; from about 27% to 50% w/w non-ether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w; from about 20% to 50% w/w triglycerides; from about 400 to about 2500 mg/kg astaxanthin; and from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

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22. (Withdrawn) A method of making a *Euphausia superba* krill oil composition comprising: contacting *Euphausia superba* with a polar solvent to provide a polar extract comprising phospholipids;

contacting *Euphasia superba* with a neutral solvent to provide a neutral extract comprising triglycerides and astaxanthin;

combining said polar extract and said neutral extract to provide *Euphausia superba* krill oil from about 3% to about 10% w/w ether phospholipids; from about 27% to 50% w/w nonether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w; from about 20% to 50% w/w triglycerides; from about 400 to about 2500 mg/kg astaxanthin; and from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

23. (Withdrawn) The method of Claim 22, further comprising the step of encapsulating the *Euphausia superba* krill oil.

24. (Withdrawn) A *Euphausia superba* krill oil produced by the method of Claim 22.

25. (Withdrawn) A method of producing a dietary supplement comprising;

contacting *Euphausia superba* with a polar solvent to provide an polar extract comprising phospholipids;

contacting *Equphasia superba* with a neutral solvent to provide a neutral extract comprising triglycerides and astaxanthin;

combining said polar extract and said neutral extract to provide *Euphausia superba* krill oil from about 3% to about 10% w/w ether phospholipids; from about 27% to 50% w/w nonether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w; from about 20% to 50% w/w triglycerides; from about 400 to about 2500 mg/kg astaxanthin; and from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids;

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encapsulating said Euphausia superba krill oil.

26. (Withdrawn) A composition comprising at least 65% (w/w) of phospholipids, said phospholipids characterized in containing at least 35% omega-3 fatty acid residues.

27. (Withdrawn) The composition according to claim 26, wherein the composition is derived from a marine or aquatic biomass.

28. (Withdrawn) The composition according to claim 26, wherein the composition is derived from krill.

29. (Withdrawn) The composition of Claim 26, wherein said composition comprises less than2% free fatty acids.

30. (Withdrawn) The composition of Claim 26, wherein said composition comprises less than 10% triglycerides.

31. (Withdrawn) The composition of Claim 26, wherein said phospholipids comprise greater than 50% phosphatidylcholine.

32. (Withdrawn) The composition of Claim 26, wherein the composition comprises at least 500 mg/kg astaxanthin esters.

33. (Withdrawn) The composition of Claim 26, wherein the composition comprises at least 500 mg/kg astaxanthin esters and at least 36% (w/w) omega-3 fatty acids.

34. (Withdrawn) The composition of Claim 26, wherein the composition comprises less than about 0.5g/100g total cholesterol.

35. (Withdrawn) The composition of Claim 26, wherein the composition comprises less than about 0.45% arachidonic acid (w/w).

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36. (Withdrawn) A krill lipid extract comprising at least 500 mg/kg astaxanthin esters and at least 36% (w/w) omega-3 fatty acids.

37. (Withdrawn) A krill lipid extract comprising at least 100 mg/kg astaxanthin esters, at least 20% (w/w) omega-3 fatty acids, and less than about 0.45% arachidonic acid (w/w).

38. (Withdrawn) A method comprising administering the composition of Claim 1 to a subject in an amount effective for reducing insulin resistance, reducing inflammation, improving blood lipid profile and reducing oxidative stress.

39. (Withdrawn) A krill lipid extract comprising greater than about 80% triglycerides and greater than about 90 mg/kg astaxanthin esters.

40. (Withdrawn) The krill lipid extract of Claim 39, characterized in containing from about 5% to about 15% omega-3 fatty acid residues.

41. (Withdrawn) The krill lipid extract of Claim 39, characterized in containing less than about 5% phospholipids.

42. (Withdrawn) The krill lipid extract of Claim 39, characterized in comprising from about 5% to about 10% cholesterol.

43. (Withdrawn) A krill meal composition comprising less than about 50g/kg total fat.

44. (Withdrawn) The krill meal composition of Claim 43 comprising from about 5 to about 20 mg/kg astaxanthin esters.

45. (Withdrawn) The krill meal composition of Claim 43 comprising greater than about 65% protein.

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46. (Withdrawn) The krill meal composition of Claim 43 comprising greater than about 70% protein.

47. (Withdrawn) An animal feed comprising the krill meal of Claim 46.

48. (Withdrawn) A method of increasing flesh coloration in an aquatic species comprising feeding said aquatic species a composition comprising the krill meal of Claim 46.

49. (Withdrawn) A method of increasing growth and overall survival rate of aquatic species by feeding the krill meal of Claim 46.

50. (Currently amended) A method of producing krill oil <u>containing phospholipids</u> comprising:

a) cooking and drying krill to provide cooked and dried krill meal; and

b) extracting contacting said cooked and dried krill meal with a polar solvent to extract a krill oil containing phospholipids from said cooked and dried krill meal, wherein said krill oil containing phospholipids comprises from about 3% to about 10% w/w ether phospholipids; from about 27% to 50% w/w non-ether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w; and from about 20% to 50% w/w triglycerides; and

c) providing a delipidated krill meal following said extraction comprising greater than 65% protein and less than 50 g/kg total fat.

51. (Cancelled)

52. (Currently amended) The method of Claim 50, wherein said <u>cooked and dried</u> krill meal is stored prior to said <u>contacting extraction</u> step.

53. (Currently amended) The method of Claim 50, wherein said <u>contacting</u> extracting step <u>further</u> comprises extraction by supercritical fluid extraction.

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54. (Previously presented) The method of Claim 53, wherein said supercritical fluid extraction is a two step process comprising a first extraction step with carbon dioxide and from 1 to 10% of a co-solvent and a second extraction step with carbon dioxide and from 10-30% of a co-solvent, wherein said co-solvent in said first and second extraction steps is a C_1 - C_3 monohydric alcohol.

55. (Currently amended) A krill oil <u>containing phospholipids</u> produced by the method of claim 50.

56. (Withdrawn) A method of production of krill oil comprising:

a) providing fresh krill;

b) treating said fresh krill to denature lipases and phospholipases in said fresh krill to provide a denatured krill product; and

c) extracting oil from said denatured krill product.

57. (Withdrawn) The method of claim 56 in which the denaturation step comprises heating of said fresh krill.

58. (Withdrawn) The method of claim 56 in which the denaturation step comprises heating said fresh krill after grinding.

59. (Withdrawn) The method of claim 56, further comprising storing said denatured krill product at room temperature or below between the denaturation step and the extraction step.

60. (Withdrawn) The method of claim 56, wherein the enzyme denaturation step is achieved by application of heat.

61. (Withdrawn) The method of claim 56, wherein the extraction step comprises use of supercritical carbon dioxide, with or without use of a polar modifier.

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62. (Withdrawn) The method of claim 56, wherein the extraction step comprises the use of ethanol.

63. (Withdrawn) The method of Claim 56, wherein the extraction step comprises ethanol extraction followed by acetone to precipitation of phospholipids.

64. (Withdrawn) The method of Claim 56, wherein said denatured krill product is a meal.

65. (Withdrawn) Oil produced by the method of Claim 56.

66. (Withdrawn) A composition comprising an oil extracted from krill having a phosphatidylcholine content of greater then about 50% (w/w).

67. (Withdrawn) The composition of Claim 66, wherein said oil has a phosphatidylcholine content of greater then about 70% (w/w).

68. (Withdrawn) The composition of Claim 66, wherein said oil has a phosphatidylcholine content of greater then about 80% (w/w).

69. (Withdrawn) The composition of Claim 66, wherein said composition comprises less than 2% free fatty acids.

70. (Withdrawn) The composition of Claim 66, wherein said composition comprises less than 10% triglycerides.

71. (Withdrawn) The composition of Claim 66, wherein the composition comprises at least 500 mg/kg astaxanthin esters.

72. (Withdrawn) The composition of Claim 66, wherein the composition comprises less than about 0.45% arachidonic acid (w/w).

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73. (Withdrawn) A composition comprising odorless krill oil.

74. (Withdrawn) The composition of Claim 73, wherein said odorless krill oil comprises less than about 10 mg/kg (w/w) trimethylamine.

75. (Withdrawn) An odorless krill oil produced by the method comprising:

extracting a neutral krill oil from a krill oil containing material by supercritical fluid extraction to provide a deodorized krill material, wherein said neutral krill oil contains odor causing compounds and

extracting a polar krill oil from said deodorized krill material by supercritical fluid extraction with a polar entrainer to provide an essentially odorless krill oil.

76. (Withdrawn) A composition comprising krill oil containing less than about 70 micrograms/kilogram (w/w) astaxanthin esters.

77. (Withdrawn) The composition of claim 76, comprising less than about 50 micrograms/kilogram (w/w) astaxanthin esters.

78. (Withdrawn) The composition of claim 76, comprising less than about 20 micrograms/kilogram (w/w) astaxanthin esters.

79. (Withdrawn) The composition of claim 76, comprising less than about 5 micrograms/kilogram (w/w) astaxanthin esters.

80. (Withdrawn) A krill oil produced by the process comprising:

pumping fresh krill from a trawl onto a ship, heating the krill to provide a krill material, and extracting oil from the krill material.

81. (Withdrawn) A method of reducing diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction or hepatic steatosis comprising:

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in a subject exposed to a high fat diet, administering to said subject exposed to a high fat diet an effective amount of a krill oil composition under conditions such that a condition selected from the group consisting of diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction and hepatic steatosis is reduced.

82. (Withdrawn) The method of Claim 81, wherein said effective amount of a krill oil composition is from 0.2 grams to 10 grams of said krill oil composition.

83. (Withdrawn) The method of Claim 81, wherein said krill oil composition comprises: from about 45% to 55% w/w phospholipids; from about 35% to 45% w/w triglycerides; and from about 400 to about 2500 mg/kg astaxanthin.

84. (Withdrawn) The method of Claim 81, wherein said krill oil composition comprises a blend of lipid fractions obtained from *Euphausia superba*.

85. (Withdrawn) A method of reducing diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction or hepatic steatosis comprising in a subject consuming a high fat diet or a normal fat diet:

administering to said subject consuming a high fat diet or a normal fat diet an effective amount of a krill oil composition under conditions such that a condition selected from the group consisting of diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction and hepatic steatosis is reduced.

86. (Withdrawn) A method of inducing diuresis in a subject comprising: administering to said subject an effective amount of a krill oil composition under conditions such that diuresis is induced.

87. (Withdrawn) A method of increasing muscle mass in a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that muscle mass is increased.

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- 88. (Withdrawn) A method of decreasing protein catabolism in a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that protein catabolism is decreased.
- 89. (Withdrawn) A method of decreasing lipid content in the heart of a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that lipid content in the heart of the subject is decreased.
- 90. (Withdrawn) A method of decreasing lipid content in the liver of a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that lipid content in the liver of the subject is decreased.

REMARKS

Claims 50 and 51-55 are pending and under examination following entry of this amendment. Claims 50, 52, 53 and 55 have been amended. Support for the amendments may be found in the specification, for example at page 7, lines 19-30 (polar solvent and phospholipid composition), page 3, line 26 - page 4, line 11 (phospholipid composition), page 5, lines 6 - 12 (meal composition), and the claims as originally filed among other places. No new matter has been added. All amendments and cancellation of claims are made without acquiescing to any of the Examiner's arguments or rejections, and solely for the purpose of expediting the patent application process in a manner consistent with the PTO's Patent Business Goals (PBG), and without waiving the right to prosecute the cancelled claims (or similar claims) in the future.

The following rejections are at issue:

1. Claims 50, 52 and 55 are rejected as anticipated by Japanese Abstract 04-057853 or US 2003/0113432;

2. Claims 50 and 52-55 are rejected as being obvious over Japanese Abstract 04-057853 and US 2003/0113432 in view of Kamiya (US 20060193962).

These rejections are addressed in order below.

1. The claims are not anticipated

Claims 50-53 and 55 are rejected as anticipated by Japanese Abstract 04-057853 or US 2003/0113432. Applicants respectfully disagree. Nevertheless, Applicants have amended the claims to clarify that the extraction process with a polar solvent yields a krill oil with a specified phospholipid content as well as a high-protein, low fat residual meal. The Japanese abstract discloses a protease treated and mechanically ground composition:

Krill shells are treated with a protease to decompose the protein in the shells and the treatment product is filtered. The residue of filtration is dried to give treated shells having a water content of 6-8% and a mean particle size of 200 μ m or lower. The treated shells are put into an extraction vessel 5.

The purpose of the process is to extract a coloring pigment from krill shells: "To prepare a reddish orange coloring matter having a high safety in a high concn. by extracting, with CO2 in a supercritical state, krill shells of which the protein has been decomposed by a protease."

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Applicants respectfully submit that the alleged prior art process, which uses only krill shells, is substantially different from the claimed process which uses a polar solvent to extract a specified phospholipid-rich oil and high protein, low fat krill meal.

In response to Applicant's previous arguments, the Examiner states:

The argument that only krill shells are used is noted, however, the krill oil is obtained from krill meal which the reference does disclose. However, newly cited WO teaches that the krill meal contains all of the components of krill and will produce the krill oil as claimed. Also parts of the krill organism can remain in the shell and be treated as well because the organism is enclosed by the carapace of the shell(s). The reference does not disclose that the krill is removed entirely from the shells and that the shells are absent of any krill remainder. Thus, in light of the newly cited reference and these arguments the rejection is sustained over at least some of the claims.

The currently amended claim address the reasoning by the Examiner with respect to the Japanese abstract as extraction from krill shells, even if they have residual material associated with the shells, would not yield a high phospholipid krill oil or low fat, high protein meal as claimed. The shells of krill are primarily composed of chitin which is a carbohydrate, not a protein. Therefore, the residual material would be primarily chitin. Furthermore, the extract from the krill shells would comprise lipids associated with the shells, primarily a pigment (astaxanthin which is normally present as an ester with a fatty acid). This process would not use a polar solvent as claimed because the pigments are poorly soluble in polar solvents. A non-polar solvent would have be utilized for the extraction. Furthermore, the extracted material would not contain etherphospholipids, phospholipids, or triglycerides in the specified amounts.

With respect to US 2003/0113432, Applicant first notes that paragraph 45 does not teach extraction of a lipid from the disclosed powder. The Examiner states that:

WO teaches Krill lipid (e.g. krill oil) at [0045], lines 1-8. An apparatus comprising a cooker and a drier is disclosed at [0051], at page 5, col. 1, lines 1-2. Heating which is akin to cooking is disclosed at page 5, line 17 at col. 2. Cooking and drying step is disclosed at [0051], lines 1-12, to provide dried krill meal. Extraction of Krill oil is disclosed at [0045], line 3. WO clearly teaches that the krill meal contains all of its components and is not extracted and since it is dried it can inherently be stored before extraction.

Applicant respectfully submits that paragraph 45 does not teach extraction of an oil from the krill powder described elsewhere in the application. This paragraph is a generic statement directed to

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krill oil in general, not krill oil extracted from a specific source material by a polar solvent. As provided in paragraph 45:

[0045] There are several indexes indicating a degree of lipid degradation. About the lipid in krill, particularly, the krill lipid having been extracted and refined, it is known that, during the preservation, a peroxide value hardly increases and only a carbonyl value increases. In other words, it is pointed out that degradation of the krill lipid differs in creation of oxides and progress rate of the decomposing reaction from those in general fish oil, etc.

As can be seen, this paragraph merely makes a statement degradation of stored krill oil is different than fish oil. There is no statement that krill lipids being discussed were extracted from the powder described in the remainder of the specification. Furthermore, the paragraphs around paragraph 45 address the lipids found in the powder, not lipids extracted from the powder. The comparison is between a powder to which antioxidants have been added as compared to a powder where no antioxidants have been added.

In any event, the current claims have been amended to clearly distinguish US 2003/0113432. US 2003/0113432 does not teach any extraction from the krill powder, much less extraction with a polar solvent to provide a krill oil with the specified phospholipid content. Likewise. US 2003/0113432 does not teach a process where a high protein, low fat meal is obtained.

Applicants further submit that neither Japanese Abstract 04-057853 nor US 2003/0113432 teach a krill oil with the lipid content defined in Claim 50.

Applicants respectfully submit that the cited references do not teach each element of the claims and thus request that the anticipation rejections be withdrawn.

2. The claims are not obvious

Claims 50 and 52-55 are rejected as being obvious over Japanese Abstract 04-057853 and US 2003/0113432 in view of Kamiya (US 20060193962) Applicants respectfully disagree. In any event, the amendments to the claims address the rejection. None of the references, alone or combined teach extraction of a krill oil with the claimed phospholipid content or production of a krill meal with the claimed protein and fat content.

As discussed above, the currently amended claim address the reasoning by the Examiner with respect to the Japanese abstract as extraction from krill shells, even if they have residual

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material associated with the shells, would not yield a high phospholipid krill oil or low fat, high protein meal as claimed. The shells of krill are primarily composed of chitin which is a carbohydrate, not a protein. Therefore, the residual material would be primarily chitin. Furthermore, the extract from the krill shells would comprise lipids associated with the shells, primarily a pigment (astaxanthin which is normally present as an ester with a fatty acid). This process would not use a polar solvent as claimed because the pigments are poorly soluble in polar solvents. A non-polar solvent would have to be utilized for the extraction. Furthermore, the extracted material would not contain etherphospholipids, phospholipids, or triglycerides in the specified amounts.

As also discussed above, US 2003/0113432 does not teach extraction of any lipid from the powders disclosed in the application, much less use a polar solvent to provide oils or meals with the specified compositions.

Kamiya does not cure these deficiencies. In particular, Kamiya is directed to extraction from Hydrangea. Neither the nature nor the target of the extraction is identified. Instead, Kamiya lists a number of different, generic extraction technologies. There is no disclosure of krill or even phospholipid extracts in general.

As can be seen, none of the references, alone or combined, teach the processes for production of a krill oil or meal with the specified compositions. Thus, any prima facie case of obviousness established by the Examiner is rebutted because the combined references do not teach element of the claims.

Furthermore, the references are not properly combinable in view of the amended claims. Japanese Abstract 04-057853 is directed to extraction of a coloring pigment from the shells of krill. US 2003/0113432 does not teach extraction of any lipid from the krill powders disclosed in the application with any type of solvent, polar or otherwise. Kamiya is directed to extraction of some unidentified compound from Hydrangea and simply lists a number of extraction methods that could be used. There is no motivation in any of the references to make a polar phospholipid extract and the resulting high protein, low fat meal from a cooked and dried krill meal as claimed. Thus, any prima facie case of obviousness established by the Examiner is rebutted because the references are not properly combinable in view of the amendments to the claims.

For these reasons, Applicant requests that the obviousness rejection be withdrawn.

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CONCLUSION

If a telephone interview would aid in the prosecution of this application, the Examiner is encouraged to call the undersigned collect at (608) 662-1277.

Dated: September 7, 2012

/J. Mitchell Jones/

John Mitchell Jones Registration No. 44,174

Casimir Jones, S.C. 2275 Deming Way, Suite 310 Middleton, WI, 53562 (608) 662-1277

PTO/SB/06 (07-06)

Approved for use through 1/31/2007. OMB 0651-0032 LLC Detent and T

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process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 12 minutes to complete including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. **SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.**

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	ED STATES PATENT	TAND TRADEMARK OFFICE	UNITED STATES DEPAR United States Patent and Address: COMMISSIONER F P.O. Box 1450 Alexandria, Virginia 22: www.uspto.gov	FOR PATENTS		
APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.		
12/057,775	03/28/2008	Inge Bruheim	NATNUT-14409/US-5/ORD	1945		
72960 Casimir Jones,	7590 06/07/2012 S C		EXAMINER WARE, DEBORAH K			
2275 DEMING	WAY, SUITE 310					
MIDDLETON,	W1 53562		ART UNIT	PAPER NUMBER		
			1651			
			MAIL DATE	DELIVERY MODE		
			06/07/2012	PAPER		

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

	Application No.	Applicant(s)								
	12/057,775	BRUHEIM ET AL.								
Office Action Summary	Examiner	Art Unit								
	DEBBIE K. WARE	1651								
The MAILING DATE of this communication app Period for Reply	ears on the cover sheet with the c	correspondence address								
 A SHORTENED STATUTORY PERIOD FOR REPLY WHICHEVER IS LONGER, FROM THE MAILING D/ Extensions of time may be available under the provisions of 37 CFR 1.13 after SIX (6) MONTHS from the mailing date of this communication. If NO period for reply is specified above, the maximum statutory period v Failure to reply within the set or extended period for reply will, by statute, Any reply received by the Office later than three months after the mailing earned patent term adjustment. See 37 CFR 1.704(b). 	ATE OF THIS COMMUNICATION 36(a). In no event, however, may a reply be tir vill apply and will expire SIX (6) MONTHS from , cause the application to become ABANDONE	N. nely filed the mailing date of this communication. D (35 U.S.C. § 133).								
Status										
 1) Responsive to communication(s) filed on <u>04 April 2012</u>. 2a) This action is FINAL. 2b) This action is non-final. 3) An election was made by the applicant in response to a restriction requirement set forth during the interview on; the restriction requirement and election have been incorporated into this action. 4) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i>, 1935 C.D. 11, 453 O.G. 213. 										
Disposition of Claims										
 5) Claim(s) <u>1-50 and 52-90</u> is/are pending in the a 5a) Of the above claim(s) <u>1-49 and 56-90</u> is/are 6) Claim(s) is/are allowed. 7) Claim(s) <u>50 and 52-55</u> is/are rejected. 8) Claim(s) is/are objected to. 9) Claim(s) are subject to restriction and/or 	withdrawn from consideration.									
Application Papers										
 10) The specification is objected to by the Examine 11) The drawing(s) filed on is/are: a) according a contract of the second seco	epted or b) objected to by the drawing(s) be held in abeyance. Se ion is required if the drawing(s) is ob	e 37 CFR 1.85(a). jected to. See 37 CFR 1.121(d).								
	priority under 35 U.S.C. § 119(a))-(d) or (f).								
 13) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: Certified copies of the priority documents have been received. Certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 										
Attachment(s) 1)	4) Interview Summary Paper No(s)/Mail D 5) Notice of Informal F 6) Other:	ate								

Office Action Summary RIMFROST EXHIBIT 1024 page 0653

DETAILED ACTION

Claims 1-50 and 52-90 are pending.

Response to Amendment

The Amendments filed April 4, 2012, were received and entered.

Information Disclosure Statement

The information disclosure statements (IDSs) submitted on March 21, 2012, February 21, 2012 and January 25, 2012, were received. The submissions are in compliance with the provisions of 37 CFR 1.97. Accordingly, the information disclosure statements are being considered by the examiner.

Election/Restrictions

Applicant's election without traverse of Group VIII, claims 50-55, 51, now canceled so remaining elected, claims 50 and 52-55, original election in the reply filed on October 31, 2011, and was acknowledged.

Claims 1-49 and 56-90 are hereby withdrawn from further consideration pursuant to 37 CFR 1.142(b) as being drawn to a nonelected inventions, there being no allowable generic or linking claim. Election was made without traverse in the reply filed on October 31, 2011.

Claim Rejections - 35 USC § 102

The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

A person shall be entitled to a patent unless -

(b) the invention was patented or described in a printed publication in this or a foreign country or in public use or on sale in this country, more than one year prior to the date of application for patent in the United States.

Claims 50, 52 and 55 are rejected under 35 U.S.C. 102(b) as being clearly anticipated by Patent Abstract of Japan 04-057853, dated Feb. 25, 1992, cited on previously enclosed PTO-1449 Form or **newly cited** US 2003/0113432, cited on enclosed PTO-892 form.

Claims drawn to method for producing oil and an oil produced thereby.

Abstract 04-057853 teaches method for extracting krill oil comprising a)providing krill meal; and extracting oil from the krill meal (powdered form of krill parts). The meal (powdered form of krill parts) can be provided from heat-treated krill parts and is storable. The extracting is carried out by supercritical extraction. An oil is produced by the method.

WO teaches Krill lipid (e.g. krill oil) at [0045], lines 1-8. An apparatus comprising a cooker and a drier is disclosed at [0051], at page 5, col. 1, lines 1-2. Heating which is akin to cooking is disclosed at page 5, line 17 at col. 2. Cooking and drying step is disclosed at [0051], lines 1-12, to provide dried krill meal. Extraction of Krill oil is disclosed at [0045], line 3. WO clearly teaches that the krill meal contains all of its components and is not extracted and since it is dried it can inherently be stored before extraction.

The claims are identical to the abstract and WO as discussed above and are considered to be clearly anticipated by the teachings therein. Krill shells are part of krill

and oil is obtained from the krill parts. The krill parts are dried and hence subjected to heating to provide for the krill meal which is subjected to supercritical extraction in two steps to obtain the oil. WO clearly teaches cooking and drying and extraction is disclosed as well which will be carried out on a prepared product having all the contents including oil or lipid. The krill lipid or oil is not different than any krill oil or lipid as disclosed in the art or Applicants have not shown a single difference. The krill oil as claimed must be different than the oil or lipid as disclosed, no matter how it is prepared. Krill meal can be stored before it is desired to extract an oil therefrom. The claims are anticipated by the cited references.

Response to Arguments

Applicant's arguments filed April 4, 2012, have been fully considered but they are not persuasive. The argument that only krill shells are used is noted, however, the krill oil is obtained from krill meal which the reference does disclose. However, newly cited WO teaches that the krill meal contains all of the components of krill and will produce the krill oil as claimed. Also parts of the krill organism can remain in the shell and be treated as well because the organism is enclosed by the carapace of the shell(s). The reference does not disclose that the krill remainder. Thus, in light of the newly cited reference and these arguments the rejection is sustained over at least some of the claims.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

The factual inquiries set forth in Graham v. John Deere Co., 383 U.S. 1, 148 USPQ 459 (1966), that are applied for establishing a background for determining obviousness under 35 U.S.C. 103(a) are summarized as follows:

1. Determining the scope and contents of the prior art.

2. Ascertaining the differences between the prior art and the claims at issue.

3. Resolving the level of ordinary skill in the pertinent art.

4. Considering objective evidence present in the application indicating obviousness or nonobviousness.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to

consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).

Claims 50 and 52-55 are rejected under 35 U.S.C. 103(a) as being unpatentable over JP as cited and discussed above and the **newly cited** WO cited above, in view of Kamiya et al (US 20060193962A1), cited on enclosed PTO-892 Form.

Claims are discussed above as if the JP abstract.

WO teaches Krill lipid (e.g. krill oil) at [0045], lines 1-8. An apparatus comprising a cooker and a drier is disclosed at [0051], at page 5, col. 1, lines 1-2. Heating which is akin to cooking is disclosed at page 5, line 17 at col. 2. Cooking and drying step is disclosed at [0051], lines 1-12, to provide dried krill meal. Extraction of Krill oil is disclosed at [0045], line 3. WO clearly teaches that the krill meal contains all of its components and is not extracted and since it is dried it can inherently be stored before extraction.

Kamiya et al, US 20060193962A1, teach extraction with supercritical fluid and solvent [0043], and the solvent can be a monohydric alcohol [0049], ranging from 1 to 20% [0059].

Claim differs from JP in that monohydric alcohol is not disclosed. It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to carry out supercritical extraction of JP and WO using a cosolvent monohydric alcohol as disclosed by Kamiya et al to produce oil from krill. To carry out more than one extract step is an obvious modification of the cited prior art. Alcohol extraction is a well known extractant as disclosed by Kamiya et al.

Each of the claim feature are disclosed and one of skill would have been motivated to carry out the process steps to provide oil with the expectation of successful results. Clearly the claim is prima facie obvious over the cited prior art.

Response to Arguments

Applicant's arguments filed April 4, 2012, have been fully considered but they are not persuasive. The newly added WO reference clearly teaches that krill oil or lipid can be extracted from cooked and/or dried krill meal as desired, or at least the same is suggested by the teachings of the steps. One of skill may not desire to extract the oil until ready to do so since it will be contained and preserved in the whole contents of the krill until ready and needed for use by those of skill in the art. Supercritical extraction is a well known process and disclosed by the cited prior art combination of references. Each element is disclosed or at least suggested by the cited prior art. The rejection is sustained.

Applicant's amendment necessitated the new ground(s) of rejection presented in this Office action. Accordingly, **THIS ACTION IS MADE FINAL**. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any

extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the date of this final action.

All claims fail to be patentably distinguishable over the state of the art discussed above and cited on the enclosed PTO-892 and/or PTO-1449. Therefore, the claims are properly rejected.

The remaining references listed on the enclosed PTO-892 and/or PTO-1449 are cited to further show the state of the art.

No claims are allowed.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to DEBBIE K. WARE whose telephone number is (571)272-0924. The examiner can normally be reached on 9:30-6:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Mike Wityshyn can be reached on 571-272-0926. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300. Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov.

Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like

Application/Control Number: 12/057,775 Art Unit: 1651 assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/Deborah K. Ware/

Deborah K. Ware

Primary Examiner

Art Unit 1651

Notice of References Cited	Application/Control No. 12/057,775	Applicant(s)/Patent Under Reexamination BRUHEIM ET AL.		
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	DEBBIE K. WARE	1651	Page 1 of 1	

U.S. PATENT DOCUMENTS

*		Document Number Country Code-Number-Kind Code	Date MM-YYYY	Name	Classification
*	А	US-2003/0113432	06-2003	Yoshitomi et al.	426/643
	В	US-			
	С	US-			
	D	US-			
	Е	US-			
	F	US-			
	G	US-			
	н	US-			
	Ι	US-			
	J	US-			
	к	US-			
	L	US-			
	М	US-			

FOREIGN PATENT DOCUMENTS

*		Document Number Country Code-Number-Kind Code	Date MM-YYYY	Country	Name	Classification
	Ν					
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NON-PATENT DOCUMENTS

*		Include as applicable: Author, Title Date, Publisher, Edition or Volume, Pertinent Pages)
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*A copy of this reference is not being furnished with this Office action. (See MPEP § 707.05(a).) Dates in MM-YYYY format are publication dates. Classifications may be US or foreign.

Notice of References Cited

Part of Paper No. 20120529



RIMFROST EXHIBIT 1024 page 0662

Doc description: Information Disclosure Statement (IDS) Filed

Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

Application Number		12057775
Filing Date		2008-03-28
First Named Inventor	Inge E	Bruheim
Art Unit		1651
Examiner Name Ware		Deborah K.
Attorney Docket Number		NATNUT-14409/US-5/ORD

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	1	200	04-534800	JP			2004-11-18	Kohyo			

ALL REFERENCES CONSIDERED EXCREMENTAL STREET 1024R 024 R 026 0663 D.W./ EFS Web 2.1.17

Receipt date: 01/25/2012

INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

Application Number		12057775	12057775 - GAU: 1651				
Filing Date		2008-03-28	2008-03-28				
First Named Inventor	Inge E	Bruheim	uheim				
Art Unit		1651					
Examiner Name	Ware	, Deborah K.					
Attorney Docket Numb	er	NATNUT-144	109/US-5/ORD				

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	1	SIKORSKI, E., "The Ut	SIKORSKI, E., "The Utilization of Krill For Food," Food Process Eng., 1:845-855 (1980)											
	2	BUDZINSKLI, E., et al., "Possibilities of processing and marketing of products made from Antarctic Krill", FAO Fish. Tech. Pap. (268) 46 pages (1985)												
	3	BUNEA R., et al, "Eva Medicine Review, Thor				the Clinical Course of Hyp 5. 4, January 1, 2004	erlipidemia," Alternative							
	4	GORDEEV, K.Y., et al. Khim. Prirod. Soed. 2 (the Main Phosp	holipids of the Antarctic Kri	II, Euphausia superba,"							
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¹ See Kind Codes of USPTO Patent Documents at <u>www.USPTO.GOV</u> or MPEP 901.04. ² Enter office that issued the document, by the two-letter code (WIPO Standard ST.3). ³ For Japanese patent documents, the indication of the year of the reign of the Emperor must precede the serial number of the patent document ⁴ Kind of document by the appropriate symbols as indicated on the document under WIPO Standard ST.16 if possible. ⁵ Applicant is to place a check mark here i English language translation is attached.														

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	Filing Date		2008-03-28		
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	Art Unit		1651		
	Examiner Name Ware		, Deborah K.		
	Attorney Docket Numb	er	NATNUT-14409/US-5/ORD		

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Receipt date: 02/21/2012	Application Number		12057775 - GAU: 165		
	Filing Date		2008-03-28		
	First Named Inventor	Inge E	Bruheim		
(Not for submission under 37 CFR 1.99)	Art Unit		1651		
	Examiner Name	Ware,	, Deborah K.		
	Attorney Docket Number		NATNUT-14409/US-5/ORD		

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WEST Search History for Application 12057775

Creation Date: 2012052914:28

Prior Art Searches

Query	DB	Op.	Plur.	Thes.	Date
''krill oil''	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES		01-03-2012
(''krill oil'') and ''krill meal''	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES		01-03-2012
(''krill oil'' and ''krill meal'') and ''supercritical fluid''	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES		01-03-2012
extract? and krill and oil and meal and supercritical	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES		01-03-2012
2004241249	PGPB	OR	YES		01-03-2012
200400241249	PGPB	OR	YES		01-03-2012
20040241249	PGPB	OR	YES		01-03-2012
(20040241249) and "supercritical"	PGPB	OR	YES		01-03-2012
(20040241249) and "solvent extraction"	PGPB	OR	YES		01-03-2012
(20040241249) and "extract"	PGPB	OR	YES		01-03-2012
(20040241249 and "extract") and "oil"	PGPB	OR	YES		01-03-2012
(20040241249 and "extract" and "oil") and "meal"	PGPB	OR	YES		01-03-2012
supercritical and extraction and krill	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES		01-03-2012
(supercritical and extraction and krill) and co-solvent	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES		01-03-2012
(supercritical and extraction and krill and co-solvent) and oil	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES		01-03-2012
supercritical and extraction and alcohol		OR	YES		01-03-2012

WEST Search History for Application 12057775

	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD			
(supercritical and extraction and alcohol) and monohydric	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES	01-03-2012
(supercritical and extraction and alcohol and monohydric) and krill and meal	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES	01-03-2012
krill and oil and cooking and drying	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES	05-29-2012
(krill and oil and cooking and drying) and extracting	PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD	OR	YES	05-29-2012

Doc description: Information Disclosure Statement (IDS) Filed

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INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

	·		12057775
			2008-03-28
	First Named Inventor Inge E		Bruheim
	Art Unit		1651
	Examiner Name Ware		
	Attorney Docket Number		NATNUT-14409/US-5/ORD

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Examiner Initial*		Foreign Document Number ³	Country Code ²		Kind Code⁴	Publication Date	Name of Patented Applicant of cited Document		Pages,Columns,Lines where Relevant Passages or Relevant Figures Appear	T 5
	1	JP-A-S52-114046	JP			1977-09-24	Kokai			
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INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

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Filing Date		2008-03-28	
First Named Inventor	Inge E	Bruheim	
Art Unit		1651	
Examiner Name Ware			
Attorney Docket Number		NATNUT-144	09/US-5/ORD

Examiner Initials*	Cite No	(book	ude name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item ok, magazine, journal, serial, symposium, catalog, etc), date, pages(s), volume-issue number(s), T lisher, city and/or country where published.			
	1 JP Office Action mailed February 23, 2012, JP Patent Application No. 2010-522444 (and English translation)					
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	Application/Control No.	Applicant(s)/Patent Under Reexamination
Search Notes	12057775	BRUHEIM ET AL.
	Examiner	Art Unit
	DEBBIE K WARE	1651

SEARCHED				
Class	Subclass	Date	Examiner	

SEARCH NOTES		
Search Notes	Date	Examiner
WEST, NPL and INV: see search history print out	12/2011-1/2012	DKW
WEST, NPL and INV: see search history print out	6/2012	DKW

	INTERFERENCE SEARCH		
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Fealey, Terence, Marietta, GA, UNITED STATES Bailes, Julian E., Morgantown, WV, UNITED STATES ΡT US 20110257267 A1 20111020 ΑT US 2010-904049 A1 20101013 (12) PRAI US 2009-251230P 20091013 (61) DT Utility FS APPLICATION LN.CNT 2397 INCL INCLM: 514/547.000 INCLS: 514/560.000; 514/549.000 NCL 514/547.000 NCLM: NCLS: 514/549.000; 514/560.000 IPC IPCI A61K0031-232 [I,A]; A61P0025-00 [I,A]; A61K0031-202 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 3 OF 27 USPATFULL on STN L4AN 2011:251469 USPATFULL SOLVENT-FREE PROCESS FOR OBTAINING PHOSPHOLIPIDS AND NEUTRAL ENRICHED ТΤ KRILL OILS ΙN Sclabos Katevas, Dimitri, Santiage, CHILE Toro Guerra, Raul R., Santiage, CHILE Chiong Lay, Mario M., Santiage, CHILE PA THAROS LTD., Santiago, CHILE (non-U.S. corporation) LONZA LTD., Basel, SWITZERLAND (non-U.S. corporation) ΡI US 20110224450 A1 20110915 AI US 2011-96644 A1 20110428 (13) RLI Continuation-in-part of Ser. No. WO 2009-IB7269, filed on 30 Oct 2009, PENDING Utility DT APPLICATION FS LN.CNT 2021 INCLM: 554/023.000 INCL INCLS: 554/008.000; 554/078.000 NCL NCLM: 554/023.000 554/008.000; 554/078.000 NCLS: IPC IPCI C11B0001-00 [I,A]; C07F0009-10 [I,A] IPCR C11B0001-00 [I,A]; C07F0009-10 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 4 OF 27 USPATFULL on STN T.4 AN 2011:212256 USPATFULL ΤI METHOD FOR PRODUCING LIPIDS ΙN Yoshikawa, Kazuhiro, Tokyo, JAPAN Mikajiri, Akihiro, Tokyo, JAPAN ΡA NIPPON SUISAN KAISHA, LTD., Tokyo, JAPAN (non-U.S. corporation) US 20110189760 ΡT A1 20110804 US 2009-120842 AI 20090924 (13) A1 WO 2009-JP66530 20090924 PCT 371 date 20110425 JP 2008-248986 PRAI 20080926 DT Utility FS APPLICATION LN.CNT 1345 INCLM: 435/271.000 INCL INCLS: 554/020.000 NCL NCLM: 435/271.000 NCLS: 554/020.000 IPC IPCI C11C0001-00 [I,A]; C11B0001-00 [I,A] IPCR C11C0001-00 [I,A]; C11B0001-00 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 5 OF 27 USPATFULL on STN T.4

2011:211870 USPATFULL ΑN ТΤ METHOD FOR CONCENTRATING LIPIDS ΤN Yoshikawa, Kazuhiro, Tokyo, JAPAN PA NIPPON SUISAN KAISHA, LTD., Tokyo, JAPAN (non-U.S. corporation) ΡT US 20110189374 A1 20110804 ΑI US 2009-120875 A1 20090924 (13) WO 2009-JP66529 20090924 20110425 PCT 371 date PRAI JP 2008-248986 20080926 DT Utility APPLICATION FS LN.CNT 961 INCL INCLM: 426/601.000 INCLS: 554/008.000 NCL 426/601.000 NCLM: NCLS: 554/008.000 IPCI TPC A23D0009-00 [I,A]; C11B0001-06 [I,A] IPCR A23D0009-00 [I,A]; C11B0001-06 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. L4ANSWER 6 OF 27 USPATFULL on STN AN 2011:198158 USPATFULL ΤI METHODS OF TREATING AND PREVENTING NEUROLOGICAL DISORDERS USING DOCOSAHEXAENOIC ACID ΙN AISEN, Paul S., Solana Beach, CA, UNITED STATES Quinn, Joseph F., Portland, OR, UNITED STATES Yurko-Mauro, Karin, Silver Spring, MD, UNITED STATES PA MARTEK BIOSCIENCES CORPORATION, Columbia, MD, UNITED STATES (U.S. corporation) A1 20110721 ΡT US 20110177061 A1 20100709 (12) US 2010-833913 ΑT US 2009-224836P PRAI 20090710 (61) US 2010-359792P 20100629 (61) DT Utility FS APPLICATION LN.CNT 2653 INCL INCLM: 424/133.100 INCLS: 514/560.000; 514/120.000; 514/547.000; 514/549.000; 514/297.000; 514/319.000; 514/479.000; 514/215.000; 424/184.100; 424/172.100; 424/152.100; 514/458.000 NCL NCLM: 424/133.100 NCLS: 424/152.100; 424/172.100; 424/184.100; 514/120.000; 514/215.000; 514/297.000; 514/319.000; 514/458.000; 514/479.000; 514/547.000; 514/549.000; 514/560.000 A61K0031-202 [I,A]; A61K0031-661 [I,A]; A61K0031-232 [I,A]; TPC IPCI A61K0031-473 [I,A]; A61K0031-445 [I,A]; A61K0031-27 [I,A]; A61K0031-55 [I,A]; A61K0039-00 [I,A]; A61K0039-395 [I,A]; A61K0031-355 [I,A]; A61P0025-28 [I,A]; A61P0025-00 [I,A] A61K0031-202 [I,A]; A61K0031-232 [I,A]; A61K0031-27 [I,A]; IPCR A61K0031-355 [I,A]; A61K0031-445 [I,A]; A61K0031-473 [I,A]; A61K0031-55 [I,A]; A61K0031-661 [I,A]; A61K0039-00 [I,A]; A61K0039-395 [I,A]; A61P0025-00 [I,A]; A61P0025-28 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 7 OF 27 USPATFULL on STN T.4 2011:146375 USPATFULL AN ΤI KRILL OIL PROCESS IN Breivik, Harald, Porsgrunn, NORWAY Thorstad, Olav, Porsgrunn, NORWAY ΡA PRONOVA BIOPHARMA NORGE AS, Lysaker, NORWAY (non-U.S. corporation) РT US 20110130458 A1 20110602 ΑI US 2009-992365 A1 20090515 (12)

WO 2009-NO184 20090515 20110211 PCT 371 date 20080515 (61) PRAT US 2008-53455P Utility DT APPLICATION FS LN.CNT 688 INCL INCLM: 514/560.000 INCLS: 426/608.000; 426/417.000 NCL 514/560.000 NCLM: 426/417.000; 426/608.000 NCLS: A61K0031-202 [I,A]; A61P0003-06 [I,A]; A61P0003-00 [I,A]; IPC IPCI A61P0009-00 [I,A]; A61P0009-04 [I,A]; A61P0009-10 [I,A]; A23D0007-00 [I,A]; A23D0009-00 [I,A] TPCR A61K0031-202 [I,A]; A23D0007-00 [I,A]; A23D0009-00 [I,A]; A61P0003-00 [I,A]; A61P0003-06 [I,A]; A61P0009-00 [I,A]; A61P0009-04 [I,A]; A61P0009-10 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 8 OF 27 USPATFULL on STN T.4 2011:117434 USPATFULL AN ТΤ POWDERED COMPOSITION CONTAINING OIL-SOLUBLE COMPONENT, FUNCTIONAL FOOD USING THE SAME, AND PACKAGED PRODUCT THEREOF ΙN Suzuki, Keiichi, Kanagawa, JAPAN Sasaki, Hidemi, Kanagawa, JAPAN Serizawa, Shinichiro, Kanagawa, JAPAN Arakawa, Jun, Kanagawa, JAPAN PA FUJIFILM CORPORATION, Minato-ku, Tokyo, JAPAN (non-U.S. corporation) ΡI US 20110104340 A1 20110505 ΑT US 2008-673977 A1 20080819 (12) WO 2008-JP65061 20080819 20100218 PCT 371 date 20070820 PRAI JP 2007-213712 JP 2007-230582 20070905 DT Utility FS APPLICATION LN.CNT 2345 INCL INCLM: 426/096.000 INCLS: 426/654.000; 426/590.000 426/096.000 NCL NCLM: NCLS: 426/590.000; 426/654.000 IPC IPCI A21D0002-16 [I,A]; A23L0002-52 [I,A] IPCR A21D0002-16 [I,A]; A23L0002-52 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 9 OF 27 USPATFULL on STN T.4 AN 2011:117391 USPATFULL METHODS OF USING KRILL OIL TO TREAT RISK FACTORS FOR ТΤ CARDIOVASCULAR, METABOLIC, AND INFLAMMATORY DISORDERS BRUHEIM, Inge, Volda, NORWAY ΙN Tilseth, Snorre, Bergen, NORWAY Cohn, Jeffery, Sydney, AUSTRALIA Griinari, Mikko, Espoo, FINLAND Mancinelli, Daniele, Orsta, NORWAY Hoem, Nils, Oslo, NORWAY Vik, Hogne, Eiksmarka, NORWAY Banni, Sebastiano, Calgliari, ITALY Aker BioMarine A.S.A., Oslo, NORWAY (non-U.S. corporation) PA ΡI US 20110104297 A1 20110505 AI US 2010-790575 A1 20100528 (12) Continuation-in-part of Ser. No. US 2008-57775, filed on 28 Mar 2008, RLI PENDING PRAI US 2007-975058P 20070925 (60)

US 2007-983446P 20071029 (60) US 2008-24072P 20080128 (61) US 2009-181743P 20090528 (61) US 2007-920483P 20070328 (60) DT Utility FS APPLICATION LN.CNT 2547 INCL INCLM: 424/522.000 INCLS: 426/002.000 424/522.000 NCL NCLM: 426/002.000 NCLS: TPC IPCI A61K0035-56 [I,A]; A61P0009-10 [I,A]; A61P0003-04 [I,A]; A61P0003-00 [I,A] IPCR A61K0035-56 [I,A]; A61P0003-00 [I,A]; A61P0003-04 [I,A]; A61P0009-10 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 10 OF 27 USPATFULL on STN T.4 AN 2011:97925 USPATFULL Methods for Treating Traumatic Brain Injury ΤI ΙN Bailes, Julian E., Morgantown, WV, UNITED STATES ΡI US 20110086914 A1 20110414 ΑT US 2010-904045 A1 20101013 (12) PRAI US 2009-251234P 20091013 (61) DT Utility FS APPLICATION LN.CNT 2356 INCL INCLM: 514/549.000 INCLS: 514/560.000 NCL NCLM: 514/549.000 NCLS: 514/560.000 A61K0031-232 [I,A]; A61K0031-20 [I,A]; A61P0025-00 [I,A] IPCI IPC IPCR A61K0031-232 [I,A]; A61K0031-20 [I,A]; A61P0025-00 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. L4ANSWER 11 OF 27 USPATFULL on STN AN 2011:92475 USPATFULL ΤI Docosahexaenoic Acid Gel Caps ΤN PANKER, Cynthia A., Jessup, MD, UNITED STATES Billard, Michael Ames, Laurel, MD, UNITED STATES Ryan, Alan, Ellicott City, MD, UNITED STATES Dangi, Bindi, Elkridge, MD, UNITED STATES ΡI US 20110082205 A1 20110407 ΑT US 2010-896763 A1 20101001 (12) PRAI US 2009-247944P 20091001 (61) DT Utility APPLICATION FS LN.CNT 2444 INCLM: 514/549.000 INCL NCL NCLM: 514/549.000 IPC IPCI A61K0031-232 [I,A]; A61P0003-06 [I,A] IPCR A61K0031-232 [I,A]; A61P0003-06 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 12 OF 27 USPATFULL on STN T.4 2010:256169 USPATFULL AN ΤI PHOSPHOLIPID AND PROTEIN TABLETS IN Tilseth, Snorre, Bergen, NORWAY Hoem, Nils, Oslo, NORWAY PA AKER BIOMARINE ASA, Oslo, NORWAY (non-U.S. corporation) ΡT US 20100227792 A1 20100909 ΑI US 2010-711822 A1 20100224 (12)

US 2009-155758P 20090226 (61) PRAT Utility DT FS APPLICATION LN.CNT 3112 INCLM: 514 2 INCL NCL NCLM: 514/005.500 NCLS: 514/691.000 A61K0038-02 [I,A] IPC IPCI IPCR A61K0038-02 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 13 OF 27 USPATFULL on STN T.4 AN 2010:255355 USPATFULL ΤТ LOW VISCOSITY PHOSPHOLIPID COMPOSITIONS ΤN Tilseth, Snorre, Bergen, NORWAY PA AKER BIOMARINE ASA, Oslo, NORWAY (non-U.S. corporation) ΡT US 20100226977 A1 20100909 US 2010-711553 20100224 (12) AI A1 Continuation-in-part of Ser. No. US 2008-201325, filed on 29 Aug 2008, RLI PENDING PRAI US 2009-155767P 20090226 (61) US 2007-968765P 20070829 (60) Utility DT APPLICATION FS LN.CNT 2394 INCL INCLM: 424/456.000 INCLS: 426/601.000; 426/417.000; 514/078.000 NCL NCLM: 424/456.000 NCLS: 426/417.000; 426/601.000; 514/078.000 TPC IPCI A61K0031-685 [I,A]; A23D0009-00 [I,A]; A23D0009-02 [I,A]; A61K0009-48 [I,A]; A61P0009-00 [I,A]; A61P0019-00 [I,A]; A61P0029-00 [I,A] IPCR A61K0031-685 [I,A]; A23D0009-00 [I,A]; A23D0009-02 [I,A]; A61K0009-48 [I,A]; A61P0009-00 [I,A]; A61P0019-00 [I,A]; A61P0029-00 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 14 OF 27 USPATFULL on STN L4 2010:228249 USPATFULL AN METHODS FOR IMPROVING COGNITIVE FUNCTION AND DECREASING HEART RATE ТΤ ΙN YURKO-MAURO, Karin, Silver Spring, MD, UNITED STATES PA MARTEK BIOSCIENCES CORPORATION, Columbia, MD, UNITED STATES (U.S. corporation) ΡT US 20100203123 Α1 20100812 ΑT US 2010-699009 Α1 20100202 (12) US 2009-149310P 20090202 (61) PRAT US 2009-183548P 20090602 (61) DT Utility APPLICATION FS LN.CNT 2358 INCL INCLM: 424/456.000 INCLS: 514/560.000; 514/549.000; 514/458.000 NCL NCLM: 424/456.000 514/458.000; 514/549.000; 514/560.000 NCLS: TPC IPCI A61K0009-64 [I,A]; A61K0031-20 [I,A]; A61K0031-22 [I,A]; A61K0031-355 [I,A]; A61P0025-00 [I,A]; A61P0009-00 [I,A] IPCR A61K0009-64 [I,A]; A61K0031-20 [I,A]; A61K0031-22 [I,A]; A61K0031-355 [I,A]; A61P0009-00 [I,A]; A61P0025-00 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 15 OF 27 USPATFULL on STN T.4

ΤТ PROCESS FOR PRODUCTION OF OMEGA-3 RICH MARINE PHOSPHOLIPIDS FROM KRILL ΤN Breivik, Harald, Porsgrunn, NORWAY ΡТ US 20100143571 A1 20100610 ΑI US 2007-515098 A1 20071115 (12) WO 2007-NO402 20071115 20100217 PCT 371 date PRAI US 2006-859289P 20061116 (60) DT Utility FS APPLICATION LN.CNT 537 INCLM: 426/643.000 INCL INCLS: 426/417.000; 554/021.000; 568/366.000; 536/020.000 NCL 426/643.000 NCLM: NCLS: 426/417.000; 536/020.000; 554/021.000; 568/366.000 IPCI A23L0001-325 [I,A]; A23K0001-10 [I,A]; A23K0001-18 [I,A]; TPC C11B0001-10 [I,A]; C07C0045-78 [I,A]; C08B0037-08 [I,A] IPCR A23L0001-325 [I,A]; A23K0001-10 [I,A]; A23K0001-18 [I,A]; C07C0045-78 [I,A]; C08B0037-08 [I,A]; C11B0001-10 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. T.4 ANSWER 16 OF 27 USPATFULL on STN AN 2009:109974 USPATFULL ΤI Polyunsaturated Fatty Acid-Containing Solid Fat Compositions and Uses and Production Thereof ΙN Namal Senanayake, S.P. Janaka, Lexington, KY, UNITED STATES Ahmed, Naseer, Lexington, KY, UNITED STATES Fichtali, Jaouad, Lexington, KY, UNITED STATES PA Martek Biosciences Corporation, Columbia, MD, UNITED STATES (U.S. corporation) A1 20090416 ΡT US 20090099260 US 2008-201728 A1 20080829 (12) ΑI US 2007-969536P PRAI 20070831 (60) Utility DT APPLICATION FS LN.CNT 2660 INCL INCLM: 514/560.000 INCLS: 426/601.000; 426/072.000 514/560.000 NCL NCLM: 426/072.000; 426/601.000 NCLS: IPC TPCT A61K0031-20 [I,A]; A23D0007-005 [I,A]; A23L0001-30 [I,A] IPCR A61K0031-20 [I,A]; A23D0007-005 [I,A]; A23L0001-30 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. T.4 ANSWER 17 OF 27 USPATFULL on STN AN 2009:67318 USPATFULL METHOD FOR MAKING KRILL MEAL ТΤ ΙN Tilseth, Snorre, Bergen, NORWAY Hostmark, Oistein, Loddefjord, NORWAY PA Aker BioMarine ASA, Oslo, NORWAY (non-U.S. corporation) ΡI US 20090061067 A1 20090305 AI US 2008-201325 A1 20080829 (12) PRAI US 2007-968765P 20070829 (60) DT Utility FS APPLICATION LN.CNT 2307 INCL INCLM: 426/602.000 INCLS: 426/417.000; 210/149.000; 426/480.000; 426/609.000; 426/648.000; 426/608.000; 366/145.000; 366/147.000 NCL NCLM: 426/602.000 210/149.000; 366/145.000; 366/147.000; 426/417.000; 426/480.000; NCLS: 426/608.000; 426/609.000; 426/648.000 IPC IPCI A23D0007-005 [I,A]; A23D0007-02 [I,A]; A23D0007-04 [I,A];

A23L0001-29 [I,A]; B01F0015-06 [I,A]; A23L0001-33 [I,A]; A23L0001-326 [I,A]; B01D0021-30 [I,A] TPCR A23D0007-005 [I,A]; A23D0007-02 [I,A]; A23D0007-04 [I,A]; A23L0001-29 [I,A]; A23L0001-326 [I,A]; A23L0001-33 [I,A]; B01D0021-30 [I,A]; B01F0015-06 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 18 OF 27 USPATFULL on STN L4AN 2006:254989 USPATFULL ΤT Natural astaxanthin extract reduces dna oxidation Chew, Boon P., Pullman, WA, UNITED STATES ΤN Park, Jean Soon, Pullman, WA, UNITED STATES ΡI US 20060217445 A1 20060928 A1 20040726 (10) ΑT US 2004-565717 WO 2004-US24314 20040726 20060123 PCT 371 date US 2003-490121P 20030725 (60) PRAI DT Utility APPLICATION FS LN.CNT 1366 INCL INCLM: 514/690.000 INCLS: 514/763.000; 514/560.000 NCL NCLM: 514/690.000 514/560.000; 514/763.000 NCLS: A61K0031-12 [I,A]; A61K0031-015 [I,A] IPC IPCI IPCR A61K0031-12 [I,A]; A61K0031-015 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. L4 ANSWER 19 OF 27 USPATFULL on STN 2006:227598 USPATFULL AN ΤI Preventive or remedy for arthritis ΤN Kamiya, Toshikazu, Ibaraki, JAPAN Nakagiri, Ryusuke, Chapel Hill, NC, UNITED STATES PA Kyowa Hakko Kogyo Co., Ltd., Tokyo, JAPAN, 100-8185 (non-U.S. corporation) ΡI US 20060193962 A1 20060831 AI US 2004-552526 A1 20040409 (10) WO 2004-JP5115 20040409 20051011 PCT 371 date PRAI JP 2003-107405 20030411 DT Utilitv FS APPLICATION LN.CNT 1047 INCL INCLM: 426/615.000 NCL NCLM: 426/615.000 A23L0001-212 [I,A] IPC IPCI A23L0001-212 [I,A]; A23K0001-14 [I,A]; A23K0001-16 [I,A]; IPCR A23L0001-30 [I,A]; A61K0031-7008 [I,A]; A61K0031-726 [I,A]; A61K0036-00 [I,A]; A61K0036-185 [I,A]; A61P0019-02 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 20 OF 27 USPATFULL on STN L4AN 2004:209092 USPATFULL ΤI Process for producing a plant extract containing plant powder Sakai, Yasushi, Tsukuba-shi, JAPAN ΤN Yokoo, Yoshiharu, Sagamihara-shi, JAPAN ΡI US 20040161524 A1 20040819 US 7521079 B2 20090421 US 2003-481519 A1 20031219 (10) ΑT WO 2002-JP6226 20020621 JP 2001-188480 20010621 PRAT DT Utility

FS APPLICATION LN.CNT 1479 INCLM: 426/655.000 INCL NCL NCLM: 426/655.000 426/433.000; 426/594.000; 426/597.000 NCLS: IPC [7] IPCI A23L0001-28 [ICM, 7] IPCI-2 A23L0001-28 [I,A] A23L0001-28 [I,A]; A23K0001-14 [I,A]; A23K0001-16 [I,A]; IPCR A23L0001-30 [I,A]; A61K0036-185 [I,A] ANSWER 21 OF 27 USPATFULL on STN T.4 AN 2004:209046 USPATFULL ΤI Preventives or remedies for arthritis ΤN Nakagiri, Rysuke, Tokyo, JAPAN Kamiya, Toshikazu, Tsuchiura-shi, JAPAN Suda, Toshio, Sunto-gun, JAPAN Miki, Ichiro, Mishima-shi, JAPAN ΡT US 20040161478 A1 20040819 US 2003-480044 A1 20031209 (10) ΑI WO 2002-JP5790 20020611 PRAI JP 2001-181947 20010615 JP 2002-70702 20020314 DT Utility APPLICATION FS LN.CNT 1301 INCL INCLM: 424/725.000 NCL NCLM: 424/725.000 IPC [7] IPCI A61K0035-78 [ICM, 7] IPCR A21D0002-36 [I,A]; A21D0013-08 [I,A]; A23K0001-14 [I,A]; A23K0001-16 [I,A]; A23L0001-30 [I,A]; A61K0036-185 [I,A]; A61P0019-02 [I,A]; A61P0029-00 [I,A] ANSWER 22 OF 27 USPATFULL on STN L4AN 2004:159281 USPATFULL ΤI Liver funcion protecting or ameliorating agent Sakai, Yasushi, Tsukuba-shi, JAPAN ΙN Kayahashi, Shun, Tsukuba-shi, JAPAN Hashizume, Erika, Tsukuba-shi, JAPAN Nakagiri, Ryusuke, Tokyo, JAPAN ΡI US 20040122085 A1 20040624 US 7332522 B2 20080219 A1 20031003 (10) ΑT US 2003-473867 WO 2002-JP3098 20020328 DT Utility APPLICATION FS LN.CNT 1146 INCLM: 514/470.000 INCL NCL NCLM: 514/457.000; 514/470.000 NCLS: 514/470.000; 549/283.000 IPC [7] A61K0031-365 [ICM, 7] IPCI IPCI-2 A61K0031-34 [I,A]; A61K0031-343 [I,A] A61K0031-34 [I,A]; A23L0001-30 [I,A]; A61K0031-343 [I,A]; IPCR A61K0031-365 [I,A]; A61K0031-366 [I,A]; A61P0001-16 [I,A]; C07D0307-88 [I,A]; C07D0311-76 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 23 OF 27 USPATFULL on STN L4 AN 2003:64375 USPATFULL ТΤ Processes for extracting carotenoids and for preparing feed materials

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Kagan, Michael, Jerusalem, ISRAEL
ΤN
       Braun, Sergei, Zur Hadassa, ISRAEL
ΡT
       US 20030044495
                           A1 20030306
       US 6818239
                           B2 20041116
       US 2002-172747
                           A1 20020617 (10)
ΑI
RLI
       Continuation of Ser. No. WO 2000-IL846, filed on 18 Dec 2000, UNKNOWN
PRAI
       GB 1999-30194
                                19991221
DT
       Utility
FS
       APPLICATION
LN.CNT 526
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INCL
NCL
       NCLM:
             426/429.000; 426/250.000
       NCLS:
             426/250.000; 426/253.000; 426/431.000; 426/478.000; 426/540.000
IPC
       [7]
       IPCI
              A23L0001-27 [ICM, 7]
       IPCI-2 A23L0001-28 [ICM, 7]; A23L0001-27 [ICS, 7]
              A23L0001-27 [I,A]; A23L0001-275 [I,A]; C07C0403-00 [I,A];
       IPCR
              C07C0403-24 [I,A]; C09B0061-00 [I,A]
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
T.4
     ANSWER 24 OF 27 USPATFULL on STN
AN
       2002:205917 USPATFULL
       Liver function protecting or improving agent
ΤΙ
ΙN
       Nakagiri, Ryusuke, Tsukuba-shi, JAPAN
       Kamiya, Toshikazu, Tsukuba-shi, JAPAN
       Hashizume, Erika, Tsukuba-shi, JAPAN
       Sakai, Yasushi, Inashiki-gun, JAPAN
       Kayahashi, Shun, Tsukuba-shi, JAPAN
                           A1 20020815
ΡT
       US 20020110605
ΑT
       US 2001-10154
                           A1 20011210 (10)
PRAI
       JP 2000-375510
                                20001211
DT
       Utility
       APPLICATION
FS
LN.CNT 1786
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INCL
NCL
       NCLM: 424/725.000
IPC
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              A61K0035-78 [ICM, 7]
       IPCI
              A21D0002-36 [I,A]; A21D0013-08 [I,A]; A23K0001-14 [I,A];
       IPCR
              A23K0001-16 [I,A]; A23L0001-212 [I,A]; A23L0001-30 [I,A];
              A61K0036-185 [I,A]; A61P0001-16 [I,A]
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 25 OF 27 USPAT2 on STN
L4
       2004:209092 USPAT2
AN
ТΤ
       Process for producing an extract of Hydrangea containing plant powder
ΤN
       Sakai, Yasushi, Tsukuba, JAPAN
       Yokoo, Yoshiharu, Sagamihara, JAPAN
       Kyowa Hakko Kogyo Co., Ltd., Tokyo, JAPAN (non-U.S. corporation)
PA
       US 7521079
ΡI
                           B2 20090421
       WO 2003000074
                                20030301
AI
       US 2002-481519
                                20020621 (10)
       WO 2002-JP6226
                                20020621
                                20031219 PCT 371 date
       JP 2001-188480
PRAI
                                20010621
DT
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FS
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LN.CNT 1371
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             426/655.000
       NCLS:
              426/433.000; 426/594.000; 426/597.000
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A23L0001-28 [ICM, 7] TPC IPCI IPCI-2 A23L0001-28 [I,A] A23L0001-28 [I,A]; A23K0001-14 [I,A]; A23K0001-16 [I,A]; TPCR A23L0001-30 [I,A]; A61K0036-185 [I,A] EXF 426/597; 426/433; 426/594 L4ANSWER 26 OF 27 USPAT2 on STN AN 2004:159281 USPAT2 ΤI Liver function protecting or ameliorating agent ΤN Sakai, Yasushi, Tsukuba, JAPAN Kayahashi, Shun, Tsukuba, JAPAN Hashizume, Erika, Tsukuba, JAPAN Nakagiri, Ryusuke, Tokyo, JAPAN PA Kyowa Hakko Kogyo Co., Ltd., Tokyo, JAPAN (non-U.S. corporation) US 7332522 ΡI B2 20080219 WO 2002080904 20021017 US 2002-473867 20020328 (10) ΑI WO 2002-JP3098 20020328 PCT 371 date 20031003 PRAI JP 2001-106600 20010405 DT Utility FS GRANTED LN.CNT 1099 INCL INCLM: 514/457.000 INCLS: 514/470.000; 549/283.000 NCL NCLM: 514/457.000; 514/470.000 NCLS: 514/470.000; 549/283.000 IPC IPCI A61K0031-365 [ICM, 7] IPCI-2 A61K0031-34 [I,A]; A61K0031-343 [I,A] IPCR A61K0031-34 [I,A]; A23L0001-30 [I,A]; A61K0031-343 [I,A]; A61K0031-365 [I,A]; A61K0031-366 [I,A]; A61P0001-16 [I,A]; C07D0307-88 [I,A]; C07D0311-76 [I,A] EXF 549/283; 549/290; 549/307; 549/289; 514/457; 514/470 CAS INDEXING IS AVAILABLE FOR THIS PATENT. L4ANSWER 27 OF 27 USPAT2 on STN AN 2003:64375 USPAT2 Processes for extracting carotenoids and for preparing feed materials ΤT ΤN Kagan, Michael, Jerusalem, ISRAEL Braun, Sergei, Zur Hadassa, ISRAEL ΡA Fermentron Ltd., Jerusalem, ISRAEL (non-U.S. corporation) ΡI US 6818239 B2 20041116 ΑT US 2002-172747 20020617 (10) RLI Continuation of Ser. No. WO 2000-IL846, filed on 18 Dec 2000 PRAI GB 1999-30194 19991221 DT Utility FS GRANTED LN.CNT 501 INCLM: 426/429.000 INCL INCLS: 426/431.000; 426/478.000; 426/250.000; 426/253.000; 426/540.000 NCL NCLM: 426/429.000; 426/250.000 NCLS: 426/250.000; 426/253.000; 426/431.000; 426/478.000; 426/540.000 IPC [7] A23L0001-27 [ICM, 7] IPCI IPCI-2 A23L0001-28 [ICM, 7]; A23L0001-27 [ICS, 7] IPCR A23L0001-27 [I,A]; A23L0001-275 [I,A]; C07C0403-00 [I,A]; C07C0403-24 [I,A]; C09B0061-00 [I,A] EXF 426/807; 426/250; 426/253; 426/635; 426/425; 426/429; 426/430; 426/431; 426/478; 426/540; 424/439; 424/451 CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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C M CT M T	1	UL U	тU	Numerical Property Search Feature
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Enter SET DETAIL ON to see search term postings or to view search error messages that display as 0* with SET DETAIL OFF.

=> s krill and oil and cook? and dry?(p)krill and krill(p)meal and extract?(p)krill(p)oil 0* FILE ADISNEWS 0* FILE ANTE 0* FILE AQUALINE 0* FILE BIOENG 0* FILE BIOTECHABS 0* FILE BIOTECHDS 0* FILE BIOTECHNO 0* FILE CEABA-VTB 0* FILE CIN 29 FILES SEARCHED... 0* FILE FOMAD 0* FILE FROSTI FILE IFIPAT 3 0* FILE KOSMET 0* FILE NTIS 0* FILE PASCAL 9 FILE USPATFULL 50 FILES SEARCHED... 3 FILE USPAT2 0* FILE WATER

FILE WPIDS 4 4 FILE WPINDEX 5 FILES HAVE ONE OR MORE ANSWERS, 56 FILES SEARCHED IN STNINDEX L1OUE KRILL AND OIL AND COOK? AND DRY? (P) KRILL AND KRILL (P) MEAL AND EXTRACT? (P)KRILL(P)OIL => file ifipat uspatfull uspat2 COST IN U.S. DOLLARS SINCE FILE TOTAL SESSION ENTRY FULL ESTIMATED COST 1.48 1.72 FILE 'IFIPAT' ENTERED AT 14:52:35 ON 29 MAY 2012 COPYRIGHT (C) 2012 IFI CLAIMS(R) Patent Services (IFI) FILE 'USPATFULL' ENTERED AT 14:52:35 ON 29 MAY 2012 CA INDEXING COPYRIGHT (C) 2012 AMERICAN CHEMICAL SOCIETY (ACS) FILE 'USPAT2' ENTERED AT 14:52:35 ON 29 MAY 2012 CA INDEXING COPYRIGHT (C) 2012 AMERICAN CHEMICAL SOCIETY (ACS) => s l1 L2 15 L1 => dup rem 12 PROCESSING COMPLETED FOR L2 L3 13 DUP REM L2 (2 DUPLICATES REMOVED) => d 13 1-13 ANSWER 1 OF 13 IFIPAT COPYRIGHT 2012 IFI on STN DUPLICATE 1 L3 12887434 IFIPAT; IFIUDB; IFICDB AN SOLVENT-FREE PROCESS FOR OBTAINING PHOSPHOLIPIDS AND NEUTRAL ENRICHED ΤI KRILL OILS ΙN Sclabos Katevas Dimitri (CL); Toro Guerra Raul R (CL); Chiong Lay Mario M (CL) THAROS LTD CL PA LONZA LTD CH (50035)ΡI US 20110224450 A1 20110915 AI US 2011-96644 20110428 (13) RLI WO 2009-IB7269 20091030 CONTINUATION-IN-PART PENDING FΙ US 20110224450 20110915 DT Utility; Patent Application - First Publication CHEMICAL FS APPLICATION Entered STN: 21 Oct 2011 ED Last Updated on STN: 13 Jan 2012 CLMN 25 ANSWER 2 OF 13 USPAT2 on STN L3 2007:272601 USPAT2 AN ТΤ Gels, gel composites, and gel articles Chen, John Y., Hillsborough, CA, UNITED STATES ΤN ΡA Applied Elastomerics, Inc., South San Francisco, CA, UNITED STATES (U.S. corporation) ΡI US 7930782 B2 20110426 20070605 (11) AI US 2007-810584 Continuation-in-part of Ser. No. US 2007-787257, filed on 12 Apr 2007, RLI Pat. No. US 7661164 Continuation-in-part of Ser. No. US 2004-912464, filed on 4 Aug 2004, Pat. No. US 7226484 Continuation-in-part of Ser.

No. US 2002-420489, filed on 21 Apr 2002, Pat. No. US 7222380 Continuation-in-part of Ser. No. US 2003-420492, filed on 21 Apr 2003, Pat. No. US 7344568 Continuation-in-part of Ser. No. US 2000-721213, filed on 21 Nov 2000, Pat. No. US 6867253 DT Utility FS GRANTED LN.CNT 5886 INCL INCLM: 005/655.500 INCLS: 005/636.000; 005/652.000; 005/654.000; 005/909.000; 602/041.000; 602/061.000; 602/062.000; 602/063.000; 623/016.110; 623/020.140; 623/021.110; 623/023.400; 623/027.000; 623/033.000; 623/036.000; 524/270.000; 524/284.000; 524/490.000; 524/491.000; 524/549.000; 524/571.000; 524/575.000; 521/050.000; 521/054.000; 521/139.000; 521/140.000; 521/148.000 005/655.500; 525/240.000 NCL NCLM: 005/636.000; 005/652.000; 005/654.000; 005/909.000; 521/050.000; NCLS: 521/054.000; 521/139.000; 521/140.000; 521/148.000; 524/270.000; 524/284.000; 524/490.000; 524/491.000; 524/549.000; 524/571.000; 524/575.000; 602/041.000; 602/061.000; 602/062.000; 602/063.000; 623/016.110; 623/020.140; 623/021.110; 623/023.400; 623/027.000; 623/033.000; 623/036.000 IPC IPCI C08L0023-16 [I,A] IPCI-2 B29C0067-20 [I,A]; B60R0021-26 [I,A]; A61F0002-80 [I,A]; B60K0028-00 [I,A]; A47C0007-00 [I,A] IPCR B29C0067-20 [I,A]; A47C0007-00 [I,A]; A61F0002-80 [I,A]; B60K0028-00 [I,A]; B60R0021-26 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. L3 ANSWER 3 OF 13 USPATFULL on STN 2010:256169 USPATFULL AN ΤI PHOSPHOLIPID AND PROTEIN TABLETS ΤN Tilseth, Snorre, Bergen, NORWAY Hoem, Nils, Oslo, NORWAY AKER BIOMARINE ASA, Oslo, NORWAY (non-U.S. corporation) PA ΡI US 20100227792 A1 20100909 ΑI US 2010-711822 A1 20100224 (12) PRAI US 2009-155758P 20090226 (61) DT Utility APPLICATION FS LN.CNT 3112 INCL INCLM: 514 2 NCL NCLM: 514/005.500 NCLS: 514/691.000 IPC IPCI A61K0038-02 [I,A] A61K0038-02 [I,A] IPCR CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 4 OF 13 USPATFULL on STN L3 2010:255355 USPATFULL AN LOW VISCOSITY PHOSPHOLIPID COMPOSITIONS ΤΙ ΙN Tilseth, Snorre, Bergen, NORWAY PA AKER BIOMARINE ASA, Oslo, NORWAY (non-U.S. corporation) ΡI US 20100226977 A1 20100909 US 2010-711553 ΑT A1 20100224 (12) Continuation-in-part of Ser. No. US 2008-201325, filed on 29 Aug 2008, RLI PENDING PRAI US 2009-155767P 20090226 (61) US 2007-968765P 20070829 (60) DT Utility FS APPLICATION LN.CNT 2394 INCL INCLM: 424/456.000

INCLS: 426/601.000; 426/417.000; 514/078.000 NCL NCLM: 424/456.000 NCLS: 426/417.000; 426/601.000; 514/078.000 IPC IPCI A61K0031-685 [I,A]; A23D0009-00 [I,A]; A23D0009-02 [I,A]; A61K0009-48 [I,A]; A61P0009-00 [I,A]; A61P0019-00 [I,A]; A61P0029-00 [I,A] IPCR A61K0031-685 [I,A]; A23D0009-00 [I,A]; A23D0009-02 [I,A]; A61K0009-48 [I,A]; A61P0009-00 [I,A]; A61P0019-00 [I,A]; A61P0029-00 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 5 OF 13 IFIPAT COPYRIGHT 2012 IFI on STN DUPLICATE 2 L3 AN 12061067 IFIPAT; IFIUDB; IFICDB ΤТ METHOD FOR MAKING KRILL MEAL Hostmark Oistein (NO); Tilseth Snorre (NO) ΤN Aker BioMarine ASA NO (79725) PA РT US 20090061067 A1 20090305 US 2008-201325 ΑI 20080829 (12)US 2007-968765P PRAI 20070829 (Provisional) US 20090061067 20090305 FΙ DT Utility; Patent Application - First Publication FS CHEMICAL APPLICATION Entered STN: 10 Mar 2009 ED Last Updated on STN: 9 Apr 2009 CLMN 51 L3 ANSWER 6 OF 13 USPATFULL on STN AN 2008:312554 USPATFULL ΤI BIOEFFECTIVE KRILL OIL COMPOSITIONS ΤN Bruheim, Inge, Volda, NORWAY Griinari, Mikko, Espoo, FINLAND Tilseth, Snorre, Bergen, NORWAY Banni, Sebastiano, Cagliari, ITALY Cohn, Jeffrey Stuart, Camperdown, AUSTRALIA Mancinelli, Daniele, Orsta, NORWAY ΡA AKER BIOMARINE ASA, Oslo, NORWAY (non-U.S. corporation) ΡI US 20080274203 A1 20081106 US 2008-57775 A1 20080328 (12) ΑT PRAI US 2007-920483P 20070328 (60) US 2007-975058P 20070925 (60) US 2007-983446P 20071029 (60) US 2008-24072P 20080128 (61) DT Utility FS APPLICATION LN.CNT 2199 INCL INCLM: 424/522.000 INCLS: 514/121.000; 514/078.000; 514/114.000; 426/601.000 NCL NCLM: 424/522.000 NCLS: 426/601.000; 514/078.000; 514/114.000; 514/121.000 IPC IPCI A61K0035-56 [I,A]; A61K0031-661 [I,A]; A61K0031-685 [I,A]; A61P0003-02 [I,A]; A23D0009-00 [I,A]; A61K0031-66 [I,A] IPCR A61K0035-56 [I,A]; A23D0009-00 [I,A]; A61K0031-66 [I,A]; A61K0031-661 [I,A]; A61K0031-685 [I,A]; A61P0003-02 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. L3 ANSWER 7 OF 13 USPATFULL on STN AN 2007:272601 USPATFULL ΤI Gels, gel composites, and gel articles ΤN Chen, John Y., Hillsborough, CA, UNITED STATES РT US 20070238835 A1 20071011 US 7930782 B2 20110426

AI US 2007-810584 A1 20070605 (11)

Continuation-in-part of Ser. No. US 2007-787257, filed on 12 Apr 2007, RLI PENDING Continuation-in-part of Ser. No. US 2004-912464, filed on 4 Aug 2004, GRANTED, Pat. No. US 7226484 Continuation-in-part of Ser. No. US 2003-613567, filed on 2 Jul 2003, GRANTED, Pat. No. US 7093316 Continuation-in-part of Ser. No. US 2003-420489, filed on 21 Apr 2003, GRANTED, Pat. No. US 7222380 Continuation-in-part of Ser. No. US 2003-420487, filed on 21 Apr 2003, GRANTED, Pat. No. US 7193002 Continuation-in-part of Ser. No. US 2003-420488, filed on 21 Apr 2003, GRANTED, Pat. No. US 7134929 Continuation-in-part of Ser. No. US 2003-420490, filed on 21 Apr 2003, GRANTED, Pat. No. US 7105607 Continuation-in-part of Ser. No. US 2003-420491, filed on 21 Apr 2003, GRANTED, Pat. No. US 7093599 Continuation-in-part of Ser. No. US 2003-420492, filed on 21 Apr 2003, PENDING Continuation-in-part of Ser. No. US 2003-420493, filed on 21 Apr 2003, GRANTED, Pat. No. US 7067583 Continuation-in-part of Ser. No. US 2004-896047, filed on 22 Jul 2004, PENDING Continuation-in-part of Ser. No. US 2002-273828, filed on 17 Oct 2002, GRANTED, Pat. No. US 6909220 Continuation-in-part of Ser. No. US 2002-334542, filed on 31 Dec 2002, GRANTED, Pat. No. US 7159259 Continuation-in-part of Ser. No. US 2002-299073, filed on 18 Nov 2002, ABANDONED Continuation-in-part of Ser. No. US 2002-199364, filed on 20 Jul 2002, GRANTED, Pat. No. US 6794440 Continuation-in-part of Ser. No. US 2002-199361, filed on 20 Jul 2002, GRANTED, Pat. No. US 7134236 Continuation-in-part of Ser. No. US 2002-199362, filed on 20 Jul 2002, GRANTED, Pat. No. US 7208184 Continuation-in-part of Ser. No. US 2002-199363, filed on 20 Jul 2002, GRANTED, Pat. No. US 7108873 Continuation-in-part of Ser. No. US 2000-721213, filed on 21 Nov 2000, GRANTED, Pat. No. US 6867253 Continuation-in-part of Ser. No. US 1998-130545, filed on 8 Aug 1998, GRANTED, Pat. No. US 6627275 Continuation-in-part of Ser. No. US 1999-230940, filed on 3 Feb 1999, GRANTED, Pat. No. US 6161555 Continuation-in-part of Ser. No. US 1997-863794, filed on 27 May 1997, GRANTED, Pat. No. US 6117176 PRAI JP 2003-204428 20030731 WO 1994-US4278 19940419 19940627 WO 1994-US7314 DTUtility FS APPLICATION LN.CNT 5757 INCL INCLM: 525/240.000 NCL 005/655.500; 525/240.000 NCLM: NCLS: 005/636.000; 005/652.000; 005/654.000; 005/909.000; 521/050.000; 521/054.000; 521/139.000; 521/140.000; 521/148.000; 524/270.000; 524/284.000; 524/490.000; 524/491.000; 524/549.000; 524/571.000; 524/575.000; 602/041.000; 602/061.000; 602/062.000; 602/063.000; 623/016.110; 623/020.140; 623/021.110; 623/023.400; 623/027.000; 623/033.000; 623/036.000 IPC C08L0023-16 [I,A] IPCI IPCI-2 B29C0067-20 [I,A]; B60R0021-26 [I,A]; A61F0002-80 [I,A]; B60K0028-00 [I,A]; A47C0007-00 [I,A] B29C0067-20 [I,A]; A47C0007-00 [I,A]; A61F0002-80 [I,A]; IPCR B60K0028-00 [I,A]; B60R0021-26 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. L3 ANSWER 8 OF 13 USPAT2 on STN 2004:24434 USPAT2 AN ΤI Gelatinous food elastomer compositions and articles for use as fishing bait IN Chen, John Y., Pacifica, CA, UNITED STATES ΡA Applied Elastomerics, Inc., South San Francisco, CA, UNITED STATES (U.S. corporation) ΡT B2 20070424 US 7208184 ΑI US 2002-199362 20020720 (10)

DT Utility FS GRANTED LN.CNT 4932 INCLM: 426/001.000 INCL INCLS: 043/042.000; 043/042.240; 424/084.000 NCL NCLM: 426/001.000 NCLS: 043/042.000; 043/042.240; 424/084.000 IPC IPCI A23L0001-00 [ICM, 7] IPCI-2 A23L0001-00 [I,A] A23L0001-00 [I,A]; A01K0085-01 [I,A]; A01K0097-04 [I,A] IPCR 426/1; 043/42; 043/42.24; 424/84 EXF CAS INDEXING IS AVAILABLE FOR THIS PATENT. L3 ANSWER 9 OF 13 USPAT2 on STN AN 2004:24385 USPAT2 ΤI Gelatinous food elastomer compositions and articles ΤN Chen, John Y., Pacifica, CA, UNITED STATES PA Applied Elastomerics, Inc., South San Francisco, CA, UNITED STATES (U.S. corporation) 20060919 ΡT US 7108873 В2 ΑI US 2002-199363 20020720 (10) RLI Continuation-in-part of Ser. No. US 2001-721213, filed on 21 Nov 2001, Pat. No. US 6867253 Continuation-in-part of Ser. No. US 2001-896047, filed on 30 Jun 2001, PENDING Continuation-in-part of Ser. No. US 1999-421886, filed on 5 Oct 1999, ABANDONED Continuation-in-part of Ser. No. US 1999-285809, filed on 1 Apr 1999, ABANDONED Continuation-in-part of Ser. No. US 1999-274498, filed on 23 Mar 1999, Pat. No. US 6420475 Continuation-in-part of Ser. No. US 1998-130545, filed on 8 Aug 1998, Pat. No. US 6627275 Continuation-in-part of Ser. No. US 1997-984459, filed on 3 Dec 1997, Pat. No. US 6324703 Continuation-in-part of Ser. No. WO 1997-US17534, filed on 30 Sep 1997, Pat. No. WO 6161555 Continuation-in-part of Ser. No. US 1997-909487, filed on 12 Jul 1997, Pat. No. US 6050871 Continuation-in-part of Ser. No. US 1997-863794, filed on 27 May 1997, Pat. No. US 6117176 Continuation-in-part of Ser. No. US 1996-719817, filed on 30 Sep 1996, Pat. No. US 6148830 Continuation-in-part of Ser. No. US 1996-665343, filed on 17 Jun 1996, PENDING Continuation-in-part of Ser. No. US 1996-612586, filed on 8 Mar 1996, Pat. No. US 6552109 Continuation-in-part of Ser. No. US 1995-581191, filed on 29 Dec 1995, Pat. No. US 5760117 Continuation-in-part of Ser. No. US 1995-581188, filed on 29 Dec 1995, ABANDONED Continuation-in-part of Ser. No. US 1995-581125, filed on 29 Dec 1995, Pat. No. US 5962572 Continuation-in-part of Ser. No. US 1994-288690, filed on 11 Aug 1994, Pat. No. US 5633286 Continuation-in-part of Ser. No. WO 1994-US7314, filed on 27 Jun 1994, Pat. No. WO 5868597 Continuation-in-part of Ser. No. WO 1994-US4278, filed on 19 Apr 1994, Pat. No. WO 6033383 DT Utility FS GRANTED LN.CNT 3521 INCL INCLM: 426/001.000 INCLS: 426/573.000; 524/505.000 NCL NCLM: 426/001.000; 424/439.000 426/573.000; 524/505.000 NCLS: A61K0047-00 [ICM, 7] IPC IPCI IPCI-2 A01K0097-04 [I,A]; A23L0001-05 [I,A] IPCR A01K0097-04 [I,A]; A23L0001-05 [I,A]; A23L0001-317 [I,A]; A23L0001-325 [I,A]; A61K0047-00 [I,A] EXF 524/505; 424/486; 426/1; 426/648; 426/656; 426/534; 426/555; 426/573 CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 10 OF 13 IFIPAT COPYRIGHT 2012 IFI on STN ЪЗ

AN

04308583 IFIPAT; IFIUDB; IFICDB

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ΤТ
      Protein and lipid sources for use in aquafeeds and animal feeds and a
      process for their preparation; Subjecting oilseed to heat treatment to
      reduce concentration of antinutritional components to obtain heat-treated
      seed; dehulling seed to produce a meat fraction, a hull fraction or a
      mixture; cold pressing to obtain plant oils andm meals
ΙN
      Shand Ian (CA); Cairns Robert E (CA); Higgs David (CA)
PA
      Canada Fisheries and Oceans Minister of CA (51835)
ΡI
      US 6955831
                      В2
                          20051018 (CITED IN 002 LATER PATENTS)
      US 20030072866 A1 20030417
      US 2002-76499
                          20020219
ΑT
                                    (10)
      US 2000-566728
                          20000509 CONTINUATION-IN-PART
RLT
                                                           ABANDONED
PRAT
     CA 2001-2334745
                           20010213
      WO 2001-CA663
                           20010508
      CA 2001-2351903
                           20010626
FΙ
      US 6955831
                          20051018
      US 20030072866
                          20030417
DT
      Utility; Granted Patent - Utility, with Pre-Grant Publication
FS
      CHEMICAL
      GRANTED
ΕD
      Entered STN: 19 Oct 2005
      Last Updated on STN: Jan 2011
MRN
      012837
             MFN: 0842
CLMN
     32
     ANSWER 11 OF 13 USPATFULL on STN
L3
AN
       2004:24434 USPATFULL
ΤI
       Gelatinous food elastomer compositions and articles for use as fishing
       bait
       Chen, John Y., Pacifica, CA, UNITED STATES
ΤN
ΡI
                           A1 20040129
       US 20040018272
       US 7208184
                           B2 20070424
       US 2002-199362
                           A1 20020720 (10)
ΑI
DT
       Utility
       APPLICATION
FS
LN.CNT 4354
INCL
       INCLM: 426/001.000
NCL
       NCLM:
             426/001.000
             043/042.000; 043/042.240; 424/084.000
       NCLS:
TPC
       [7]
              A23L0001-00 [ICM, 7]
       IPCI
       IPCI-2 A23L0001-00 [I,A]
       IPCR
              A23L0001-00 [I,A]; A01K0085-01 [I,A]; A01K0097-04 [I,A]
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 12 OF 13 USPATFULL on STN
L3
AN
       2004:24385 USPATFULL
ТΤ
       Gelatinous food elastomer compositions and articles
       Chen, John Y., Pacifica, CA, UNITED STATES
ΙN
                           A1 20040129
PT
       US 20040018223
       US 7108873
                           В2
                               20060919
AI
       US 2002-199363
                           A1 20020720 (10)
DT
       Utility
FS
       APPLICATION
LN.CNT 3229
       INCLM: 424/439.000
INCL
NCL
       NCLM:
              426/001.000; 424/439.000
       NCLS:
             426/573.000; 524/505.000
IPC
       [7]
       IPCI
              A61K0047-00 [ICM,7]
       IPCI-2 A01K0097-04 [I,A]; A23L0001-05 [I,A]
       IPCR
              A01K0097-04 [I,A]; A23L0001-05 [I,A]; A23L0001-317 [I,A];
              A23L0001-325 [I,A]; A61K0047-00 [I,A]
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CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 13 OF 13 USPATFULL on STN L3 2003:165578 USPATFULL AN ΤT Process for making dried powdery and granular krill ΙN Yoshitomi, Bunji, Tokyo, JAPAN Shigematsu, Yoshiaki, Tokyo, JAPAN PA NIPPON SUISAN KAISHA, LTD., Tokyo, JAPAN (non-U.S. corporation) ΡI US 20030113432 A1 20030619 US 2002-283063 A1 20021030 (10) ΑI Continuation of Ser. No. US 2001-807953, filed on 25 Apr 2001, PENDING RLI JP 1998-311730 PRAI 19981102 DTUtility FS APPLICATION LN.CNT 481 INCL INCLM: 426/643.000 NCL NCLM: 426/643.000 IPC [7] IPCI A23L0001-325 [ICM, 7] IPCR A23B0004-03 [I,A]; A23L0001-325 [I,A]; A23L0001-326 [I,A]; A23L0001-33 [I,A]

=> d 13 13 kwic

- L3 ANSWER 13 OF 13 USPATFULL on STN
- TI Process for making dried powdery and granular krill
- AB A dried powdery and granular krill product containing all components of krill. The proteolytic enzymes originally contained in krill materials are perfectly disabled. The product is produced by a process including only heating as means for denaturing protein and disabling the proteolytic enzymes originally contained in krill materials. The product is produced by a process including no chemicals treatment to remove water and disable or inactivate the proteolytic enzymes in any production steps, and generating no wastewater. The production process comprises the steps of lightly dehydrating krill, coarsely crushing the krill, and drying the coarsely crushed krill under heating. Thus, water is removed from the krill by only heating, and degradation of the lipid in the krill product is prevented without using an anti-oxidant. Application fields are enlarged and the preservation characteristic is improved. The so-called zero-emission. .
- SUMM [0002] The present invention relates to a dried powdery and granular krill product which contains all components of krill and in which lipid degradation is sufficiently prevented with no need of an anti-oxidant.
- SUMM [0004] Krill are animal plankton living primarily in the Arctic and Antarctic Oceans, and about 80 kinds of krill have been known up to date. Of those many kinds of krill, Antarctic Krill (Euphasia superba) living in the Antarctic Ocean are found in abundance as one of natural resources. Therefore, survey of the resource and development of the method of catching the krill have been extensively conducted in the period of 1970 to 1985, including studies for developing methods of processing the krill to be useful in practical applications.
- SUMM [0005] Krill are comparable to fish, flesh and fowl in point of nutritive value, but there are several problems in processing the krill for practical applications. One of the problems is that krill lose freshness in short time. If krill are left to stand after being caught, the heads and chests of the krill start changing into black color in 1-2 hours even at a low atmospheric temperature of about 0° C. Further, shells of the heads and chests of krill are so vulnerable to external pressure that the krill are easily broken down upon impacts applied at the time of catching, whereupon the enzymes

present in the internal organs flow out and decompose muscles. Those phenomena occur under actions of the enzymes present in krill. It is thought that tyrosinase is responsible for the former color-changing phenomenon, and protease is responsible for the latter muscle-decomposing. . .

- SUMM [0006] Accordingly, those enzymes require to be disabled or inactivated when processing krill. In other words, it has been required immediately after catching krill to quickly freeze the krill down to below -40° C., thereby inactivating the enzymes, or to heat the krill up to above 80° C., thereby disabling the enzymes, followed by preserving the krill.
- SUMM [0007] Known krill products include raw frozen and peeled krill products which are subjected to quick freezing and then preserved in a frozen condition, boiled krill products which are heated and then preserved in a frozen condition, and krill meal which is heated and dried and then preserved at the normal temperature. The following Tables 1 and 2 list classifications of those products depending on how krill are processed, and features and points to be improved of the products.
- SUMM . . Japan, the product price greatly depends on the transportation cost. There is hence a desire for extracting excellent characteristics of krill more efficiently and realizing krill products having a higher value added.

TABLE 1

| Pı | rocessing | | Processing | Object | Product Exa | imples |
|----------------|---------------------------------------|--|---|---|---|---------------|
| Pi | uick freez:
reserve in
ondition | 2. | Inactivate | enzymes | Raw frozen
stripped kr | |
| He | eating, Pre | | Disable en: | zymes | Boiled kril | .1 |
| He
Pi
te | eating &
reserve at
emperature | drying, | | Disable en: | zymes | Krill meal |
| SUMM | • • • | Points to be | e improved | | | |
| | | Products hav
taste and fe
raw krill. | | | s
l
state.
apon
ality | |
| enzym
prote | | Heating disa
enzymes and
protein stak
meat-like fe | makes
ble to give | Flavor and taste components
flow out during boiling. Cold
chain is required because of
high water content. | | |
| Krill | meal | Heating disa
enzymes and
protein stak
can be store
normal temp
of low water | ables
makes
ole. Meal
ed at
. because | Digestibility
protein den
heating. Wat
components
stickwater. | ty lowers du
aturation du
ter-soluble | iring |
| SUMM | [0010] Ja | | | nt Publicatio | on No. 57-11 | 876 discloses |

SUMM [0010] Japanese Unexamined Patent Publication No. 57-11876 discloses a method of impeding activity of the proteolytic enzymes in krill and utilizing the krill as protein materials. With the disclosed method, a krill paste is degenerated with alcohol to effect fixation (denaturation) of protein and degeneration of the enzymes at the same time. The processed krill paste is then washed with water to remove alcohol. The disclosed method however has the following problems.

- SUMM [0013] 3. Polar lipid is removed together with alcohol during washing with water. Most of the lipid in krill is phospholipid and is rich in polyunsaturated fatty acids (PUFAs). Thus these PUFAs are removed.
- SUMM . . . square. The shrimp materials thus processed are dried under heating to thereby provide dried shrimp granules. Considering specific properties of krill, however, it is inferred that even if krill are dried under heating after being processed in a similar manner as in the prior art, ground krill are very difficult to dry into a satisfactory condition.
- SUMM [0018] From intensive studies, the inventors found that when krill are processed in a similar manner as in the prior art, lipid, protein and water contained in the krill are brought into an emulsified state, and the processed krill are very difficult to dry even with a heating and drying machine. Such a difficulty is related to the fact that most of the lipid in krill is phospholipid, as described above, and therefore emulsification is further increased. In other words, water in the krill is stabilized in structure with emulsification and becomes still harder to evaporate under heating.
- SUMM [0019] In addition, when krill are crushed into the form of ground meat, the proteolytic enzymes present in the internal organs of the krill develop activity, and a temperature rise during the grinding process increases the activity of those enzymes. As a consequence, proteolysis in the krill is promoted and specific taste is deteriorated.
- SUMM [0021] An object of the present invention is therefore to effectively utilize krill as one of valuable aquatic resources, and to provide a dried powdery and granular krill product and a method of producing the dried powdery and granular krill product, which contains all components of krill and has a good preservation ability while activity of the enzymes in the krill is totally disabled.
- SUMM [0022] The present invention resides in a dried powdery and granular krill product that contains all components of krill. Because of containing all components of krill, the present product has a function capable of sufficiently preventing degradation of the lipid in the krill product without using an anti-oxidant. In the dried powdery and granular krill product, the proteolytic enzymes originally contained in krill materials are perfectly disabled. Accordingly, the present invention also resides in a dried powdery and granular krill product which contains all components of krill and in which the proteolytic enzymes originally contained in krill materials are perfectly disabled. The present product is produced by a process including only heating as means for denaturing protein and disabling the proteolytic enzymes originally contained in krill materials. Accordingly, the present invention further resides in a dried powdery and granular krill product which contains all components of krill, in which the proteolytic enzymes originally contained in krill materials are perfectly disabled, and which is produced by a process including only heating as means for denaturing protein and disabling the proteolytic enzymes originally contained in krill materials.
- SUMM [0023] The dried powdery and granular krill product of the present invention is produced by a process including no chemicals treatment to remove water and disable or. . . the proteolytic enzymes in any production steps, and generating no wastewater. The production process comprises the steps of lightly dehydrating krill, coarsely crushing the krill, and drying the coarsely crushed krill under heating.
- SUMM [0024] The dried powdery and granular krill product of the present invention is subjected to no chemical treatment using chemicals, etc. in any production steps, and is. . Also, there is no step in the production process in which wastewater is generated. Thus, water is removed from the krill by only heating. Moreover, application fields are enlarged and the preservation characteristic is improved. The so-called zero-emission method and product, . .
- SUMM [0025] The production method of the present invention comprises steps of

removing seawater from krill, coarsely crushing the krill, and drying the coarsely crushed krill under heating. In the conventional process of producing krill meal, krill are first boiled in water in the same amount as the krill, and are then subjected to separation into solid and liquid components. The solid component is heated and dried using a. . . drier. The liquid component obtained from the solid/liquid separation is called stickwater and preserved separately. For this reason, the conventional krill meal contains less water-soluble components than the krill product of the present invention, and therefore has disadvantages in not providing satisfactory flavor and taste in the extracted form, . . the conventional production process is disadvantageous in that protein is excessively denatured by heating applied in both the boiling and heating/drying steps, and digestibility of the product is reduced.

- DRWD [0026] FIG. 1 is a graph showing activity of the proteolytic enzymes remaining in raw krill and the product of the present invention; and DETD [0028] There are 80 or more kinds of krill as described above, but the kind of krill used in the present invention is not restricted. In addition to krill, mysids are also usable.
- DETD [0029] Krill primarily used in an embodiment are Antarctic Krill (Euphasia superba) which have been employed in industrial fields.
- DETD [0031] Krill used as materials are put into a fish tank at once after being caught. The krill are then put in a dehydrator to remove seawater, etc. attaching to the krill surfaces. The type of the dehydrator is not particularly restricted, but outer shells of krill are so fragile that the shells are easily broken down under pressure of 40-140 g/cm.sup.2 and the internal components flow. . . Therefore, the type of the dehydrator is preferably selected so that an excessive physical load will not be applied to krill.
- DETD [0032] The dehydrated krill are chopped to improve thermal efficiency in the heating and drying process. The type of a machine used for chopping the krill is not particularly restricted. The grain size of the chopped krill is selected to a coarsely crushed state, i.e., about 1.5-2.5 cm square, at which outer shells and muscular tissues of the krill materials remain. This process can be performed with, e.g., a known mincing apparatus, which is usually employed for grinding meat.
- DETD [0033] The chopped krill are dried under heating. The type of a machine for use in this process is also not particularly restricted. While a known heating and drying machine such as a steam type disk dryer, for example, can be used, the machine is preferably adjustable in heating time, heating temperature, degree of agitation, and so forth. Because the internal components of krill as one of natural resources change depending on the season, it is desired to adjust the parameters of the machine in match with the change of the internal components of krill for obtaining products with constant quality.
- DETD [0034] The heating time and the heating temperature are set to such an extent that the muscular protein of krill and the proteolytic enzymes in krill are denatured and degenerated under heating, and that the water content is reduced down to below 10% from a point of ensuring good preservation. It is important that the heating and drying process is not performed at overly high temperatures and for an overly long time, and is performed at the necessary. . . values to satisfy the above-described conditions. Excessive heating lowers digestibility due to extreme denaturation, reduces astaxanthin, natural dye, present in krill, reduces vitamins, and oxidizes lipid. On the other hand, if heating is insufficient, activity of the proteolytic enzymes in krill remains, which leads to a deterioration of product quality. If the water content is over ten and several percents, the krill product gathers mold during preservation.
- DETD [0035] The dried krill are very fragile, including the shells, and therefore can be easily crushed any desired grain size.

DETD [0036] The krill product of the present invention can be used as a main material of feed for cultured fish in place of. . .

- DETD . . . above in connection with the prior art is attributable to crushing of raw materials into the form of ground meat, krill materials are first chopped into pieces having a size of 20-30% of the body length (about 1.5-2.5 cm square) and are then put into a heating and drying machine in the present invention. As a result, the krill materials are avoided from being emulsified and the drying efficiency is enhanced. Further, strong activity of the proteolytic enzymes present in the internal organs of krill is suppressed and an adverse influence upon flavor and taste of the krill product is reduced. In addition, the chopped krill do not adhere to the heating surface and can be heated appropriately, thus greatly contributing to improvement of product quality.
- DETD [0038] Moreover, since the dried krill product obtained in accordance with the method of the present invention has a large grain size and maintains a fair part of shapes of the krill materials, it is also possible to produce products utilizing the shapes of the krill materials advantageously. Additionally, the dried krill can be simply crushed into a desired grain size as required.
- DETD [0040] FIG. 1 shows comparatively activity of the proteolytic enzymes remaining in raw krill and the krill product of the present invention. DETD . . as a substrate. As will be seen from FIG. 1, the activity of the remaining proteolytic enzymes in the raw krill is increased with lapse of the reaction time, while the activity of the remaining
- proteolytic enzymes in the krill product of the present invention is hardly changed. This suggests that the proteolytic enzymes remain not alive in the krill product of the present invention and they are perfectly disabled in the production process, and that a possibility of quality deterioration of the krill product during the preservation is low.
- DETD [0042] Preservation characteristics of the krill product of the present invention will be described with reference to Tables 3 and 4 below.
- DETD [0043] For comparison, the results listed in Table 3 were obtained by preparing two groups of the krill product of the present invention, in one of which ethoxyquin that is most generally used as an anti-oxidant in meal, etc. was added to the krill product and in the other of which no ethoxyquin was added, and then measuring a change of product quality by. . .
- DETD [0045] There are several indexes indicating a degree of lipid degradation. About the lipid in krill, particularly, the krill lipid having been extracted and refined, it is known that, during the preservation, a peroxide value hardly increases and only a carbonyl value increases. In other words, it is pointed out that degradation of the krill lipid differs in creation of oxides and progress rate of the decomposing reaction from those in general fish oil, etc.
- TABLE 3

Acid value

| | with | Pero | xid | e value | Carbonyl | l value |
|----------|---------|------|-----|---------|----------|---------|
| no anti- | anti- | | | wit | h | with |
| oxidant | oxidant | no. | • | • | | |

DETD . . from Table 4, a phenomenon of the lipid degrading at apparently different rates during the preservation was found between the krill product of the present invention and a control prepared by perfectly removing all the water-soluble components originally present in krill from the krill product of the present invention. Although the material responsible for the above phenomenon is not yet known, it is believed that the water-soluble components originally present in krill have some anti-oxidizing action. For this reason, in the krill product of

the present invention which contains all the components of krill in an enriched condition, lipid degradation can be prevented satisfactorily without using any anti-oxidant.

TABLE 4

| DETD
DETD | Peroxide value Carbonyl
[0048] 1. Process Flow Including Plant for Drying Krill
[0049] An outline of the process flow is as shown in FIG. 2. Krill
materials are first conveyed by a krill supply apparatus from a fish
tank to a material tank, and are then supplied to a dehydrator in a
proper lot. The use of a dehydrator basically intends to remove seawater
contained in the krill materials. Since it is expected that the amount
of water contained in krill varies depending on the materials, a
diaphragm is adjusted to provide a proper dehydration rate, taking into
account the performance are then supplied to a drier. The
materials are boiled in the drier under heating with vapor, followed by
further drying. At the time when reaching a predetermined water
content, the drying is stopped and a resulting dried semifinished
product is ejected. The dried semifinished product is conveyed to a
product tank, |
|--------------|--|
| DETD | [0050] The conventional production process for krill meal is
represented by raw krill→boiling→centrifugal separation
or solid/liquid separation→extraction of
solid→ drying→crushing→packaging. The liquid
component was removed in the centrifugal separation step, and the useful |
| DETD | <pre>components of krill contained in the liquid component were discarded.
It can be said from one aspect that the krill meal was a product
resulted from drying the sludge.
[0051] By contrast, the process flow for producing the krill product
of the present invention is represented by raw krill→removal of
water attached to krill→boiling→ drying→crushing
→packaging. The centrifugal separation step is not included. In
the boiling and drying steps, the enzymes in krill are disabled and
the krill components are stabilized through thermal degeneration.
Thus, the components originally contained in the krill are all kept in</pre> |
| DETD | the product without being discarded externally. An apparatus for
implementing the above process is featured in omitting a step of
squeezing boiled krill using a decanter or a press. The krill
drying apparatus used in the present invention differs from the
conventional meal producing apparatus in that a cooker and a drier
are combined in an integral structure.
[0053] Table 5 lists component analytical values of the krill product
of the present invention. For comparison, Table 5 also lists component
analytical values of the krill meal produced by the conventional
process. In particular, the krill product of the present invention
contains free amino acids as much as more than twice the amount
contained in the conventional krill meal. The free amino acids
deeply take part in developing flavor and taste of the product when |

eaten, attractant of feed. . . DETD [0054] Since the squeezing step subsequent to boiling of the krill materials is omitted, the components developing flavor and taste are not lost and the krill product of the present invention has good flavor. Further, the production process of the present invention generates no appreciable wastewater and provides a high yield.

TABLE 5

Krill meal Product of invention

Water

| Coarse protein | 64.0 | 65.1 |
|-------------------|-------|--------|
| (Free amino acid) | (2.9) | (7.54) |
| Coarse fat | 7.0 | 7.0 |
| Coarse | | |

- DETD [0055] According to the present invention, a method is provided which can effectively utilize krill, as one of important aquatic resources, in a perfect manner without any loss due to efflux of krill components. The dried powdery and granular krill product obtained by the present invention contains all the components originally contained in the krill, and strong activity of the enzymes specific to the krill is disabled. Therefore, the krill product of the present invention can be widely applied to not only the feed industry, but also the food industry.
- CLM What is claimed is: 1. A dried powdery and granular krill product containing all components of krill.
- CLM What is claimed is: 2. A dried powdery and granular krill product according to claim 1, wherein the proteolytic enzymes originally contained in krill materials are perfectly disabled.
- CLM What is claimed is: 3. A dried powdery and granular krill product according to claim 1 or 2, wherein said product is produced by a process including only heating as means for denaturing protein and disabling the proteolytic enzymes originally contained in krill materials.
- CLM What is claimed is: 4. A dried powdery and granular krill product according to claim 1, 2 or 3, wherein said product is produced by a process including no chemicals treatment.
- CLM What is claimed is: 5. A dried powdery and granular krill product according to any one of claims 1 to 4, wherein said product is produced by a process comprising the steps of lightly dehydrating krill, coarsely crushing the krill, and drying the coarsely crushed krill under heating.

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Bruheim et al.Serial No.:12/057,775Filed:March 28, 200Entitled:BIOEFFECTIV

Bruheim et al.Art12/057,775ExaMarch 28, 2008CoBIOEFFECTIVE KRIL OIL COMPOSITIONS

Art Unit: 1651 Examiner: Ware Confirmation: 1945

RESPONSE TO OFFICE ACTION MAILED JANUARY 6, 2012

EFS WEB FILED

Commissioner for Patents P.O. Box 1450 Alexandria, Virginia 22313-1450

Examiner Ware:

This communication is responsive to the Office Action mailed January 6, 2012. The Commissioner is hereby authorized to charge any fees during the entire pendency of this application, including fees due under 37 C.F.R. §§ 1.16 and 1.17 that may be required, including any required extension of time fees, or credit any overpayment to Deposit Account 50-4302, referencing Attorney Docket No. **NATNUT-14409/US-5/ORD**. This paragraph is intended to be a CONSTRUCTIVE PETITION FOR EXTENSION OF TIME in accordance with 37 C.F.R. § 1.136(a)(3).

-1-

IN THE CLAIMS

1. (Withdrawn) A composition comprising:

from about 3% to 10% ether phospholipids on a w/w basis;

from about 35% to 50% non-ether phospholipids on w/w basis, so that the total amount of ether phospholipids and non-ether phospholipids in the composition is from about 48% to 60% on a w/w basis;

from about 20% to 45% triglycerides on a w/w basis; and from about 400 to about 2500 mg/kg astaxanthin.

2. (Withdrawn) The composition of Claim 1, wherein said ether phospholipids are selected from the group consisting of alkylacylphosphatidylcholine, lyso-alkylacylphosphatidylcholine, alkylacylphosphatidylethanolamine, and combinations thereof.

3. (Withdrawn) The composition of Claim 1, wherein said ether lipids are greater than 90% alkylacylphosphatidylcholine.

4. (Withdrawn) The composition of Claim 1, wherein said non-ether phospholipids are selected from the group consisting of phosphatidylcholine, phosphatidylserine, phosphatidylethanolamine and combinations thereof.

5. (Withdrawn) The composition of Claim 1, wherein said composition comprises a blend of lipid fractions obtained from *Euphausia superba*.

6. (Withdrawn) The composition of Claim 1, wherein said composition comprises from about 25% to 30% omega-3 fatty acids as a percentage of total fatty acids and wherein from about 80% to 90% of said omega-3 fatty acids are attached to said phospholipids.

7. (Withdrawn) A capsule containing the composition of Claim 1.

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8. (Withdrawn) A composition comprising:
from about 3% to 10% ether phospholipids on a w/w basis; and
from about 400 to about 2500 mg/kg astaxanthin.

9. (Withdrawn) The composition of Claim 8, further comprising from about 35% to 50% non-ether phospholipids on w/w basis, so that the total amount of ether phospholipids and non-ether phospholipids in the composition is from about 38% to 60% on a w/w basis.

10. (Withdrawn) The composition of Claim 8, further comprising from about 20% to 45% triglycerides on a w/w basis.

11. (Withdrawn) The composition of Claim 8, wherein said ether phospholipids are selected from the group consisting of alkylacylphosphatidylcholine, lyso-alkylacylphosphatidylcholine, alkylacylphosphatidylethanolamine, and combinations thereof.

12. (Withdrawn) The composition of Claim 11, wherein said ether lipids are greater than 90% alkylacylphosphatidylcholine.

13. (Withdrawn) The composition of Claim 8, wherein said non-ether phospholipids are selected from the group consisting of phosphatidylcholine, phosphatidylserine, phosphatidylethanolamine and combinations thereof.

14. (Withdrawn) The composition of Claim 8, wherein said composition comprises a blend of lipid fractions obtained from *Euphausia superba*.

15. (Withdrawn) The composition of Claim 10, wherein said composition comprises from about 25% to 30% omega-3 fatty acids as a percentage of total fatty acids and wherein from about 80% to 90% of said omega-3 fatty acids are attached to said phospholipids.

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16. (Withdrawn) A capsule containing the composition of Claim 8.

17. (Withdrawn) A blended krill oil composition comprising: from about 45% to 55% w/w phospholipids; from about 20% to 45% w/w triglycerides; and from about 400 to about 2500 mg/kg astaxanthin.

18. (Withdrawn) The composition of Claim 17, wherein said blended krill oil product comprises a blend of lipid fractions obtained from *Euphausia superba*.

19. (Withdrawn) The composition of Claim 17, wherein said composition comprises from about 25% to 30% omega-3 fatty acids as a percentage of total fatty acids and wherein from about 80% to 90% of said omega-3 fatty acids are attached to said phospholipids.

20. (Withdrawn) A *Euphausia superba* krill oil composition comprising:

from about 3% to about 10% w/w ether phospholipids;

from about 27% to 50% w/w non-ether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w;

from about 20% to 50% w/w triglycerides;

from about 400 to about 2500 mg/kg astaxanthin; and

from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

21. (Withdrawn) A dietary supplement comprising encapsulated *Euphausia superba* krill oil comprising from about 3% to about 10% w/w ether phospholipids; from about 27% to 50% w/w non-ether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w; from about 20% to 50% w/w triglycerides; from about 400 to about 2500 mg/kg astaxanthin; and from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

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22. (Withdrawn) A method of making a *Euphausia superba* krill oil composition comprising: contacting *Euphausia superba* with a polar solvent to provide a polar extract comprising phospholipids;

contacting *Euphasia superba* with a neutral solvent to provide a neutral extract comprising triglycerides and astaxanthin;

combining said polar extract and said neutral extract to provide *Euphausia superba* krill oil from about 3% to about 10% w/w ether phospholipids; from about 27% to 50% w/w nonether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w; from about 20% to 50% w/w triglycerides; from about 400 to about 2500 mg/kg astaxanthin; and from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

23. (Withdrawn) The method of Claim 22, further comprising the step of encapsulating the *Euphausia superba* krill oil.

24. (Withdrawn) A *Euphausia superba* krill oil produced by the method of Claim 22.

25. (Withdrawn) A method of producing a dietary supplement comprising;

contacting *Euphausia superba* with a polar solvent to provide an polar extract comprising phospholipids;

contacting *Equphasia superba* with a neutral solvent to provide a neutral extract comprising triglycerides and astaxanthin;

combining said polar extract and said neutral extract to provide *Euphausia superba* krill oil from about 3% to about 10% w/w ether phospholipids; from about 27% to 50% w/w nonether phospholipids so that the amount of total phospholipids in the composition is from about 30% to 60% w/w; from about 20% to 50% w/w triglycerides; from about 400 to about 2500 mg/kg astaxanthin; and from about 20% to 35% omega-3 fatty acids as a percentage of total fatty acids in said composition, wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids;

encapsulating said Euphausia superba krill oil.

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26. (Withdrawn) A composition comprising at least 65% (w/w) of phospholipids, said phospholipids characterized in containing at least 35% omega-3 fatty acid residues.

27. (Withdrawn) The composition according to claim 26, wherein the composition is derived from a marine or aquatic biomass.

28. (Withdrawn) The composition according to claim 26, wherein the composition is derived from krill.

29. (Withdrawn) The composition of Claim 26, wherein said composition comprises less than2% free fatty acids.

30. (Withdrawn) The composition of Claim 26, wherein said composition comprises less than 10% triglycerides.

31. (Withdrawn) The composition of Claim 26, wherein said phospholipids comprise greater than 50% phosphatidylcholine.

32. (Withdrawn) The composition of Claim 26, wherein the composition comprises at least 500 mg/kg astaxanthin esters.

33. (Withdrawn) The composition of Claim 26, wherein the composition comprises at least 500 mg/kg astaxanthin esters and at least 36% (w/w) omega-3 fatty acids.

34. (Withdrawn) The composition of Claim 26, wherein the composition comprises less than about 0.5g/100g total cholesterol.

35. (Withdrawn) The composition of Claim 26, wherein the composition comprises less than about 0.45% arachidonic acid (w/w).

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36. (Withdrawn) A krill lipid extract comprising at least 500 mg/kg astaxanthin esters and at least 36% (w/w) omega-3 fatty acids.

37. (Withdrawn) A krill lipid extract comprising at least 100 mg/kg astaxanthin esters, at least 20% (w/w) omega-3 fatty acids, and less than about 0.45% arachidonic acid (w/w).

38. (Withdrawn) A method comprising administering the composition of Claim 1 to a subject in an amount effective for reducing insulin resistance, reducing inflammation, improving blood lipid profile and reducing oxidative stress.

39. (Withdrawn) A krill lipid extract comprising greater than about 80% triglycerides and greater than about 90 mg/kg astaxanthin esters.

40. (Withdrawn) The krill lipid extract of Claim 39, characterized in containing from about 5% to about 15% omega-3 fatty acid residues.

41. (Withdrawn) The krill lipid extract of Claim 39, characterized in containing less than about 5% phospholipids.

42. (Withdrawn) The krill lipid extract of Claim 39, characterized in comprising from about 5% to about 10% cholesterol.

43. (Withdrawn) A krill meal composition comprising less than about 50g/kg total fat.

44. (Withdrawn) The krill meal composition of Claim 43 comprising from about 5 to about 20 mg/kg astaxanthin esters.

45. (Withdrawn) The krill meal composition of Claim 43 comprising greater than about 65% protein.

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46. (Withdrawn) The krill meal composition of Claim 43 comprising greater than about 70% protein.

47. (Withdrawn) An animal feed comprising the krill meal of Claim 46.

48. (Withdrawn) A method of increasing flesh coloration in an aquatic species comprising feeding said aquatic species a composition comprising the krill meal of Claim 46.

49. (Withdrawn) A method of increasing growth and overall survival rate of aquatic species by feeding the krill meal of Claim 46.

50. (Currently amended) A method of producing krill oil comprising:

- a) <u>cooking and drying krill to providing provide cooked and dried krill meal; and</u>
- b) extracting <u>a krill</u> oil from said <u>cooked and dried</u> krill meal.

51. (Cancelled)

52. (Original) The method of Claim 50, wherein said krill meal is stored prior to said extraction step.

53. (Original) The method of Claim 50, wherein said extracting step comprises extraction by supercritical fluid extraction.

54. (Currently amended) The method of Claim 53, wherein said supercritical fluid extraction is a two step process comprising a first extraction step with carbon dioxide and from 1 to 10% of a co-solvent and a second extraction step with carbon dioxide and from 10-30% of a co-solvent, wherein said co-solvent in said first and second extraction steps is a C_1 - C_3 monohydric alcohol.

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55. (Currently amended) An <u>krill</u> oil produced by the method of claim 50.

56. (Withdrawn) A method of production of krill oil comprising:

a) providing fresh krill;

b) treating said fresh krill to denature lipases and phospholipases in said fresh krill to provide a denatured krill product; and

c) extracting oil from said denatured krill product.

57. (Withdrawn) The method of claim 56 in which the denaturation step comprises heating of said fresh krill.

58. (Withdrawn) The method of claim 56 in which the denaturation step comprises heating said fresh krill after grinding.

59. (Withdrawn) The method of claim 56, further comprising storing said denatured krill product at room temperature or below between the denaturation step and the extraction step.

60. (Withdrawn) The method of claim 56, wherein the enzyme denaturation step is achieved by application of heat.

61. (Withdrawn) The method of claim 56, wherein the extraction step comprises use of supercritical carbon dioxide, with or without use of a polar modifier.

62. (Withdrawn) The method of claim 56, wherein the extraction step comprises the use of ethanol.

63. (Withdrawn) The method of Claim 56, wherein the extraction step comprises ethanol extraction followed by acetone to precipitation of phospholipids.

64. (Withdrawn) The method of Claim 56, wherein said denatured krill product is a meal.

65. (Withdrawn) Oil produced by the method of Claim 56.

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66. (Withdrawn) A composition comprising an oil extracted from krill having a phosphatidylcholine content of greater then about 50% (w/w).

67. (Withdrawn) The composition of Claim 66, wherein said oil has a phosphatidylcholine content of greater then about 70% (w/w).

68. (Withdrawn) The composition of Claim 66, wherein said oil has a phosphatidylcholine content of greater then about 80% (w/w).

69. (Withdrawn) The composition of Claim 66, wherein said composition comprises less than 2% free fatty acids.

70. (Withdrawn) The composition of Claim 66, wherein said composition comprises less than 10% triglycerides.

71. (Withdrawn) The composition of Claim 66, wherein the composition comprises at least 500 mg/kg astaxanthin esters.

72. (Withdrawn) The composition of Claim 66, wherein the composition comprises less than about 0.45% arachidonic acid (w/w).

73. (Withdrawn) A composition comprising odorless krill oil.

74. (Withdrawn) The composition of Claim 73, wherein said odorless krill oil comprises less than about 10 mg/kg (w/w) trimethylamine.

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75. (Withdrawn) An odorless krill oil produced by the method comprising:

extracting a neutral krill oil from a krill oil containing material by supercritical fluid extraction to provide a deodorized krill material, wherein said neutral krill oil contains odor causing compounds and

extracting a polar krill oil from said deodorized krill material by supercritical fluid extraction with a polar entrainer to provide an essentially odorless krill oil.

76. (Withdrawn) A composition comprising krill oil containing less than about 70 micrograms/kilogram (w/w) astaxanthin esters.

77. (Withdrawn) The composition of claim 76, comprising less than about 50 micrograms/kilogram (w/w) astaxanthin esters.

78. (Withdrawn) The composition of claim 76, comprising less than about 20 micrograms/kilogram (w/w) astaxanthin esters.

79. (Withdrawn) The composition of claim 76, comprising less than about 5 micrograms/kilogram (w/w) astaxanthin esters.

80. (Withdrawn) A krill oil produced by the process comprising:

pumping fresh krill from a trawl onto a ship, heating the krill to provide a krill material, and extracting oil from the krill material.

81. (Withdrawn) A method of reducing diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction or hepatic steatosis comprising:

in a subject exposed to a high fat diet, administering to said subject exposed to a high fat diet an effective amount of a krill oil composition under conditions such that a condition selected from the group consisting of diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction and hepatic steatosis is reduced.

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82. (Withdrawn) The method of Claim 81, wherein said effective amount of a krill oil composition is from 0.2 grams to 10 grams of said krill oil composition.

83. (Withdrawn) The method of Claim 81, wherein said krill oil composition comprises: from about 45% to 55% w/w phospholipids; from about 35% to 45% w/w triglycerides; and from about 400 to about 2500 mg/kg astaxanthin.

84. (Withdrawn) The method of Claim 81, wherein said krill oil composition comprises a blend of lipid fractions obtained from *Euphausia superba*.

85. (Withdrawn) A method of reducing diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction or hepatic steatosis comprising in a subject consuming a high fat diet or a normal fat diet:

administering to said subject consuming a high fat diet or a normal fat diet an effective amount of a krill oil composition under conditions such that a condition selected from the group consisting of diet-induced hyperinsulinemia, insulin insensitivity, muscle mass hypertrophy, serum adiponectin reduction and hepatic steatosis is reduced.

86. (Withdrawn) A method of inducing diuresis in a subject comprising: administering to said subject an effective amount of a krill oil composition under conditions such that diuresis is induced.

- 87. (Withdrawn) A method of increasing muscle mass in a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that muscle mass is increased.
- 88. (Withdrawn) A method of decreasing protein catabolism in a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that protein catabolism is decreased.

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- 89. (Withdrawn) A method of decreasing lipid content in the heart of a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that lipid content in the heart of the subject is decreased.
- 90. (Withdrawn) A method of decreasing lipid content in the liver of a subject, comprising: administering to said subject an effective amount of a krill oil composition under conditions such that lipid content in the liver of the subject is decreased.

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REMARKS

Claims 50 and 51-55 are pending following entry of this amendment. Claim 52 has been cancelled without prejudice. Claims 50, 54 and 55 have been amended. Support for the amendments may be found in the specification, for example at page 42, lines 1-4, and the in the claims as originally filed among other places. No new matter has been added. All amendments and cancellation of claims are made without acquiescing to any of the Examiner's arguments or rejections, and solely for the purpose of expediting the patent application process in a manner consistent with the PTO's Patent Business Goals (PBG), and without waiving the right to prosecute the cancelled claims (or similar claims) in the future.

The following rejections are at issue:

- 1. Claims 50-55 are rejected as being indefinite;
- 2. Claims 50-53 and 55 are rejected as anticipated by Japanese Abstract 04-057853;
- Claim 54 is rejected as being obvious over Japanese Abstract 04-057853 in view of Kamiya (US 20060193962).

These rejections are addressed in order below.

1. The claims are definite

Claims 50-55 are rejected as being indefinite. The claims have been amended to correct the antecedent basis issues notes by the Examiner for oil and co-solvent as well as the second step. Applicants respectfully request that this rejection be withdrawn.

2. The claims are not anticipated

Claims 50-53 and 55 are rejected as anticipated by Japanese Abstract 04-057853. Applicants respectfully disagree. Nevertheless, Applicants have amended the claims to clarify that the meal is a cooked and dried meal. The Japanese abstract discloses a protease treated and mechanically ground composition:

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Krill shells are treated with a protease to decompose the protein in the shells and the treatment product is filtered. The residue of filtration is dried to give treated shells having a water content of 6-8% and a mean particle size of 200 μ m or lower. The treated shells are put into an extraction vessel 5.

There is no cooking step as currently claimed. Furthermore, the purpose of the process is to extract a coloring pigment from krill shells: "To prepare a reddish orange coloring matter having a high safety in a high concn. by extracting, with CO2 in a supercritical state, krill shells of which the protein has been decomposed by a protease." Applicants respectfully submit that the alleged prior art process, which uses only krill shells, is substantially different from the claimed process which uses the krill organism. The prior art process, which utilizes shells, will not produce a krill oil as claimed. Applicants respectfully request that the rejection be withdrawn and the claims passed to allowance.

3. The claims are not obvious

Claim 54 is rejected as being obvious over Japanese Abstract 04-057853 in view of Kamiya (US 20060193962). Applicants respectfully disagree. In any event, the amendments to the claims address the rejection. Kamiya does not cure the deficiencies noted for Japanese Abstract 04-057853 above. Namely, Kamiya does not teach extraction of krill oil from a cooked and dried krill meal. Accordingly, the combined references do not teach each element of the claims. Any prima facie case of obviousness allegedly established by the Examiner is therefore rebutted. Applicants respectfully request that the rejection be withdrawn and the claims passed to allowance.

CONCLUSION

If a telephone interview would aid in the prosecution of this application, the Examiner is encouraged to call the undersigned collect at (608) 662-1277.

Dated: _____ April 4, 2012___

/J. Mitchell Jones/

John Mitchell Jones Registration No. 44,174

Casimir Jones, S.C. 2275 Deming Way, Suite 310 Middleton, WI, 53562 (608) 662-1277

-15- RIMFROST EXHIBIT 1024 page 0716

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| EFS ID: | 12463979 | | | | | |
| Application Number: | 12057775 | | | | | |
| International Application Number: | | | | | | |
| Confirmation Number: | 1945 | | | | | |
| Title of Invention: | BIOEFFECTIVE KRILL OIL COMPOSITIONS | | | | | |
| First Named Inventor/Applicant Name: | Inge Bruheim | | | | | |
| Customer Number: | 72960 | | | | | |
| Filer: | John Mitchell Jones/Vickie Hoeft | | | | | |
| Filer Authorized By: | John Mitchell Jones | | | | | |
| Attorney Docket Number: | NATNUT-14409/US-5/ORD | | | | | |
| Receipt Date: | 04-APR-2012 | | | | | |
| Filing Date: | 28-MAR-2008 | | | | | |
| Time Stamp: | 14:32:46 | | | | | |
| Application Type: | Utility under 35 USC 111(a) | | | | | |

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| | Multipart Description/PDF files in .zip description | | | | | | |
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7,775 | | ing Date
28/2008 | To be Mailed |
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| | SEARCH FEE
(37 CFR 1.16(k), (i), c | or (m)) | N/A | | N/A | | N/A | | | N/A | |
| | EXAMINATION FE
(37 CFR 1.16(o), (p), c | | N/A | | N/A | | N/A | | | N/A | |
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| Application Number | | 12057775 |
|----------------------------|--|-----------------------|
| Filing Date | | 2008-03-28 |
| First Named Inventor Ingel | | Bruheim |
| Art Unit | | 1651 |
| Examiner Name Ware | | |
| Attorney Docket Number | | NATNUT-14409/US-5/ORD |

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⁴ Kind of document by the appropriate symbols as indicated on the document under WIPO Standard ST.16 if possible. ⁵ Applicant is to place a check mark here if English language translation is attached. | | | | | | | | |

| | Application Number | | 12057775 | |
|--|------------------------|--------|-----------------------|--|
| | Filing Date | | 2008-03-28 | |
| INFORMATION DISCLOSURE | First Named Inventor | Inge E | Bruheim | |
| STATEMENT BY APPLICANT
(Not for submission under 37 CFR 1.99) | Art Unit | | 1651 | |
| | Examiner Name | Ware | | |
| | Attorney Docket Number | | NATNUT-14409/US-5/ORD | |

CERTIFICATION STATEMENT

Please see 37 CFR 1.97 and 1.98 to make the appropriate selection(s):

That each item of information contained in the information disclosure statement was first cited in any communication from a foreign patent office in a counterpart foreign application not more than three months prior to the filing of the information disclosure statement. See 37 CFR 1.97(e)(1).

OR

That no item of information contained in the information disclosure statement was cited in a communication from a foreign patent office in a counterpart foreign application, and, to the knowledge of the person signing the certification after making reasonable inquiry, no item of information contained in the information disclosure statement was known to any individual designated in 37 CFR 1.56(c) more than three months prior to the filing of the information disclosure statement. See 37 CFR 1.97(e)(2).

See attached certification statement.

The fee set forth in 37 CFR 1.17 (p) has been submitted herewith.

X A certification statement is not submitted herewith.

SIGNATURE

A signature of the applicant or representative is required in accordance with CFR 1.33, 10.18. Please see CFR 1.4(d) for the form of the signature.

| Signature | /J. Mitchell Jones/ | Date (YYYY-MM-DD) | 2012-03-21 |
|------------|---------------------|---------------------|------------|
| Name/Print | J. Mitchell Jones | Registration Number | 44174 |

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1.発明の名 | 林遵 | *=*** | 昭和
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19日本国特許广

公開特許公報

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| 21特願昭 | 50-49765 | | | | | |
| 22出願日 | 昭50. (1975) | 4.25 | | | | |
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井物産館内 電話 (
<u>九 義 男</u>
) 氏名 朝 内 <i>鬼</i>
50 049765 | 591) 0261番
—— <u> </u> | | |

明 継 書

1.発明の名称

(2400) 氏名

金丸特許事務所内(6145)

南氷洋産オキアミの加工方法

5

2.特許請求の範囲

南水洋産オキアミに外力を加えることにより肉部は穀中に残し肝臓を体外に押し出して除去し、 肝臓を除去したオキアミを採肉装置により処理す ることにより肉部を得、ついでこの肉部を加熱し て熟発問蛋白を取得することを特徴とする南氷洋 産オキアミの加工方法。

3発明の詳細な説明

本発明は南氷洋産オキアミの加工方法に関する。 南氷洋産オキアミ(Euphausia superba,以下単 にオキアミという)は、南氷洋に生棲するアミの 一種であり小エビに似た体長約15~50mの大 型の動物性ブランクトンである。

オキアミの資源量は10~15億トンと推定さ れており、成長が速く2年で成熟するので、全世 界の漁獲量に近い 5,000 ~ 7,000 万トン程度を 毎年漁獲しても再生産に悪影響を及ぼさない程の 真大な量があると言われており、従つてこれを動 物性タンパク質値として利用するための加工方法 の開発が急がれている。

オキアミを食品として利用するために現在実用 化され、あるいは開発されている加工方法として は(1) 設付きのまま煮熟し、ついで冷凍あるいは乾 録する方法、(2) 設付のまま細断した後圧搾破で肉 部 むよび内臓を同時に搾り出し、これを加熱して 熱碟固蛋白質等を得る方法、(3) オキアミの肝臓の強 力な消化酵素を利用してオキアミ・ソリユーブル を製造する方法、および(4)魚類の FPC (Fieh Protein Concentrate) と同様の方法でイソプロ バノール等を密剤として使用してオキアミ FPC を製造する方法、等がある。これらの方法中(2)の 熱礙固蛋白を得る方法はオキアミの蛋白質を能率 よく採取しかつ大量に処理するのに適した方法で ある。

しかしながら、これらのいずれの加工方法においても処理工程中に内臓を除去することが困難で

あるため、最終製品中に内蔵液が混入することを 避けられず、特に肝臓にはエク味や苦味があり油 帽分も含有されており、また強力な消化酵素が含 有されているので、甲酸類等有の炎白で残快味の あるうま味の硬退した製品しか得られず、また消 化酵素の作用により歩留りが低下し、かつ冷凍保 存を行つた場合においても製品の品質が低下して いくという欠点がある。

本 発明は前記(2)の方法によりオキアミの蛋白度 を 能率 よく採取しかつ大量に処理して熱鍵固蛋白 を 製造するに あたり、 肝縁液を予め除去すること により、 美味 でかつ 長期保存性のあるオキアミ熱 瞬間蛋白を製造する方法を提供することを目的と する。

つぎに肝臓液を除去した数付オキアミを、その ままあるいは予め道径5 m目程度の内税機で破砕 した淡、例えば圧搾機で圧搾して数部を除去し粥 状义は濃厚なジュース状の内部を得る。この場合、 求用した圧搾酸の短額によつては肉部に若干の彼 離な数が遅入することがあるので、必要に応じて そのままの状態あるいは滑水を加えた後沪過を行 なつて混入している数を除去する。

つぎにかく得られた粥状义は濃厚なジュース状 の内部を、90 C程度に加熱することによりオキ すこの蛋白質の大部分を占める熟薬固性蛋白を熟 要固させ、痰黄色の魂肉状憂菌蛋白と痰黄色透明 な液部(プロス)とを生成させる。ついでこれを そのまま沪過するかあるいは遠心分離して凝固蛋 白とプロスとに分離する。かぐ得られた疑問蛋白 は肝臓凝を含有していないためエグ味や苦味がな く甲酸類の身肉に特有のうま味を有しており、肝 臓液味去処理を行わない方法で得られた数固蛋白 に酸べて極めて美味である。プロスは必要に応じ て更に速心分離または沪過を行ないついで滅圧機 特開 昭51-125774(2)

オキアミを大型トロール旗船で捕獲し、船上に 引湯げて僕み違ねた場合、オキアミの鮮皮が優め て良好な場合でも、約40m程隻覺み重ねるだけ で下積みのオキアミの頭胸部にある厳辞、特に肝 臓部がおしつぶされて黄色の肝臓液が排出される。 本発明者らは実験結果から、生鮮オキアミに対し 約40~1409/cm² 程度の強かな外力を加える かあるいは1000°G 程度の遠心力を数分間加え ることにより、肝臓液が体外に容易に排出される ことを認めた。

提供される。

本発明においては前記(2)のごとき方法に従ってオ キアミの肉部を採取するにあたり、先ず上記した ことき方法によりオキアミの体内から肝臓療を排 出させ除去する。肝臓硬を排出させたオキアミの 体の表面には肝臓液がまだ僅かに付着しているの で、清水または梅水で簡単に洗浄するかあるいは シャワー等で洗浄してこれを除去することにより 始んど完全に肝臓液の付着していないオキアミを 得る。

縮依で濃縮して甲酸類特有の美味なエキスを得る。 なおこのエキスを繰回蛋白中に適当量混入させる ことにより凝固蛋白の味を著るしく向上させるこ とができる。

本発明の方法により得られる疑問蛋白は、肝臓 成分を含まないために冷康保存中の品質保持性が 凌れており、肝臓液除去処理を行わないで製造し た蒸凝菌蛋白が約 6 か月で呈味を預ねるのに対し、 本光明による製品は1年間の保存にも耐えること ができる。

南氷洋におけるオキアミの漁期は12月~2月 の搬氷期に限られており、従つて1年以上の保存 寿命を要求されるオキアミ製品の製造においては 本光明の上記の効果は値めて重要な効果である。 なお、圧搾機で肉部から分離した般部は煮熟後乾 妹して優良な飼料として利用し得る。

実 쪤 例

生鮮な崩氷洋産オキアミ25kgをバスケット型 速心分離機に装入し、速心力1000はで2分間 回転させることにより肝臓およびその他の内臓を 始んど完全に除去して、頭胸部が扁平になりしか も尾腹部に肉を有するオキアミ12.5 kpを得た。 このオキアミを0.5 mの崩除を有する回転圧搾式 骨肉分離機により圧搾分離することにより潮状の 内部7.5 kp(原料のオキアミの60多の収率)を 得、これを直ちに不銹鈍製の加熱容器に移し、十 分階件しながら、直火で75 にまで加熱した(こ の間に60 C附近で蛋白が一部鹸肉した)。つぎ に加熱容器を沸勝湯煎中に移し、 滑拌しながら 加 熟を続けた。83 C で急速に蛋白の凝固が始まり、 90 C で醸固が完了したことが認められたが 90 C になお 2 分間保汚した。

つぎに容器を流水槽に浸着して内容物を70℃ 程度まで冷却し、ついでガーゼ布で沪遏した。か く得られた疑菌蛋白を再びパスケット型速心分離 燃に終入し、速心力2000倍程度で3分間脱水 することにより、水分60多の談黄色現肉状のオ キアミ熱緩固蛋白4.2㎏を得た(収率34多)。こ の壊薬固蛋白には苦味やエグ味が全くなく、肝臓 液除去処理をしない製品に比べてすぐれた甲数類 特開昭51-125774 (3) 特有のうま味があつた。 熱柴菌蛋白を分離した後 の夜部は更に速心力 3 0 0 0 G で 3 分間速心分離 し、ついで滅圧濃縮機により 4 5 ブリックス (水分 6 0 多前後)まで濃縮して比較的粘度の低いエキ ス 2 5 0 9 を得た(収率 2 多)このエキスにも前 記熱楽菌蛋白と同様、苦味やエグ味が全くなく、 甲酸類に特有のうま味があつた。

上記の熟練固直白を小型冷凍パンに装入して成 型し、-35℃で急速冷凍を行ない-25℃で深 存試験を実施した結果、本発明による肝縁後除去 処理を行わないものは3ケ月で褐変し始め、6ケ 月で味に変化が認められたのに対し、本発明の方 法により製造されたものは1年経過しても味の変 化は殆んど認められずまた油焼臭も少なかつた。

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5.添附書類の目録

| (1) 明 | 煭 | 퐌 | 上進 | |
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| (2) [2] | | | 1.100 | |
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| 砂蛋白質液及び蛋白質粉末の製造方法 | | | | | 明 | 者 | 江口通
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朙 オキアミは広く海洋に分布する動物性プラン クトンの一種でエビに似た体長1~7㎝程度の 2 発明の名称 甲殼類で、主にナガス鯨及び白ナカス鯨の餌と 蛋白質液及び蛋白質粉末の製造方法 して知られている。梅に南極海に棲息するユー ※許請求の範囲 ファーシア・スペルバ (Euphasia superba)種は. (1) 乾燥オキアミを水叉は塩水に浸漬して肉質 推定資源量数億トンといわれ、鯨の減少ととも 部分を容出させたのちオキアミの甲酸を除去 に増加の傾向にある。この豊富な資源を食用に し、かくして得られる溶出液を加熱し、つい 供し、将来予想される蛋白質源の不足に備える で生成した凝固物を除去することを特徴とす ことは重要な意義を有することである。このた る蛋白質液の製造方法。 め、各方面においてオキアミの食品化への研究 (2) 乾燥オキアミを水又は塩水に浸渍して肉質 が行なわれるようになつてきたが、未だ満足な 部分を溶出させたのちオキアミの甲殻を除去 成果は得られていない。このように研究が遅れ ている原因として、嗜好的なもののほかに可食 し、かくして得られる密出液を加熱し、生成 した慶園物を除去して得られる蛋白質液にた 化。保存性に次のような間歇点が指摘されてい 質を加え混合溶解したのち噴霧乾燥すること 3. を特徴とする蛋白質粉末の製造方法。 (1) オキアミはエビにくらべて小形であるらえ 3. 発明の詳細を説明 甲殼が体全体の30%もあり、しかも柔軟な 本発明は、乾燥オキアミから脂肪分を含まな ため、脱殻がきわめて困難である。このため、 い蛋白質液又は蛋白質粉末を製造する方法に関 ムキエビのような製品とすることが難かしく.

3

するものである。

従つて蛋白質部分のみを有効に利用するには、

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肉質部分を審出せざるを得ないが,そのため の有効な手段は未だ見出されていない。

オキアミはそのままの状態でも食用とする ことが不可能ではないが、オキアミ特有の臭 (アミン臭)と味を有する。これは所謂、悪 味悪臭ではないが小量の摂取でもすぐ飽きの くるものであつて、大量摂取にはむかないも のである。これらの臭や味はオキアミの脂質 中に含まれる高度不飽和脂肪酸等に起因する ところが大きく、さらにこの高度不飽和脂肪 酸の酸化は、保存性、嗜好性を悪くする原因 ともなつている。とのような魚体中の高度不 🎙 飽和脂肪酸を取り除く方法としては、例えば, n-ヘキサン,シクロヘキサン, アセトン等の 有機蓄媒により脱脂する方法が一般に用いら れているが、かかる方法では原料からくる時 長が失われ、旨味がなくなつてしまう他,有 機磨媒を用いるため,製造コストが高くなり. また食品衛生法上、天然食品として取扱うと とができなくなるという致命的な欠陥をもつ

という知見を得るに至つた。

本発明はかかる新知見にもとづいて完成さ れたもので、乾燥オキアミを所定時間水又は 塩水に浸漬して肉質部分を溶出させたのち、 オキアミの甲殻を除去し、かくして得られる 溶出液を加熱して蛋白藤固物を生成させ、つ いでこの離固物を除去して、脂肪分を含まな い蛋白質液を得ることから成る方法であり、 見つ、かくして得られる蛋白質液に糠質を加 えて攪拌下に加温して均一に混合溶解したの ち、常法により噴霧乾燥を行つて蛋白質粉末 を製造する方法である。

本 発明で用いられる乾燥オキアミは、 補獲 直後のオキアミをそのまま乾燥したもの、 又 は 補獲後そのまま陳結した生康結オキアミを 解凍後乾燥したもの等が用いられ、オキアミ を予め煮熟した後乾燥したものは本発明には 適さない。また、本発明によれば、原料とし て用いられる乾燥オキアミの水分含量が低い 程対固形分収率が低くなり、また水分含量が ている。したがつて、オキアミ中の高度不顧 和脂肪酸等の脂質を有機審媒を使用するとと なく除去するととができればオキアミの食品 化にとつて極めて好ましいことであるが、未 だこのよりな方法は知られていない。

(8) オキアミは、通常捕獲後直ちに生のまま、 あるいは薫熟して冷凍保蔵される。しかし、 オキアミは普通約80多の水分を含んでおり、 これがために陳結製品の解弾時には大量のド リップ(drip)が生じ、オキアミのもつている 多量の可容性蛋白質がかなりドリップへ移行 されて失われるので、その有効な利用方法を 開発することも急称である。

本発明者らはかかる点を解決するべく研究 を重ね,先に生及び凍結オキアミから脂肪分 のない良質な高留白質液及び高蛋白質粉末を 得る方法を発明し, 特顧昭 50-152874とし て特許出願したが更に研究を進めた結果,水 分含量の少ない乾燥オキアミからも水又は塩 水を用いて水裕性蛋白質を有効に抽出しうる

高い程原料全体に対する収率が低下する。1. たがつて、本発明に用いられる原料の乾燥オ キアミとしては、水分含量 10~50 %のものが 最も好ましいが、場合によつては水分含量が この範囲を越えるものも使用することができ る。

これらの乾燥オキアミを浸漬する場合、凍 結されているものを原料とするときは、水を 加えて解凍後そのまま浸漬すれば良い。浸漬 は通常室區以下の温度で、5~24時間行う。 浸漬中に帯拌等を行えば更に短時間で目的を 達成することができる。浸滑液の量はオキア ミが充分ひたる程度の畳以上あれば充分であ り、浸漬中に帯拌等を行つてもさしつかえな い。また、浸漬液として塩水を用いる場合は、 3 多程度の比較的低濃度の塩水を用いるとと が製品中の塩分を少なくする上で好ましい。 つぎに、浸漬液からオキアミの甲敷を沪過あ るいは金網ですくう等の方法で除去し、必要 により、更に甲殼に付着している肉質部分を

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洗い落すことにより、オキアミの器出版を得 ることができる。

つぎに、密出液を加熱するには、15~30 分間蒸煮すれば充分である。との加熱により、 高度不飽和脂肪酸等の脂肪分が一部の蛋白質 と共に暴励物として析出するが、このものは、 速心分離器、加圧沪過器(フィルターブレス)、 減圧沪過器(オリバーフィルター)等で容易に 除去することができる。以上の方法により脂 肪分を含まない蛋白質液を得ることができる。 この蛋白質液は、必要により活性炭で脱色す ることにより更に良質な蛋白質液にすること ができる。

本発明によれば、上記の如くして得られる 準白質液に所定量の種質を加え、必要により 加温し、均一に混合溶解したのち、常法によ り噴終乾燥することにより蛋白度粉末にする ことができる。この場合に加える糖質として は、乳糠、麦芽糖の如き糖類、可溶性テンプ ンの如き澱粉類、デキストリン、粉あめの如

好性の面でもすぐれている。また、本発明に よれば、生康結品を原料とする方法に比べて 水分含量の低い原料を用いるので輸送上のみ ならず貯蔵上からも極めて経済的であり、ま た、処理操作の上でも対原料収率が著しく向 上するので極めて有利である。オキアミは固 形分の約70%が粗蛋白質であるが、本発明 方法によれば、この内の約6割程度を蛋白質 被として待ることができる。更に本発明方法 には、アルカリ、酸、有機溶媒等の化学薬品 の添加あるいはこれらを便用する処理が全く ない。したがつて、本発明方法により得られ た蛋白質 変又は蛋白質粉末は食品衛生法上間 題がないという優れた利点を有する。

つきに実施例を示し,本発明を更に詳細に 説明する。

実 施 例 1

水分含量15%のオキアミ100万を真水 1460万に12時間10°C以下で受賞すること によりオキアミの肉質部分を浸滑液中に落出 特問 昭52-114046(3)

き 影 粉 中間 分 解物, あるいは, これらの 影 粉 中間 分 解物 を 愛 元 し て 得 ら れ る 糖 類 ア ル コ ー ル 等 が 用 い ら れ る 。 加 え る 糖 質 の 骨 は , 糖 質 の 種 類 に よ り 多 少 差 は あ る が , 通 常 蛋 白 質 液 中 の 罪 白 質 含 量 に 対 し 0.7 ~ 3 倍 量 用 い る の が 好 ま し い 。 糖 質 の 量 が 少 な す ぎ る と 蛋 白 質 粉 末 の 吸 湿 性 が 著 し く な り , 保 存 性 等 の 面 か ら 好 ま し く な い 。

本発明によつて得られた蛋白質液は外額が 帮ね後黄色でエビの芳香及び味覚を有するた め調味料の主原料として有用である。また、 本発明によつて得られる蛋白質粉末は、外観 が悪ね後灰白色でソフトなエビ臭と味覚を有 し、口あたりも滑らかで溶け易いので種々の 食品における主原料並びに創原料として有用 であり、且つ栄養価の向上、味覚の改善等を 目的として、従来の水産ねり製品あるいは風 味を珍重するスナック食品、香辛料等にも添 加、使用することができる。また、脂肪分を 含まないため酸化が起りにくく、保存性、嗜

せしめる。つぎに、オキアミの魚体をすくい 上けて取り除き、更にこの魚体を真水235 で2回洗浄しオキアミの溶出液1660%を得 る。この溶出液を30分間蒸煮することによ つて紫灰色の酸固物が析出する。つぎに、 酸 固物は、沪布を施した沪渦器で沪し取ること によつて、脂肪分を含まない蛋白質※1620 い(蛋白質含質2.2%)を得ることができる。 つづいて、この蛋白質液をカーボン800gで 脱色することにより、殆ど無色の蛋白質液を 得ることができる。

上記の方法により得られた登白質液, 蛋白 質粉末及び通常の乾燥により得られた乾燥オ キアミ紛末の成分組成を第1表に示す。

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第1表

| | | | | 蛋白質液 | 蛋白質粉末 | 乾燥オキアミ粉末 |
|---|---|---|---|---------|-------|----------|
| 水 | | | 分 | 95.6 \$ | 3.8% | 4.0 \$ |
| 粗 | 蛋 | þ | 質 | 2.2 | 33.9 | 70.4 |
| 脂 | | | 眆 | 0 | 0.1 | 5.9 |
| 厌 | | | ዎ | 0.2 | 3.1 | 11.4 |
| 食 | | | 塩 | 0.2 | 3.1 | 2.4 |
| 糖 | | | 質 | - | 56.0 | - |

美 実施例 2

水分含量40%のオキアミ100%を真水 940%に12時間10°0以下で浸漬すること によりオキアミの肉質部分を浸漬液中に溶出 せしめる。つぎに、オキアミの魚体をすくい 上げて取り除き、更にこの魚体を真水155% で2回洗浄しオキアミの務出液1060%を得 る。この器出液を30分間蒸煮することによ つて紫灰色の酸固物が析出する。つぎに、酸 間物は、炉布を施した沪過器で炉し取ること によつて、脂肪分を含まない蛋白質液995% 特開 昭52-114046(4)

(蛋白質含量 2.8 %)を得るととができる。 つづいて, この蛋白質液をカーボン 800 gで 脱色することにより, 殆ど無色の蛋白質液を 得ることができる。

つぎに、ここに得られた蛋白質液(蛋白質 量27.9 %)に対し、デキストリン54 %(蛋白 質量に対し1.94倍量)を加え、攪拌下に加温 して均一に容解した後、噴霧乾燥することに よつて良質な蛋白質粉末84.5 %を得ること ができる。

梅許出願人 日研化学株式会社

-242-

| Electronic A | cknowledgement Receipt |
|--------------------------------------|-------------------------------------|
| EFS ID: | 12358705 |
| Application Number: | 12057775 |
| International Application Number: | |
| Confirmation Number: | 1945 |
| Title of Invention: | BIOEFFECTIVE KRILL OIL COMPOSITIONS |
| First Named Inventor/Applicant Name: | Inge Bruheim |
| Customer Number: | 72960 |
| Filer: | John Mitchell Jones/Vickie Hoeft |
| Filer Authorized By: | John Mitchell Jones |
| Attorney Docket Number: | NATNUT-14409/US-5/ORD |
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| 5 | Foreign Reference | JP52114046.pdf | 6c5aba15820bc9acefea0035e8f0292417c6
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<u>der 35 U.S.C. 371</u>
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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Inge Bruheim, et alConfirmation:1945Serial No.:12/057,775Group No.:1651Filed:03-28-2008Examiner:Ware, Deborah K.Entitled:BIOEFFECTIVE KRILL OIL COMPOSITIONSFiled:

INFORMATION DISCLOSURE STATEMENT LETTER

EFS Web Filed Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir or Madam:

The citations listed in the attached **IDS Form SB08A** may be material to the examination of the aboveidentified application, and are therefore submitted in compliance with the duty of disclosure defined in 37 C.F.R. §§ 1.56 and 1.97.

Applicants wish to bring to the Examiner's attention that we are not providing copies of US Patents as instructed under 37 CFR 1.98(a)(2). The Examiner is requested to make these citations of official record in this application.

Applicants wish to bring to the Examiner's attention that the references supplied in this IDS are from the JP Office Action mailed February 23, 2012 from related JP Patent Application No. 2010-52244.

This Information Disclosure Statement under 37 C.F.R. §§ 1.56 and 1.97 is not to be construed as a representation that a search has been made, that additional information material to the examination of this application does not exist, or that any one or more of these citations constitutes prior art.

The Commissioner is hereby authorized to charge any required fees or credit any overpayments to Attorney Deposit Account No.: **50-4302**, referencing Attorney Docket No.: **NATNUT-14409/US-5/ORD**.

Dated: <u>March 21, 2012</u>

<u>/J. Mitchell Jones/</u> J. Mitchell Jones Registration No. 44,174 CASIMIR JONES, S.C. 2275 Deming Way, Suite 310 Middleton, WI 53562 608.662.1277

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INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

| Application Number | | 12057775 | | | |
|------------------------|--------|-----------------------|--|--|--|
| Filing Date | | 2008-03-28 | | | |
| First Named Inventor | Inge E | Bruheim | | | |
| Art Unit | | 1651 | | | |
| Examiner Name | Ware, | Deborah K. | | | |
| Attorney Docket Number | | NATNUT-14409/US-5/ORD | | | |

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|--|----------------------|-----------------------------------|-----------------------|--|
| | Filing Date | | 2008-03-28 | |
| | First Named Inventor | First Named Inventor Inge Bruheim | | |
| (Not for submission under 37 CFR 1.99) | Art Unit | | 1651 | |
| | Examiner Name | Ware | , Deborah K. | |
| | Attorney Docket Numb | er | NATNUT-14409/US-5/ORD | |

| | 1 December 8, 2011 Office Action, KR Patent Application No. 10-2010-7006897 and its English translation | | | | | | | | |
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|--|------------------------|--------|-----------------------|--|
| | Filing Date | | 2008-03-28 | |
| | First Named Inventor | Inge E | ruheim | |
| | Art Unit | | 1651 | |
| | Examiner Name | Ware, | Deborah K. | |
| | Attorney Docket Number | | NATNUT-14409/US-5/ORD | |

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Please see 37 CFR 1.97 and 1.98 to make the appropriate selection(s):

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See attached certification statement.

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A signature of the applicant or representative is required in accordance with CFR 1.33, 10.18. Please see CFR 1.4(d) for the form of the signature.

| Signature | /J. Mitchell Jones/ | Date (YYYY-MM-DD) | 2012-02-20 |
|------------|---------------------|---------------------|------------|
| Name/Print | J. Mitchell Jones | Registration Number | 44174 |

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(19)日本国特許庁(JP) (12) 公表特許公報(A)

(11)特許出願公表番号

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| A23J 7/0 | 0 | A 2 3 J 7/00 | 4 D 0 5 6 |
| A61K 31/6 | 61 | A61K 31/661 | 4H059 |
| A61P 3/0 | 2 | A 6 1 P 3/02 | |
| B01D 11/0 | 4 | B01D 11/04 | С |
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| (86) 国際出願番号 | PCT/IB01/00841 |
| (87) 国際 公開番号 | WO01/076715 |
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| (32)優先日 | 平成12年4月12日(2000.4.12) |
| (33)優先権主張国 | ドイツ (DE) |
| (31)優先権主張番号 | 60/271, 209 |
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| (33)優先権主張国 | 米国(US) |
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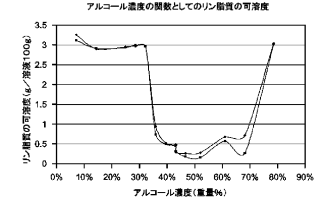
(71)出願人 ウエストファリア セパレイター インダ ストリー ゲーエムベーハー ドイツ, デー-59302 エルデ, ヴェ ルナーーハビッヒーストラーセ 1 (72)発明者 フルシュカ, ステファン, エム. ドイツ, 59302 エルデ, エーネープ ラウクジーペーストラーセ 7 (72)発明者 キルシュナー, ステファン ドイツ, 33334 キュータースロフ, エルゼンキルムストラーセ 61 (74)代理人 弁理士 山田 行一 (外1名)

最終頁に続く

(54) 【発明の名称】 油と極性脂質を含有する天然物質の分画方法

(57)【要約】

本発明は、極性脂質リッチ物質、好ましくはリン脂質の 製造プロセスに関する。好ましくは、該極性脂質リッチ 物質を、脱油した天然物質から、水とアルコールを用い た抽出によって分離と回収し、得られた混合物を密度分 離を用いて分離する。また本発明は、リン脂質の抽出と 回収の前に天然物質を脱油するための改良されたプロセ スも含む。



-1-

【特許請求の範囲】

【請求項1】 低油含有量極性脂質含有物質の分画プロセスであって、

(a)前記低油含有量極性脂質含有物質を、水と水溶性有機溶媒に混合するス テップと、

(b)該混合物を密度分離にかけて軽相と重相に分けるステップと を有するプロセス。

【請求項2】 前記水溶性有機溶媒が、極性溶媒を有する請求項1に記載の プロセス。

【請求項3】 前記水溶性有機溶媒が、アルコールを有する請求項1に記載 のプロセス。

【請求項4】 前記水溶性有機溶媒が、C₁~C₈のアルコールを有する請求 項1に記載のプロセス。

【請求項5】 前記水溶性有機溶媒が、イソプロパノール、エタノール、又はこれらの混合物を有する請求項1に記載のプロセス。

【請求項6】 密度分離にかけられる前記物質が、水溶性有機溶媒と水の混 合物中に可溶化/分散され、前記水溶性有機溶媒が、存在する水溶性有機溶媒と 水の合計量の約5~約35重量%を占める請求項1~5のいずれかに記載のプロ セス。

【請求項7】 密度分離にかけられる前記物質が、水溶性有機溶媒と水の混 合物中に可溶化/分散され、前記水溶性有機溶媒が、存在する水溶性有機溶媒と 水の合計量の約68~約98重量%を占める請求項1~6のいずれかに記載のプ ロセス。

【請求項8】 該プロセスの実行中に温度が65℃を超えない請求項1~7 のいずれかに記載のプロセス。

【請求項9】 プロセス中のpHが、pH4~約pH10である請求項1~ 8のいずれかに記載のプロセス。

【請求項10】 前記混合ステップと密度分離ステップが少なくとも1回繰 り返される請求項1~9のいずれかに記載のプロセス。

【請求項11】 前記低油含有量極性脂質含有物質が、卵と、魚と、甲殻類

と、微生物と、脳組織と、牛乳と、肉と、脂肪種子をはじめとする植物物質との 少なくとも1種から得られる請求項1~10のいずれかに記載のプロセス。

【請求項12】 前記低油含有量極性脂質含有物質中にもともと存在する前 記極性脂質の少なくとも60%が、極性脂質リッチ軽相中で回収される請求項1 ~11のいずれかに記載のプロセス。

【請求項13】 油/極性脂質/タンパク質含有混合物の分画プロセスであって、

(a)前記混合物から油を分離し、油リッチ画分と極性脂質/タンパク質リッ チ画分を形成するステップと、

(b)前記極性脂質/タンパク質リッチ画分に、水溶性有機溶媒を加えるステ ップと、

(c)前記水溶性有機溶媒と極性脂質/タンパク質リッチ画分を密度分離にかけて、極性脂質リッチ画分とタンパク質リッチ画分を形成するステップと を有するプロセス。

【請求項14】 前記ステップ(a)の油の分離が、

(a) 前記油/極性/タンパク質含有混合物をホモジナイズするステップと、

(b) 前記混合物に水溶性有機溶媒と水を加えるステップと、

(c)該混合物を、油リッチ画分と極性脂質/タンパク質リッチ画分に分離す るステップと

を有する請求項13に記載のプロセス。

【請求項15】 前記ステップ(a)の油の分離が、

(a)前記混合物に水溶性有機溶媒と水を加えるステップと、

(b)前記油/極性脂質/タンパク質含有混合物を、ホモジナイズするステップと、

(c)該混合物を、油リッチ画分と極性脂質/タンパク質リッチ画分に分離す るステップと

を有する請求項13又は14に記載のプロセス。

【請求項16】 前記ステップ(a)の油の分離が、

(a) 前記油/極性脂質/タンパク質含有混合物をホモジナイズするステップ

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と、

(b) 前記混合物に水溶性有機溶媒と水を加えるステップと、

(c)該混合物をホモジナイズするステップと、

(d) 該混合物を、油リッチ画分と極性脂質/タンパク質リッチ画分に分離す るステップと

を有する請求項13に記載のプロセス。

【請求項17】 前記油/極性脂質/タンパク質含有混合物が、卵から得られる請求項13~16のいずれかに記載のプロセス。

【請求項18】 水溶性有機溶媒が、密度分離後に極性脂質リッチ画分とタンパク質リッチ画分から回収される請求項13~17のいずれかに記載のプロセス。

【請求項19】 前記ステップ(a) で形成される前記極性脂質/タンパク 質リッチ画分が、約30~50重量%の極性脂質と、約50~70重量%のタン パク質を有する請求項13~18のいずれかに記載のプロセス。

【請求項20】 前記ステップ(a)で形成される前記油リッチ画分が、約75~約95重量%のトリアシルグリセロールを有する請求項13~19のいず れかに記載のプロセス。

【請求項21】 前記油/極性脂質/タンパク質含有混合物が更にコレステ ロールを有し、前記コレステロールの大部分が、ステップ(a)の分離に従って 前記油リッチ画分に相当する請求項13~20のいずれかに記載のプロセス。

【請求項22】 ステップ(b)で加えられる前記水溶性有機溶媒が、水溶 性有機溶媒/水混合物を形成し、前記水溶性有機溶媒が、存在する水溶性有機溶 媒と水の合計量の約20~約35重量%を占める請求項13~21のいずれかに 記載のプロセス。

【請求項23】 ステップ(b)で加えられる前記水溶性有機溶媒が、水溶 性有機溶媒/水混合物を形成し、前記水溶性有機溶媒が、存在する水溶性有機溶 媒と水の合計量の約68~約98重量%を占める、請求項13~21のいずれか に記載のプロセス。

【請求項24】 前記水溶性有機溶媒が、向流洗浄と、蒸発と、乾燥とのい

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ずれかにより回収される請求項13~23のいずれかに記載のプロセス。

【請求項25】 前記極性脂質リッチ画分を乾燥して水溶性有機溶媒を回収 し、残留タンパク質を沈殿させるために約80重量%を超える水溶性有機溶媒を 含む水溶性有機溶媒/水混合物で洗浄し、さらに乾燥して水溶性有機溶媒を回収 する請求項13~24のいずれかに記載のプロセス。

【請求項26】 前記水溶性有機溶媒の添加により、前記タンパク質の少な くとも一部が沈殿し、この沈殿物が密度分離によって回収される請求項25記載 のプロセス。

【請求項27】 水の添加により、前記極性脂質リッチ画分から残留タンパ ク質が除去される請求項13~26のいずれかに記載のプロセス。

【請求項28】 前記水溶性有機溶媒が極性溶媒を含む、請求項13~27 のいずれかに記載のプロセス。

【請求項29】 前記水溶性有機溶媒が、アルコールを有する請求項13~ 27のいずれかに記載のプロセス。

【請求項30】 前記水溶性有機溶媒が、C₁~C₈のアルコールを有する請 求項13~27のいずれかに記載のプロセス。

【請求項31】 前記水溶性有機溶媒が、イソプロパノール、エタノール、 又はこれらの混合物のいずれかを有する請求項13~27のいずれかに記載のプ ロセス。

【請求項32】 プロセス中のpHが、pH4~約pH10である請求項1 3~31のいずれかに記載のプロセス。

【請求項33】 前記混合物が、卵と、魚と、甲殻類と、微生物と、脳組織 と、牛乳と、肉と、脂肪種子をはじめとする植物物質との少なくとも1種から得 られる請求項13~32のいずれかに記載のプロセス。

【請求項34】 前記混合物中に当初より存在する前記極性脂質の少なくと も60%が、極性脂質画分中で回収される請求項13~33のいずれかに記載の プロセス。

【請求項35】 プロセス中に温度が65℃を超えない請求項13~34の いずれかに記載のプロセス。 【請求項36】 水溶性有機溶媒の使用により、極性脂質含有混合物から極 性脂質を回収するためのプロセスであって、前記回収を助けるために水溶性有機 溶媒の水溶液中への極性脂質の比較的高い可溶度を利用し、該水溶性有機溶媒は 該水溶液中のうちの35重量%未満又は68重量%を超える量を占める、上記プ ロセス。

【請求項37】 前記混合物が、卵と、魚と、甲殻類と、微生物と、脳組織 と、牛乳と、肉と、脂肪種子をはじめとする植物物質との少なくとも1種から得 られる請求項36に記載のプロセス。

【請求項38】 前記極性脂質が、リン脂質を有する請求項1~37のいず れかに記載のプロセス。

【請求項39】 前記プロセスの少なくとも一部が、低酸素雰囲気中で行われる請求項1~38のいずれかに記載のプロセス。

【請求項40】 油/極性脂質/タンパク質含有混合物を分画するプロセス であって、

(a)前記油/極性脂質/タンパク質含有混合物に水溶性有機溶媒を加えるス テップと、

(b)該油/極性脂質/タンパク質含有混合物をホモジナイズするステップと

(c)前記混合物から油を分離し、油リッチ画分と極性脂質/タンパク質リッ チ画分を形成するステップと

を有するプロセス。

【請求項41】 前記ステップ(a)と、前記ステップ(b)と、前記ステ ップ(c)が、

(a)前記油/極性脂質/タンパク質含有混合物をホモジナイズするステップ と、

(b) 水溶性有機溶媒と水を、前記混合物に加えるステップと、

(c)該混合物を、油リッチ画分と極性脂質/タンパク質リッチ画分に分離す るステップと

を有する請求項40に記載のプロセス。

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【請求項42】 前記ステップ(a)と、前記ステップ(b)と、前記ステ ップ(c)が、

(a) 水溶性有機溶媒と水を、前記混合物に加えるステップと、

(b)前記水溶性有機溶媒と油/極性脂質/タンパク質含有混合物を、ホモジ ナイズするステップと、

(c)該混合物を、油リッチ画分と極性脂質/タンパク質リッチ画分に分離す るステップと

を有する請求項40又は請求項41に記載のプロセス。

【請求項43】 前記ステップ(a)と、前記ステップ(b)と、前記ステ ップ(c)が、

(a)前記油/極性脂質/タンパク質含有混合物を、ホモジナイズするステップと、

(b) 水溶性有機溶媒と水を前記混合物に加えるステップと、

(c)得られた混合物をホモジナイズするステップと、

(d)得られた混合物を油リッチ画分と極性脂質/タンパク質リッチ画分に分離するステップと、

を有する請求項40に記載のプロセス。

【請求項44】 前記混合物が、卵と、魚と、甲殻類と、微生物と、脳組織 と、牛乳と、肉と、脂肪種子をはじめとする植物物質との少なくとも1種から得 られる請求項40~43のいずれかに記載のプロセス。

【請求項45】 前記ホモジナイズが、約100バール~約1000バール の圧力で行われる請求項14~44のいずれかに記載のプロセス。

【請求項46】 前記ホモジナイズが、約150バール~約350バールの 圧力で行われる請求項14~45のいずれかに記載のプロセス。

【請求項47】 前記油/極性脂質/タンパク質含有混合物が、水溶性有機 溶媒と水の混合物の中に可溶化/分散され、該混合物中において前記水溶性有機 溶媒は、存在する水溶性有機溶媒と水の合計量の約5%~約35重量%を占める 請求項40~46のいずれかに記載のプロセス。

【請求項48】 極性脂質リッチ画分を得るために行われる前記密度分離が

2つのステップで行われ、第1ステップでは、存在する水溶性有機溶媒と水の合 計量における該水溶性有機溶媒の割合が約5~約35重量%を占め、密度分離に より第1極性脂質リッチ画分が得られ、第2ステップでは、存在する水溶性有機 溶媒と水の合計量における該水溶性有機溶媒の割合が約68~約98重量%を占 め、密度分離により第2極性脂質リッチ画分が得られ、該第2極性脂質リッチ画 分が第1極性脂質リッチ画分よりも高い割合で極性脂質を有する請求項1~39 のいずれかに記載のプロセス。

【請求項49】 極性脂質リッチ画分中のコレステロールを減らすための方 法であって、

(a) コレステロールをゼロないし少量しか含まない油を、前記極性脂質リッ チ画分に添加するステップと、

(b)該コレステロールを該油相中に隔離させるために該混合物を脱油することにより、該極性脂質リッチ画分の中のコレステロール量を減らすステップとを有する方法。

【請求項50】 前記請求項1~49のプロセスのいずれかによって生成される、油含有の、極性脂質含有の、又はタンパク質含有の、生成物。

【発明の詳細な説明】

[0001]

【発明の属する技術分野】

本発明は、抽出、分離と回収の分野、特に天然物質等の混合物から極性脂質リ ッチ画分を抽出、分離と回収する分野に関する。物質中の他の画分を同時に回収 することができ、これらの画分(例えばタンパク質リッチ画分など)は、抽出プ ロセスで穏やかな条件が使用されるので、これらの元々の機能の大部分または全 てを保持する。

[0002]

【従来の技術】

極性脂質は例えば、リン脂質(例えばホスファチジルコリン、ホスファチジル エタノールアミン、ホスファチジルイノシトール、ホスファチジルセリン、ホス ファチジルグリセロールまたはジホスファチジルグリセロール等)、セファリン 、スフィンゴリピド(スフィンゴミエリンとグリコスフィンゴリピド)、グリセ ロ糖脂質等を挙げることができる。リン脂質は、以下の主な構造ユニットで構成 される:脂肪酸、グリセロール、リン酸、アミノアルコールと炭水化物。これら は一般に、植物、微生物と動物の膜構造において重要な役割を担う構造脂質であ ると考えられている。これらの化学構造によって、極性脂質は双極性を示し、極 性溶媒と非極性溶媒の両方において可溶性または部分的な可溶性を示す。本明細 書において極性脂質という用語は、天然の極性脂質に限らず、化学修飾された極 性脂質も含む。油という用語は様々な意味を有するが、本明細書で用いられる際 には、トリアシルグリセロール画分を指すものとする。

[0003]

極性脂質(特にリン脂質)の重要な特徴の1つは、これらが一般的には多不飽 和脂肪酸(PUFA:2以上の不飽和結合を有する脂肪酸)を含むことである。 多くの植物、微生物と動物系において、これらは特に、 ω -3と ω -6シリーズ の高度不飽和脂肪酸(HUFA:4以上の不飽和結合を有する脂肪酸)に富んで いる。これらの高度不飽和脂肪酸は、トリアシルグリセロールの状態では不安定 であると考えられるが、リン脂質に組み込まれると強化された安定性を示す。

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[0004]

市販されているPUFAリッチリン脂質の主な起源は、大豆とキャノーラ種子 である。これらの生物物質(バイオマテリアル)は、遺伝子修飾されない限りは 、はっきり認められる量のHUFAを含まない。リン脂質(一般的にはレシチン と呼ばれる)は、これらの脂肪種子から植物油抽出プロセスの副産物として常套 的に回収される。例えば、大豆またはキャノーラ油の製造において、豆(種子) をまず熱処理した後に砕き、すりつぶし、と/またはフレーク状にし、その後へ キサン等の非極性溶媒で抽出する。ヘキサンは、これらの種子から、様々な量の 極性脂質(レシチン)と一緒にトリアシルグリセロールリッチ画分を取り除く。 次に、通常の油精製プロセスの一部として物理的もしくは化学的に抽出油の脱ガ ム(レシチン除去)を行い、沈殿したレシチンを回収する。しかし、このプロセ スは以下の2つの欠点を有する: (1) ヘキサンで抽出する前に種子を熱処理し なければならず、これは処理コストを増大させ且つタンパク質画分を変質させる ことにより、副産物としてのその価値を低下させる;と(2) ヘキサン等の非極 性溶媒の使用は、対処しなければならない毒性と引火性の問題も呈する。

【0005】

「脱ガム」プロセスで抽出された粗レシチンは、最大で約33%の油(トリア シルグリセロール)を含み得る。粗レシチンからこの油を分離するための1つの 好適な方法は、アセトンによる抽出である。油(トリアシルグリセロール)はア セトン中に可溶性であり、レシチンは不溶性である。遠心分離によりアセトン溶 液を沈殿物(レシチン)から分離し、該沈殿物をまずは流動層乾燥機で乾燥して から真空乾燥オーブンで乾燥して、該産物を乾燥しているときに残留アセトンを 回収する。一般には乾燥温度50から70℃が用いられる。得られた乾燥レシチ ンは、約2から4重量%の油(トリアシルグリセロール)を含む。70℃を超え るプロセス温度はリン脂質の熱分解につながり得る。しかし、70℃未満の温度 であっても、アセトンの存在により、該リン脂質の感覚刺激的品質を損なわせ得 る生成物の形成につながる。これら副産物は、生成物にカビ臭と、辛い後味を付 与し得る。

[0006]

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ヘキサン等の非極性溶媒の使用を回避するため、とアセトンベース・プロセス のマイナスの副作用を回避するために、超臨界流体(特に超臨界CO。)の使用 を含む多くのプロセスが提唱された。例えば、米国特許第4,367,178号 は、粗大豆レシチン調製物から油を除去することにより該調製物を一部精製する ための超臨界CO₂の使用を開示している。ドイツ国特許第DE-A30 11 185号と DE-A 32 29 041号は、超臨界CO。とエタンをそれぞれ用いた粗レシチンの脱 油方法を開示している。プロパンなどの少量の炭化水素を超臨界CO。に加えて 共留剤として作用させることを含む他の超臨界プロセスが提案されている。しか し、超臨界流体抽出システムは非常に資本支出が大きく、また連続的な操作がで きない。さらに、抽出時間が長く、抽出前にバイオマテリアルを乾燥しなければ ならず、またこれにより、酸化防止剤を用いて得られた乾燥産物を安定化するこ とがさらに難しくなる。これらの要因の全てにより、超臨界プロセスは、極性脂 質物質またはこれらの物質の混合物を抽出と回収するための最も費用のかかる選 択肢のうちの1つとなっている。そのため、低圧力での液化炭化水素での抽出を 用いた他のプロセスが記載されている。例えば、米国特許第2,548,434 号は、脂肪種子物質を脱油しと低圧力(35~45バール)且つ高温(79~9 3℃)で液体炭化水素を用いて粗レシチンを回収するための方法を開示している 。米国特許第5,597,602号は、さらに低い圧力と温度で動作する同様の プロセスについて記載している。しかし、これらの改良を行っても、超臨界流体 抽出は依然として非常にコスト高であり、大規模な商業規模で食品用途でのリン 脂質の製造には現在使用されていない。

[0007]

【発明が解決しようとする課題】

HUFAリッチ極性脂質の主な流通物質は卵黄である。産業規模では卵リン脂 質の回収のために、主に2つの方法が用いられている。いずれの方法も、抽出前 に卵黄の乾燥を必要とする。第1のプロセスでは、乾燥した卵黄粉末をアセトン でまず抽出してトリアシルグリセロールを除去する。次にこれを純粋アルコール で抽出してリン脂質を除去する。第2のプロセスでは、純粋アルコールを用いて 乾燥卵黄から油/レシチン画分を抽出する。次に油/レシチン相をアセトンで抽 出して、トリアシルグリセロールを除去し、レシチン画分を残す。これらの方法 にはどちらにも幾つかの欠点がある: (1)処理前にまず卵黄を乾燥しなければ ならない(コストの高いステップ)、とさらにこの乾燥プロセスはタンパク質に ダメージを与えたりまたはこれを変性させて、その食品成分としての価値を大き く低下させ得る; (2)効果的にするために、これらのプロセスで用いられるア ルコールとアセトン濃度は80%を超える、好ましくは90%を超えるものでな ければならない。これより高い純度の溶媒はさらに高価であり、高濃度の溶媒の 使用はタンパク質の変性につながり、これらの価値を低下させる;と(3)2つ のタイプの溶媒を回収するためには別々の溶媒回収条件が利用可能でなければな らず、これは設備コストを増大させる。これら3つの欠点の全ては卵黄からの極 性脂質リッチ画分の分離と回収コストの大きな増大につながる。

【0008】

カナダ国特許第1,335,054号は、エタノール、高温、濾過と低温結晶 化を用いた、新鮮な液体卵黄からの抽出によりタンパク質、油とレシチン画分を 得るためのプロセスについて記載している。しかしこの方法は、幾つかの欠点を 有する:(1)高濃度のエタノールの使用によるタンパク質の変性;(2)この プロセスがエタノールに限定される;(3)このプロセスでは、まずタンパク質 を除去してから、油画分からレシチンを回収する。レシチン生成物の純度は開示 されていない。

【0009】

現在の技術的状況を鑑みると、動作コストが低く、関連する副産物の価値を保 護し、且つ極性脂質生成物中のHUFAの総合品質を保護する、食品級極性脂質 生成物のための改良された抽出技法が依然として必要である。

【0010】

【課題を解決するための手段】

本発明に従って、従来技術の欠点の全ては含まない、天然のバイオマテリアル から極性脂質を回収するための改良されたプロセスが提供される。本発明は、こ れまで可能と考えられていたものよりもかなり低い濃度のアルコールを用いて部 分的にまたは完全に脱油されたバイオマテリアルから極性脂質と/または極性脂

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質含有混合物を回収するためのプロセスである。また本発明は、本発明中にその 概要が記されている方法による抽出/回収前にバイオマテリアルを脱油するため の改良されたプロセスも提供する。

[0011]

本発明の1つの実施形態に従って、低油含有量極性脂質含有物質(を分画する ためのプロセスが提供される。このプロセスは、低油含有量極性脂質含有物質に 水と水溶性有機溶媒を混合するステップ、と該混合物を(例えば重量や遠心力を 用いた)密度分離にかけてこれを軽相と重相に分けるステップを含む。好ましく は、該軽相は極性脂質リッチ画分を含み、と該重相はタンパク質リッチ画分を含 む。「低油含有」とは、その極性脂質含有物質が、トリアシルグリセロールを乾 燥重量で約20%未満、好ましくは約15%未満、より好ましくは約10%未満 、と最も好ましくは約5%未満有することを意味する。低油含有極性脂質リッチ 物質は、極性脂質リッチ物質から油を除去することによって、または油含有量の 低い極性脂質リッチ物質を選択することによって、得ることができる。例えば、 (脂肪種子を除く) ある植物物質とある微生物は、油含有量の低い極性脂質リッ

チ物質として使用することができる。好ましくは、低油含有量極性脂質含有物質 中にもともと存在する極性脂質の少なくとも60%とより好ましくは少なくとも 80%が極性脂質リッチ軽相中で回収される。

[0012]

本発明の他の実施形態に従って、油/極性脂質/タンパク質含有混合物の分画 プロセスが提供される。このプロセスは、該混合物から油を分離して油リッチ画 分と極性脂質/タンパク質リッチ画分を形成するステップ、該極性脂質/タンパ ク質リッチ画分に水溶性有機溶媒を加えるステップ、と該水溶性有機溶媒と極性 脂質/タンパク質リッチ画分を(例えば重力や遠心力を用いた)密度分離にかけ て、極性脂質リッチ画分とタンパク質リッチ画分を形成するステップ、を含む。 好ましくは、該混合物中にもともと存在する極性脂質の少なくとも60%とより 好ましくは少なくとも80%が極性脂質リッチ画分中で回収される。

[0013]

本発明の他の実施形態に従って、水溶性有機溶媒を用いて極性脂質含有混合物

から極性脂質を回収するためのプロセスであって、該回収を助けるために該水溶 性有機溶媒の水溶液中への極性脂質の比較的高い可溶度を利用し、ここで、該水 溶性有機溶媒は該水溶液中のうちの35重量%未満または68重量%を超える量 を占める、上記プロセスが提供される。

[0014]

本発明の他の実施形態に従って、油/極性脂質/タンパク質含有混合物の分面 プロセスが提供される。このプロセスは、該油/極性脂質/タンパク質含有混合 物に水溶性有機溶媒を加えるステップ、該水溶性有機溶媒と油/極性脂質/タン パク質含有混合物をホモジナイズにかけるステップ、と該混合物から油を分離し て油リッチ画分と極性脂質/タンパク質リッチ画分を形成するステップを含む。

[0015]

本発明の実施形態の利点は、他の既知の方法よりもコストが非常に低いことで ある。本発明の実施形態の利点は、抽出タンパク質等の他の副産物を劣化から保 護することにより副産物としてのこれらの販売価値を増大させることである。本 発明の実施形態の利点は、極性脂質の中のHUFAを劣化から保護することであ る。これらの利点は、本発明の主なアスペクトの幾つかによるものである: (1) 脱油の前にバイオマテリアルを乾燥する必要がない; (2)このプロセスは低 濃度のアルコールを使用する; (3)関連する副産物の品質と機能が劣化(例え ば高温または高濃度溶媒によるタンパク質の変性;脂質の酸化;望ましくない副 産物の形成など)から保護される;と(4)(設備と処理ステップの両方の点に おいて)プロセス全体が非常に単純である。好ましくは、プロセスのステップは 、不活性もしくは非反応性ガス(例えば窒素、二酸化炭素、アルゴンなど)の使 用、溶媒蒸気の使用、部分的もしくは完全な真空の使用、またはこれらの任意の 組合せを含み得る低酸素雰囲気下で行われる。

[0016]

本発明は、図面を参照することによってより簡単に理解することができる。

【0017】

【発明の実施の形態】

極性脂質(リン脂質を含む)は、その二極性の性質により、湿潤剤と乳化剤と

しての商業的に非常に高い関心を集めている。またこれらの特性は、リン脂質中のHUFAの安定性を高めるのみならず、これらのバイオアベイラビリティーを向上させるのにも役立ち得る。これらの特性により、リン脂質は、栄養補助製品、食品、子供用食品、と医薬用途で使用するための理想的形体の成分となっている。

【0018】

本発明者等は、極性脂質が高濃度のアルコール(例えば68%を超えるアルコ ール濃度)だけでなく、低濃度のアルコール(アルコール約35%未満)にも非 常に良く溶けることを、思いがけず発見した(図1)。本発明の目的のために、 リン脂質は、本願明細書中で記載されるタイプの設備によって遠心分離にかけた 時に固まったり連続相(しばしば上清または軽相とも呼ばれる)から分離したり しない場合、「可溶性」であると記載される。アルコール約35~約68重量% のアルコール濃度範囲では、極性脂質は非常に低い可溶度を示す。本発明は、極 性脂質のこの特徴(低アルコール濃度での高い可溶度/分散度)を利用し、これ を幾つかの方法で利用して、天然バイオマテリアルから極性脂質(特にリン脂質) を低コストで抽出と回収するためのプロセスを開発することができる。

[0019]

HUFA含有極性脂質に富む天然のバイオマテリアルとしては、魚、甲殻類、 微生物、卵、脳組織、牛乳、肉、と脂肪種子を含む植物物質が挙げられる。本明 細書において「魚、甲殻類、微生物、卵、脳組織、牛乳、肉、と脂肪種子を含む 植物物質」という用語は、これらを遺伝子改変したものも含むものとする。これ らの物質中のリン脂質の含有量は一般には低く、通常は0.1~約4湿量%程度 である。その結果、これらのリン脂質を回収するには大量の物質を処理すること が必要である。従来の抽出技術はコストが高くつくため、リン脂質と特にHUF A富化リン脂質は非常に高価であり、そのため子供用食品、医薬品と化粧品産業 での使用に限られていた。本発明の利点の1つは、極性脂質(特にリン脂質)を 費用効率のよい方法で抽出することである。

[0020]

本発明のプロセスの1つの実施形態の第1ステップにおいて、低油含有物質が

選択されるか、または該物質を好適な脱油プロセスによって(ただし好ましくは タンパク質の変性を生じさせない脱油プロセスによって)脱油する。これには、 高温(約65℃を超える)または高濃度溶媒(例えば約50%を超える)を用い ないプロセスが含まれる。好ましくは、国際特許出願公開番号第WO96/05 278号(米国特許第5,928,696号)に概説されている脱油プロセスが 用いられる。好ましくは、この脱油プロセスに解決の鍵となる変更が加えられる 。本発明者等は、アルコールと水を加える前にバイオマテリアルをホモジナイズ すること、またはアルコールと水を加える前にホモジナイズをすることによって 、ただし最も好ましくはアルコールと水を加える前後両方でホモジナイズをする ことによって、ホモジナイズを行わないものに比べて油回収率が最大で85%改 善されることを、思いがけず発見した(図2)。本明細書中において、「ホモジ ナイズ」とは、圧力下において該混合物を小さなオリフィスに通過させたりコロ イダルミルを用いる等の高速剪断プロセス、または他の高速剪断プロセス等を含 む。好ましくは、混合物を小さなオリフィスから押し出す場合、ホモジナイズは 、約100バール~約1000バール、より好ましくは約150~約350バー ルの圧力で行われる。これは思いがけない結果である。というのは、当業者はこ のタイプの混合物をホモジナイズすれば、非常に破壊しにくい非常に強いエマル ションが形成されてプロセス効率を下げると思うからである。

[0021]

プロセス全体を通して低濃度のアルコールを用いるレシチン回収プロセスの概 要が図3に記載されている。この実施例では極性脂質リッチバイオマテリアルと して液体卵黄を用いる。しかし、他の極性脂質含有バイオマテリアル(例えば魚 、甲殻類、微生物、脳組織、牛乳、肉、と脂肪種子を含む植物物質等)は、この プロセスを若干改良して同様の方法で処理することもできることを理解されたい 。このプロセスの第1ステップでは、この物質を、任意の周知の脱油プロセスに よって(だだし好ましくはタンパク質の変性を生じさせない脱油プロセスによっ て)脱油する。より効率良く油を回収するためには、そのバイオマテリアルの中 の遊離油と同様に脂肪含有細胞状粒子中の油を分離することができるように、ホ モジナイズによって物質を剪断して該脂肪含有細胞状粒子を破壊する。次にアル

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コールと水を卵黄に加えて、この混合物を再びホモジナイズする。この水溶液中 のアルコール濃度は約5~約35重量%、好ましくは約20~約35重量%、と 最も好ましくは約25~約30重量%であってもよい。次にこの遊離油を遠心力 によって密度の差により分離する。これによって以下の2つの画分が回収される :(1)約50~70%のタンパク質(乾燥重量%)と約30~50乾燥重量% の極性脂質を含む画分、この混合物は卵黄に比べてコレステロール含有量が非常 に低い;と(2)その卵黄のトリアシルグリセロールの約85%を含む卵油。該 タンパク質/レシチン画分に低濃度アルコールを更に混ぜると、レシチンが分散 し、これをその後遠心力によってタンパク質から分離する。タンパク質とレシチ ン生成物の向流洗浄/遠心分離または逆流洗浄/分離を用いて、生成物の純度と プロセス全体の経済的な面を向上させることができる。このプロセスではタンパ ク質は変性せず、(その機能性のおかげで)このプロセスの副産物としての高い 再販価値を保持する。これにより、生成される全ての生成物の全体的なコストを 下げる。

[0022]

このプロセスにおいて必要とされる設備は単純であるため、このプロセス全体 は、低酸素雰囲気(例えばこのプロセスの好適な実施形態においては窒素)下で 非常に簡単に行うことができ、さらに該極性脂質中のHUFAを酸化から保護す る。例えば、気密デカンタを用いてこの混合物から油を分離することができる。 好適なデカンタは、ドイツ、エルデのWestfalia Separator Industry GmbH から 入手可能なCA226-28Gas Tightモデルであり、このモデルは、遠心分離場(にお いて固体含有量の多い懸濁液から油を連続的に分離することができる。タンパク 質から極性脂質を分離するために有用な気密分離装置は、ドイツ、エルデのWest falia Separator Industry GmbH から入手可能なSC6-06-576 Gas Tightモデルで あり、このモデルは、遠心分離場において固体含有量の多い懸濁液から固体を連 続的に分離することができる。

[0023]

また、このプロセスの改良されたバージョンも開発された。このプロセスにお いて、低アルコール濃度を用いた脱油とレシチン洗浄ステップは、先に概説した

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プロセスに似ている。ただし、レシチン相を乾燥した後は、該レシチン相は濃縮 アルコールで洗浄される。タンパク質は高濃度のアルコールに溶けないので、こ れらは沈殿し(レシチンは溶けるが)、沈殿したタンパク質を(例えば重力また は遠心力を用いた)密度分離によって分離する。次にタンパク質低減レシチンを 水とアルコールの蒸発によって濃縮する。このプロセスのバリエーションの利点 は、高品質と低品質のレシチン画分の両方を産生するための選択肢を提供するこ と、と高品質のレシチンを提供する際に、該タンパク質のごく一部しか変性しな いことである。

[0024]

またこのプロセスを、脱油ステップの後に高濃度のアルコールを用いるために も改良した。バイオマテリアルを脱油した後の処理ステップは、低アルコール濃 度プロセスとほぼ同様であるが、希釈アルコールの代わりに濃縮アルコールを加 える。脱油の後、極性脂質/タンパク質即席産物の濃縮と乾燥が行われる。濃縮 /乾燥ステップは、極性脂質を再び溶解するために加える必要がある濃縮アルコ ールの量を減少させるために必要である。乾燥極性脂質/タンパク質相を濃縮ア ルコールで洗浄し、タンパク質を沈殿させる。沈殿したタンパク質を、(重力ま たは遠心力を用いた)密度分離により、向流洗浄システムにおいて分離する。ア ルコールと水の蒸発によって、タンパク質低減極性脂質を濃縮する。このプロセ スの利点は、必要とされる熱エネルギーが低いことである。主な欠点は、該タン パク質の全てが変性し、価値が低くなることである。

[0025]

理論で限定しようとする訳ではないが、上記プロセスの根底にあるメカニズム の幾つかについては以下にさらに詳しく記載されるものと考えられる。ホモジナ イズに関して、細胞物質の破壊はここで起こるものと思われる。目的は、全ての 成分を均質に分配すること、すなわち均質な多分散系(タンパク質、油、リポタ ンパク質、連続相水)を作製することであって、該多分散系は、水性もしくは純 粋アルコールを加えたときに、局所的な不可逆的タンパク質変性を生じさせるこ となく、ただちに均一に(即ち均質に)分配されることができるようなものであ る。温度は、油相に溶けるレシチンの量をできるだけ少なくするために、可能な

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限り低く保たれる。タンパク質の1次と2次構造を破壊せずに4次と3次構造を 破壊するためには、ホモジナイズプロセスで用いられる圧力は、好ましくは10 00バール未満、とより好ましくは600バール未満でなければならない。アル コール濃度は好ましくは30重量%、より好ましくは約28%である。アルコー ルが過度に低いと、タンパク質の膨張がひどくなり、小さな遊離脂肪球がタンパ ク質中に取り込まれることもある。リポタンパク質の形態で結合した脂肪は極性 脂質(リン脂質)の放出を妨げないので、その割合はここでは詳しく考慮しない

[0026]

原則として、アルコール濃度が高ければ、タンパク質の収縮は増すが、水性相 がより非極性に近い程、より極性の高い脂質が該油相中に溶解すると考えられる 。したがって、適度な濃度と温度は、例えば少数の予備実験(遠心分離テスト) を各物質毎に行うことによって、見つけなければならない。

[0027]

物質の天然の水分含有量を考慮に入れると、水性アルコールを加えて、約25 ~30%の好適な最終濃度のアルコールを作製し、分散液を再びホモジナイズす る。収縮したタンパク質分子と脂肪液滴は互いに分離する。こうしてこれらの中 間にある中間相(脂肪球の表面に存在する極性脂質層)を破壊する。従って、油 は該分散液中に遊離相としてより存在し易くなる。一方ではこの油中水エマルシ ョンにおいて平衡を確立するために、極性脂質は再び脂肪球の周りを取り囲み、 また他方では油滴が凝集してより大きな油滴となる。このため、遠心分離場の追 加的力が用いられる。その後、今や大きくなった油滴は合体する(すなわち分離 可能な連続相を形成する)ことができる。

[0028]

ホモジェナイザーを用いた手法は、これによって非常に小さな油滴が生成され るので、当業者には驚くべきことである。過去の方法では、エマルションの等級 が大きな内部表面積によって大きくなるので、分離する前に油滴のサイズを小さ くすることはしなかった。反対に、油が凝集して大きな油滴になるように攪拌ま たは混練を慎重に行っていた。中でも、粘度も減少させるために、この捏揉プロ

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セスにおいて熱が有用であった。約300パール以上にホモジナイズ圧力を増加 させることにより、より多くの油を分離することもできるという驚くべき効果は 、タンパク質、極性脂質と油(実際には非極性脂質相)と溶媒相との相互作用に より説明することができる。

[0029]

従って、油分離が必ず生じ、これにより一般には(剪断により破壊された)液 滴の表面張力と表面状態がその元の平衡を取り戻す。これは、ホモジナイズした スラリーを好ましくはすぐに密度分離装置(好ましくは適当な設計と幾何学的考 慮のなされた遠心分離機)に入れ、そこで非極性脂質(油)と、タンパク質、水 とアルコールを含む極性脂質とに分離することを意味する。粘度の低下は等級に は必要ないが、ホモジナイズを行わない油回収では必要である(国際特許出願公 開WO96/05278号に記載)。ホモジナイズしたスラリーを遠心分離場に 直接移しかえることは、この融合を助けるために重要であろう。

[0030]

好ましくはデカンタ(遠心分離機を含む他のタイプの密度分離装置もこの目的 のために首尾良く使用される)で1段階もしくは2段階で油を分離した後、理想 的には、後でタンパク質相の中で水でアルコール濃度を低下させたときに、該混 合物の極性が増大し、これによりレシチンが遊離水/アルコール相の中で再び結 合して油が「放出」するが、該遊離水/アルコール相の中には油滴が見られない ように、全ての遊離油画分(脂質と非極性脂質)を分離する。通常は、アルコー ル濃度が低下すると(すなわち油可溶度が該極性脂質相においてなくなると)、 この極性脂質/タンパク質/アルコール混合物の中の油が遊離する。驚くべきこ とに、2倍のホモジナイズと遠心分離の後に、アルコール濃度がたった15%で あっても、ほんの僅かな遊離油しか遠心分離することができないことが分かった

[0031]

コレステロールを含むステロールは、油相よりも極性脂質相に対して大きな親 和性を有し得る。これにより、油相よりも極性脂質相の中のステロール含有量が 高くなる。油もしくは極性脂質相中へのステロールの移動は、混合物のpHを変 える、温度を変更する、または該水性相の極性を増減するために塩等の加工助剤 を加えることにより、操作することができる。極性脂質リッチ画分の中のコレス テロールを減らすための他の方法は、コレステロールを全くもしくは殆ど含まな い油を極性脂質リッチ画分に加えて脱油プロセスを繰り返すことである。このよ うにして、コレステロールを油相中に分離することができる。

【0032】

【実施例】

実施例1

低アルコール抽出プロセス:100kgの液体卵黄(42kg乾燥物質を含む)をホモジナイズした後、エタノール(純度96%、35.4kg)と水(30 .7kg)をこの卵黄に加えた。得られたアルコール濃度は全体で約20重量% (アルコールと水のみに関しては27重量%)であった。次にこの混合物を再び ホモジナイズし、デカンタ遠心分離機を用いて該混合物を遠心分離にかけ、油相 とアルコール/水相を生成した。この脱油ステップにより、17kgの卵黄油と 149kgのアルコール/水相ができた。次に、このアルコール/水相を、セパ レータ遠心分離機を用いた向流洗浄プロセスを用いて、同じ低濃度のアルコール で3回洗浄した。このプロセスにより、2つの画分が得られた:(1)リン脂質 リッチ画分(脂質相)、これを乾燥すると全部で17kgの乾燥物質(リン脂質) 8kgを含む)を含む生成物が得られた;と(2)タンパク質リッチ画分、これ を乾燥すると、12kgの乾燥物質(タンパク質11kgとリン脂質0.3kg を含む)が得られた。卵黄1つあたり平均重量約16.0g(それぞれ卵黄1つ あたりリン脂質約1.7gを含む)として、卵黄100kgで約10.6kgの リン脂質が得られる。このプロセスによってリン脂質リッチ画分中で回収される リン脂質8.0kgは、約76%のリン脂質画分の回収効率である。

【0033】

実施例2

高アルコールのポリッシングステップを伴う低アルコール抽出プロセス:10 0kgの液体卵黄(42kg乾燥物質を含む)をホモジナイズした後、エタノー ルと水を加えて該混合物をアルコール/水相中の最終アルコール濃度30重量%

とした。次にこの混合物を再びホモジナイズし、デカンタ遠心分離機を用いて該 混合物を遠心分離にかけ、油相とアルコール/水相を生成した。この脱油ステッ プにより、16kgの卵黄油と134kgのアルコール/水相(26kgの乾燥 物質を含む)ができた。次に、72kgのエタノールと170kgの水をこのア ルコール/水相に加えて、これを混合し、セパレータ遠心分離機で遠心分離にか けた。これにより、2つの画分が得られた: (1) 11kgの乾燥物質を含む脂 質相(299kg);と(2)15kgの乾燥物質を含む固相(78kg)。画 分1は少量のタンパク質とリン脂質とを含んでおり、画分2は主にタンパク質を 含んでいた。次に画分1を乾燥して重量11.2kgとし、20kgのエタノー ル(96%)をこの画分に加えた。次にこの混合物をセパレータ遠心分離機で処 理し、10kgの乾燥物質を含む液相を得た。次にこの液相を乾燥して最終的な 重量を10.5 kgとした(10.0 kg乾燥物質-リン脂質画分)。画分2の 中の固体78kgも乾燥して、全量16kg(または乾燥物質15kgータンパ ク質画分)とした。卵黄1つあたり平均重量約16.0g(それぞれ卵黄1つあ たりリン脂質約1.7gを含む)として、卵黄100kgで約10.6kgのリ ン脂質が得られる。このプロセスで回収されるリン脂質10.0kgは、約90 %を超えるリン脂質画分の最小回収効率である。

[0034]

実施例3

高アルコール極性脂質抽出プロセスを伴う低アルコール脱油プロセス:100 kgの液体卵黄(45kg乾燥物質を含む)をホモジナイズした後、エタノール と水を加えて、該混合物をアルコール/水相中の最終アルコール濃度30重量% とした。次にこの混合物を再びホモジナイズし、デカンタ遠心分離機を用いて該 混合物を遠心分離にかけ、油相とアルコール/水相を生成した。この脱油ステッ プにより、17kgの卵黄油と139kgのアルコール/水相(28kgの乾燥 物質を含む)ができた。次にこのアルコール/水相を乾燥し(109kgのアル コールと水を回収)、物質30kgを得た(28kg乾燥物質)。エタノール(純度96%)90kgをこの物質に加え、この混合物をセパレータ遠心分離機で 処理して、液相(リン脂質を含む)とタンパク質を含む固相を得た。液相(全量 80kg、乾燥物質10.4kgを含む)を乾燥して、乾燥物質(リン脂質)1 0.4kgを含む生成物10.6kgを得た。

固相(全量40kg)を乾燥して、18.5kgの生成物、つまりタンパク質(乾燥物質17.6kgを含む)を得た。卵黄1つあたり平均重量約16.0g(それぞれ卵黄1つあたりリン脂質約1.7gを含む)として、卵黄100kgで約10.6kgのリン脂質が得られる。このプロセスで回収されるリン脂質10.0kgは、約90%を超えるリン脂質両分の最小回収効率である。

[0035]

本発明は、様々な実施形態において、本明細書中に実質的に説明と記載された 成分、方法、プロセス、システムと/または装置、例えば様々な実施形態、サブ コンビネーション、とこれらのサブセット等を含む。当業者であれば、本明細書 の開示内容を理解すれば、本発明をどのように実施と使用すべきかが分かるであ ろう。本発明は、様々な実施形態において、本明細書中またはこれらの様々な実 施形態に説明と/または記載されていない事項を含まない装置とプロセスの提供 も含む。例えば、性能を高めたり、簡単にしたり、と/または実施コストを削減 したりするために従来装置またはプロセスにおいて用いられるようなこのような 事項を含まない場合が挙げられる。

【0036】

本発明についてのこれまでの説明は、例示と説明のために提供された。これま での記載は、本発明を本明細書中に記載された形態に限定するものではない。本 発明の説明は、1以上の実施形態ならびにそれをある種改良/改変したものの説 明を含んでいるが、他の改良と変更が本発明の範囲内に含まれる(例えば本明細 書の開示内容を理解した後の当業者の能力と知識の範囲内にあるものなど)。許 容される程度に他の実施形態(例えば、特許請求の範囲に記載されたものに代わ る、これと交換可能なと/またはこれと同等な構造、機能、範囲またはステップ 等。このような代替的な、交換可能なと/または同等な構造、機能、範囲または ステップが本明細書中に開示されているか否かにかかわらない)を含む権利を得 るものとする。また、任意の特許を受けることができる主題に公然と限定とする ものではない。 【図面の簡単な説明】

【図1】

アルコール濃度の関数としての、リン脂質(極性脂質の形態)の可溶度を表す グラフである。

【図2】

卵黄の脱油に対するホモジナイズ化の影響を表すグラフである。

【図3】

低濃度アルコールに基づく(極性脂質抽出プロセスの例としての)リン脂質抽 出プロセスを表すグラフである。

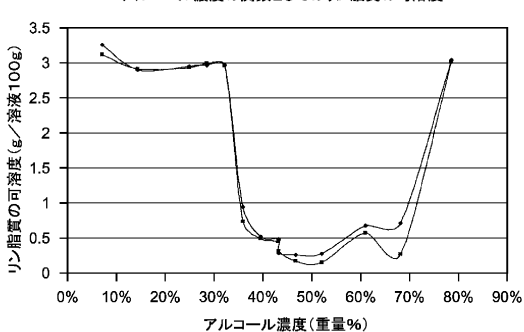
【図4】

低濃度アルコールに基づく(極性脂質抽出プロセスの例としての)リン脂質抽 出プロセスであって、高濃度アルコールを用いたステップを伴うリン脂質のポリ ッシングステップを追加した場合の結果を表すグラフである。

【図5】

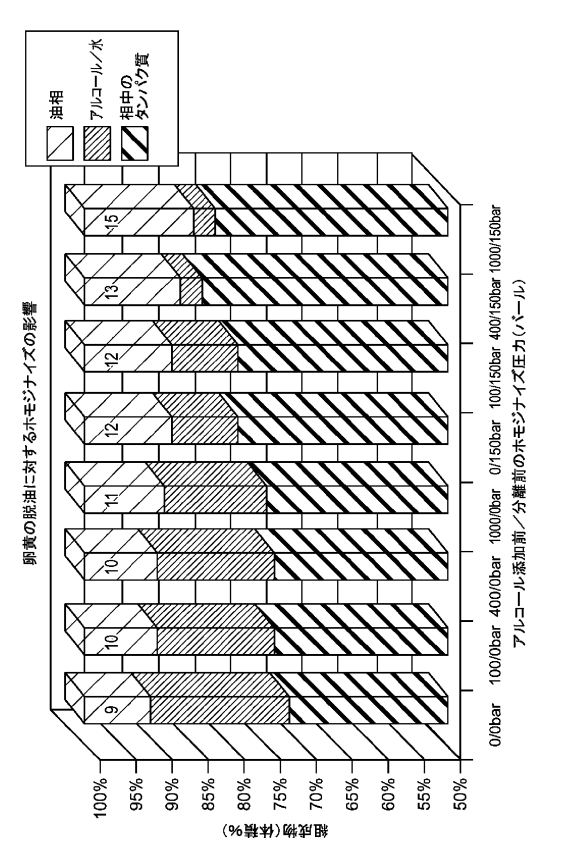
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抽出プロセスのリン脂質回収部分を通して高濃度アルコールの使用に基づく(極性脂質抽出プロセスの例としての)リン脂質抽出プロセスを表すグラフである 図1】



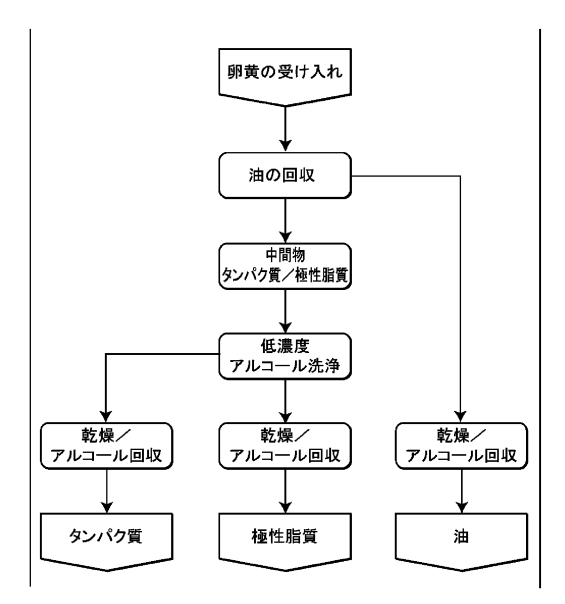
アルコール濃度の関数としてのリン脂質の可溶度

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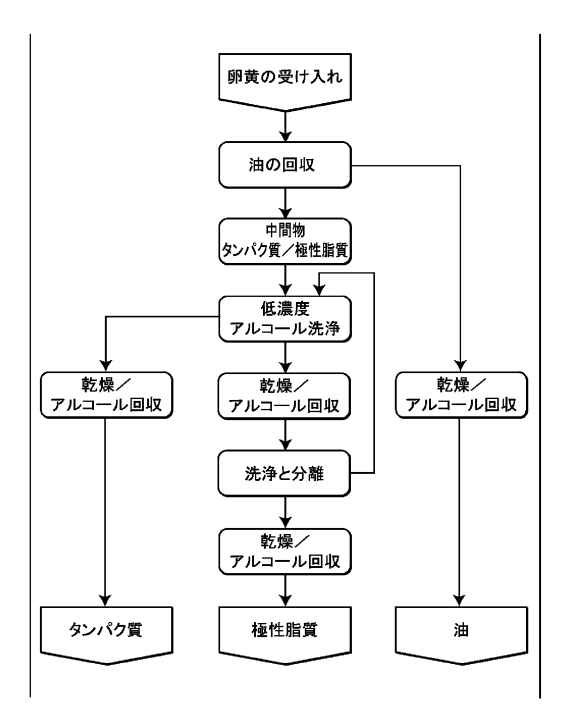
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脱油と極性脂質回収段階の双方に対して 低濃度アルコールを用いた極性脂質抽出プロセス



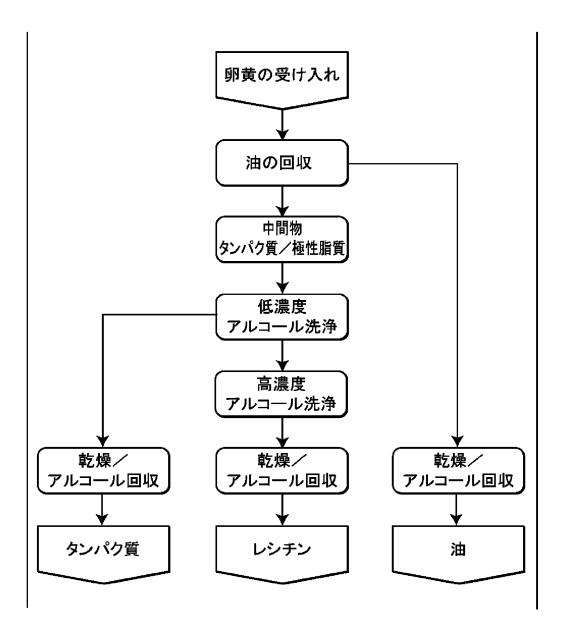
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脱油と極性脂質回収段階の双方に対して 低濃度アルコールを用いた極性脂質抽出プロセス



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脱油段階では低濃度アルコールを用い、極性脂質回収段階では 高濃度アルコールを用いる極性脂質抽出プロセス



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| Forth PCT/SA | 210 (sontinuation of execute these) (July 1992) | |

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page 2 of 2

INTERNATIONAL SEARCH REPORT

| Box I Observations where certain claims were found unsearchable (Continu | ation of item 1 of first sheet) |
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| This International Search Report has not been established in respect of certain claims under A | uticle 17(2)(a) for the following reasons: |
| Claims Nos.:
because they relate to subject matter not required to be searched by this Authority, n | arnely: |
| Claims Nos.:
because they relate to parts of the International Application that do not comply with th
an extent that no meaningful International Search can be carried cut, specifically: | ne prescribed requirements to such |
| 3. Ciaims Nos.:
because they are dependent claims and are not drafted in accordance with the second | nd and third sentences of Rule 6.4(a). |
| Box II Observations where unity of invention is lacking (Continuation of item | 2 of first sheet) |
| This International Searching Authority found multiple inventions in this international application | n, as follows: |
| see additional sheet | |
| 1. χ As all required additional search less were timely paid by the applicant, this Internation searchable claims. | onal Search Report covers all |
| As all searchable claims could be searched without effort justifying an additional fee. of any additional fee. | this Authority did not invite payment |
| 3. As only some of the required additional search fees were timely paid by the applicant covers only those claims for which fees were paid, specifically claims Nos.: | t, this International Search Report |
| 4, No required additional search fees were timely paid by the applicant. Consequently, restricted to the invention first mentioned in the claims; it is covered by claims Nos.: | this International Search Report is |
| Remark on Protest The additional search fees were | accompanied by the applicant's protest. |

Form PCT/ISA/210 (continuation of first sheet (1)) (July 1998)

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210 This International Searching Authority found multiple (groups of) inventions in this international application, as follows: 1. Claims: 1-48,50 Process for fractionating a material which contains polar lipids comprising the steps of blending said material with a water soluble organic solvent followed by the use of differential density to separate light and heavy phases. 2. Claim : 49 Method for cholesterol reduction in polar lipid fractions comprising the steps of adding an oil which has little or no cholesterol to a polar lipid fraction and removing the oil from the polar lipid fraction with a concommitant reduction in cholesterol.

| | | ATIONAL SEAR(| | | | al Application No
01/00841 |
|---|---|------------------|--|---|--|--|
| Patent document
cited in search report | | Publication date | | Patent family
member(s) | PC1/18 | Publication
date |
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Form PCT/ISA/210 (patent family annax) (July 1992)

| (51) Int. Cl. ⁷ 識別記号 | |
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| (81)指定国 EP(AT, BE, CH, CY, | |
| DE, DK, ES, FI, FR, GB, GR, IE, I | |
| T, LU, MC, NL, PT, SE, TR), OA(BF | |
| , BJ, CF, CG, CI, CM, GA, GN, GW, | |
| ML, MR, NE, SN, TD, TG), AP(GH, G | |
| M, KE, LS, MW, MZ, SD, SL, SZ, TZ | |
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| VN, YU, ZA, ZW | |
| (72)発明者 ラッセンフォーフェル, ユルゲン | |
| ドイツ, 59302 エルデ, パッペルヴ | |
| ク 9 | |
| (72)発明者 ヴィト, ヴィリ | |
| ドイツ, 49545 テックレンブルク, | |
| クリーヴク 34 | |
| Fターム(参考) 4C086 AAO4 DA40 ZC22 | |
| 4D056 AB12 AB14 AC06 BA09 CA26 | |
| CA28 DA01 DA02 DA05 DA10 | |
| 4H059 AA04 BA12 BA33 BA83 BB02 | |
| BB03 BC03 BC05 BC06 BC43 | |
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| Electronic Acknowledgement Receipt | | | | | |
|--------------------------------------|-------------------------------------|--|--|--|--|
| EFS ID: | 12113042 | | | | |
| Application Number: | 12057775 | | | | |
| International Application Number: | | | | | |
| Confirmation Number: | 1945 | | | | |
| Title of Invention: | BIOEFFECTIVE KRILL OIL COMPOSITIONS | | | | |
| First Named Inventor/Applicant Name: | Inge Bruheim | | | | |
| Customer Number: | 72960 | | | | |
| Filer: | John Mitchell Jones/Vickie Hoeft | | | | |
| Filer Authorized By: | John Mitchell Jones | | | | |
| Attorney Docket Number: | NATNUT-14409/US-5/ORD | | | | |
| Receipt Date: | 21-FEB-2012 | | | | |
| Filing Date: | 28-MAR-2008 | | | | |
| Time Stamp: | 16:18:36 | | | | |
| Application Type: | Utility under 35 USC 111(a) | | | | |

Payment information:

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Inge Bruheim, et alConfirmation:1945Serial No.:12/057,775Group No.:1651Filed:03-28-2008Examiner:Ware, Deborah K.Entitled:BIOEFFECTIVE KRILL OIL COMPOSITIONSFiled:

INFORMATION DISCLOSURE STATEMENT LETTER

EFS Web Filed Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir or Madam:

The citations listed in the attached **IDS Form SB08A** may be material to the examination of the aboveidentified application, and are therefore submitted in compliance with the duty of disclosure defined in 37 C.F.R. §§ 1.56 and 1.97.

Applicants wish to bring to the Examiner's attention that we are not providing copies of US Patents as instructed under 37 CFR 1.98(a)(2). The Examiner is requested to make these citations of official record in this application.

Applicants wish to bring to the Examiner's attention that the references supplied in this IDS are from the December 8, 2011 Office Action from related KR Patent Application No. 10-2010-7006897.

This Information Disclosure Statement under 37 C.F.R. §§ 1.56 and 1.97 is not to be construed as a representation that a search has been made, that additional information material to the examination of this application does not exist, or that any one or more of these citations constitutes prior art.

The Commissioner is hereby authorized to charge any required fees or credit any overpayments to Attorney Deposit Account No.: **50-4302**, referencing Attorney Docket No.: **NATNUT-14409/US-5/ORD**.

Dated: February 20, 2012

<u>/J. Mitchell Jones/</u> J. Mitchell Jones Registration No. 44,174 CASIMIR JONES, S.C. 2275 Deming Way, Suite 310 Middleton, WI 53562 608.662.1277 Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

| Application Number | | 12057775 |
|----------------------|--------|-----------------------|
| Filing Date | | 2008-03-28 |
| First Named Inventor | Inge E | Bruheim |
| Art Unit | | 1651 |
| Examiner Name | Ware, | Deborah K. |
| Attorney Docket Numb | er | NATNUT-14409/US-5/ORD |

| | | | | | U.S. | PATENTS | | | Remove | |
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INFORMATION DISCLOSURE Application Number 12057775 Filing Date 2008-03-28 First Named Inventor Inge Bruheim Art Unit 1651 Examiner Name Ware, Deborah K. Attorney Docket Number NATNUT-14409/US-5/ORD

| | 2 | 07/080 | 0515 | WO | | 2007-07-19 | Aker Biomarine ASA | | |
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| | Application Number | | 12057775 |
|--|----------------------|--------|-----------------------|
| | Filing Date | | 2008-03-28 |
| INFORMATION DISCLOSURE | First Named Inventor | Inge E | Bruheim |
| STATEMENT BY APPLICANT
(Not for submission under 37 CFR 1.99) | Art Unit | | 1651 |
| | Examiner Name | Ware, | , Deborah K. |
| | Attorney Docket Numb | er | NATNUT-14409/US-5/ORD |

| CERTIFICATION | STATEMENT |
|---------------|-----------|
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Please see 37 CFR 1.97 and 1.98 to make the appropriate selection(s):

That each item of information contained in the information disclosure statement was first cited in any communication from a foreign patent office in a counterpart foreign application not more than three months prior to the filing of the information disclosure statement. See 37 CFR 1.97(e)(1).

OR

That no item of information contained in the information disclosure statement was cited in a communication from a foreign patent office in a counterpart foreign application, and, to the knowledge of the person signing the certification after making reasonable inquiry, no item of information contained in the information disclosure statement was known to any individual designated in 37 CFR 1.56(c) more than three months prior to the filing of the information disclosure statement. See 37 CFR 1.97(e)(2).

See attached certification statement.

X The fee set forth in 37 CFR 1.17 (p) has been submitted herewith.

A certification statement is not submitted herewith.

SIGNATURE

A signature of the applicant or representative is required in accordance with CFR 1.33, 10.18. Please see CFR 1.4(d) for the form of the signature.

| Signature | /J. Mitchell Jones/ | Date (YYYY-MM-DD) | 2012-01-24 |
|------------|---------------------|---------------------|------------|
| Name/Print | J. Mitchell Jones | Registration Number | 44174 |

This collection of information is required by 37 CFR 1.97 and 1.98. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 1 hour to complete, including gathering, preparing and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. **SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450**.

The Privacy Act of 1974 (P.L. 93-579) requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b)(2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

The information provided by you in this form will be subject to the following routine uses:

- The information on this form will be treated confidentially to the extent allowed under the Freedom of Information Act (5 U.S.C. 552) and the Privacy Act (5 U.S.C. 552a). Records from this system of records may be disclosed to the Department of Justice to determine whether the Freedom of Information Act requires disclosure of these record s.
- 2. A record from this system of records may be disclosed, as a routine use, in the course of presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
- 4. A record in this system of records may be disclosed, as a routine use, to a contractor of the Agency having need for the information in order to perform a contract. Recipients of information shall be required to comply with the requirements of the Privacy Act of 1974, as amended, pursuant to 5 U.S.C. 552a(m).
- 5. A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
- 6. A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (i.e., GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
- 8. A record from this system of records may be disclosed, as a routine use, to the public after either publication of the application pursuant to 35 U.S.C. 122(b) or issuance of a patent pursuant to 35 U.S.C. 151. Further, a record may be disclosed, subject to the limitations of 37 CFR 1.14, as a routine use, to the public if the record was filed in an application which became abandoned or in which the proceedings were terminated and which application is referenced by either a published application, an application open to public inspections or an issued patent.
- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.

| Electronic Patent / | App | olication Fee | e Transmi | ttal | |
|---|-----|--------------------|---------------|--------|-------------------------|
| Application Number: | 120 | 057775 | | | |
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File Listing: Document File Size(Bytes)/ Multi Pages **Document Description File Name** Number Message Digest Part /.zip (if appl.) 73326 14409US5IDSLetter01242012. 1 **Transmittal Letter** no 1 pdf fa62420ce082e95fcd8bf3867ed55ff79d97 98b Warnings: Information: 1831904 2 **Foreign Reference** JPA2004534800.pdf no 60 01b0eb7d7150ad8436d4ab1c6ff7e6796d 8a34e Warnings: Information: 1126330 3 **Foreign Reference** WO2007080515.pdf 27 no 14a09da6f685f2301096da43381d3f9e2b de013 Warnings: Information: 2606929 4 Non Patent Literature BUDZINSKI1985.pdf 51 no dea47d9074c0ec8bab93e6ad83f9ef4ae5. c099 Warnings: Information: 97505 5 Non Patent Literature Bunea2004print.pdf 9 no 44f351c4bf6f11a65d72a0891f26ef366741b Warnings: Information: 473015 6 Non Patent Literature SIKORSKI.pdf 11 no 3415d6af73aed943aa6d2bfe78ea2413416 8714c Warnings: Information: 612564 Information Disclosure Statement (IDS) 7 14409US5ORDIDS01242012.pdf no 4 Form (SB08) 25fdb919eb9d44342fc47c97df4005df3d3fl 413 Warnings: Information: 366063 8 Non Patent Literature Gordeev1990.pdf 5 no 4e7b4812c0af3939a51244e75fa282c670a RIMFROST EXHIBIT 1024

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Inge Bruheim, et alConfirmation:1945Serial No.:12/057,775Group No.:1651Filed:03-28-2008Examiner:Ware, Deborah K.Entitled:BIOEFFECTIVE KRILL OIL COMPOSITIONSFiled:

INFORMATION DISCLOSURE STATEMENT LETTER

EFS Web Filed Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir or Madam:

The citations listed in the attached **IDS Form SB08A** may be material to the examination of the aboveidentified application, and are therefore submitted in compliance with the duty of disclosure defined in 37 C.F.R. §§ 1.56 and 1.97.

Applicants wish to bring to the Examiner's attention that we are not providing copies of US Patents as instructed under 37 CFR 1.98(a)(2). The Examiner is requested to make these citations of official record in this application.

This Information Disclosure Statement under 37 C.F.R. §§ 1.56 and 1.97 is not to be construed as a representation that a search has been made, that additional information material to the examination of this application does not exist, or that any one or more of these citations constitutes prior art.

The Commissioner is hereby authorized to charge any required fees or credit any overpayments to Attorney Deposit Account No.: **50-4302**, referencing Attorney Docket No.: **NATNUT-14409/US-5/ORD**.

Dated: January 24, 2012

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(54) 【発明の名称】心臓血管疾患、関節炎、皮膚ガン、糖尿病、月経前症候群および経皮送達の予防および/または 治療のためのオキアミおよび/または海洋生物の抽出物

(57)【要約】

本発明は心臓血管疾患、慢性関節リウマチ、皮膚ガン、糖尿病、月経前症候群および経皮 送達増強の予防および/または治療に関する。本発明の方法は療治効果的な量のオキアミ 油および/または海洋生物油を患者に投与することを含む。本発明はまた、これら疾患の 予防および/または治療のための組成物に関する。

【特許請求の範囲】

【請求項1】

患者におけるコレステロールを低下させるための組成物であって、効果的な量のオキアミ 油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オキアミ 油および/または海洋生物油が、以下の工程:

(2)

a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れて、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し;

b)これを液体内容物と固体内容物に分離し;

c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し;

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d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 脂質画分を抽出し;

e)これを液体内容物と固体内容物に分離し;

f) 分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および

g) 固体内容物を回収する工程

からなるプロセスから得られる組成物。

【請求項2】

患者におけるコレステロールを低下させるための組成物であって、効果的な量のオキアミ 油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オキアミ 油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキサンエン酸、ホスフ ァチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスファチジル エタノールアミン、スフィンゴミエリン、α-トコフェロール、アスタキサンチンおよび フラボノイドを含む組成物。

【請求項3】

リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、β-カロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 30 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項2に記載の 組成物。

【請求項4】

患者におけるコレステロールを低下させる方法であって、請求項1~3のいずれか一項に 記載される組成物の効果的な量を前記患者に投与することを含む方法。

【請求項5】

前記投与が経口的に行われる、請求項4に記載の方法。

【請求項6】

前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範囲の量で投与される、請求項4に記載の方法。

【請求項7】

前記量が4.8グラムである、請求項6に記載の方法。

【請求項8】

患者におけるコレステロールを低下させるための、請求項1~3のいずれか一項に記載さ れる組成物の使用。

【請求項9】

患者におけるコレステロールを低下させる医薬品を製造するための、請求項1~3のいず れか一項に記載される組成物の使用。

【請求項10】

患者の動脈における血小板接着およびプラーク形成を阻害するための組成物であって、効 50

果的な量のオキアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに 含み、前記オキアミ油および/または海洋生物油が、以下の工程: a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し; b)これを液体内容物と固体内容物に分離し; c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し: d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 10 脂質画分を抽出し; e)これを液体内容物と固体内容物に分離し; f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および g) 固体内容物を回収する工程 からなるプロセスから得られる組成物。 【請求項11】 患者の動脈における血小板接着およびプラーク形成を阻害するための組成物であって、効 果的な量のオキアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに 含み、前記オキアミ油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキ 20 サンエン酸、ホスファチジルコリン、ホスファチジルイノシトール、ホスファチジルセリ ン、ホスファチジルエタノールアミン、スフィンゴミエリン、 α – トコフェロール、アス タキサンチンおよびフラボノイドを含む組成物。 【請求項12】 リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、βーカロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項11に記載 の組成物。 【請求項13】 30 患者の動脈における血小板接着およびプラーク形成を阻害する方法であって、請求項11 ~ 1 2 のいずれか一項に記載される組成物の効果的な量を前記患者に投与することを含む 方法。 【請求項14】 前記投与が経口的に行われる、請求項13に記載の方法。 【請求項15】 前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範 囲の量で投与される、請求項13に記載の方法。 【請求項16】 前記量が4.8グラムである、請求項15に記載の方法。 40 【請求項17】 患者の動脈における血小板接着およびプラーク形成を阻害するための、請求項11~13 のいずれか一項に記載される組成物の使用。 【請求項18】 患者の動脈における血小板接着およびプラーク形成を阻害する医薬品を製造するための、 請求項11~13のいずれか一項に記載される組成物の使用。 【請求項19】 患者における高血圧を防止するための予防薬組成物であって、予防効果的な量のオキアミ 油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オキアミ 油および/または海洋生物油が、以下の工程: 50

a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し; b) これを液体内容物と固体内容物に分離し; c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し; d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたは t-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 脂質画分を抽出し; e)これを液体内容物と固体内容物に分離し; 10 f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および g) 固体内容物を回収する工程 からなるプロセスから得られる予防薬組成物。 【請求項20】 患者における高血圧を防止するための予防薬組成物であって、予防効果的な量のオキアミ 油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オキアミ 油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキサンエン酸、ホスフ ァチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスファチジル エタノールアミン、スフィンゴミエリン、α-トコフェロール、アスタキサンチンおよび 20 フラボノイドを含む予防薬組成物。 【請求項21】 リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、βーカロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項20に記載 の組成物。 【請求項22】 患者における高血圧を防止するための方法であって、請求項19~21のいずれか一項に 記載される組成物の予防効果的な量を前記患者に投与することを含む方法。 30 【請求項23】 前記投与が経口的に行われる、請求項22に記載の方法。 【請求項24】 前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範 囲の量で投与される、請求項22に記載の方法。 【請求項25】 前記量が4.8グラムである、請求項24に記載の方法。 【請求項26】 患者における高血圧を防止するための、請求項19~21のいずれか一項に記載される組 40 成物の使用。 【請求項27】 患者における高血圧を防止する医薬品を製造するための、請求項19~21のいずれか一 項に記載される組成物の使用。 【請求項28】 関節炎を症状的に抑制または治療するための療治用組成物であって、療治効果的な量のオ キアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オ キアミ油および/または海洋生物油が、以下の工程: a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し;

b)これを液体内容物と固体内容物に分離し;

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c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し; d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 脂質画分を抽出し; e)これを液体内容物と固体内容物に分離し; f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および g)固体内容物を回収する工程 10 からなるプロセスから得られる療治用組成物。 【請求項29】 前記関節炎が、慢性関節リウマチおよび変形性関節症からなる群から選択される、請求項 28に記載の方法。 【請求項30】 関節炎を症状的に抑制または治療するための療治用組成物であって、療治効果的な量のオ キアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オ キアミ油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキサンエン酸、 ホスファチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスファ チジルエタノールアミン、スフィンゴミエリン、α-トコフェロール、アスタキサンチン 20 およびフラボノイドを含む療治用組成物。 【請求項31】 リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、β-カロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項30に記載 の組成物。 【請求項32】 前記関節炎が、慢性関節リウマチおよび変形性関節症からなる群から選択される、請求項 30に記載の組成物。 30 【請求項33】 患者における関節炎を症状的に抑制または治療するための方法であって、請求項29~3 2のいずれか一項に記載される組成物の療治効果的な量を前記患者に投与することを含む 方法。 【請求項34】 前記投与が経口的に行われる、請求項33に記載の方法。 【請求項35】 前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範 囲の量で投与される、請求項33に記載の方法。 【請求項36】 40 前 記 量 が 4.8 グ ラ ム で あ る 、 請 求 項 3 5 に 記 載 の 方 法 。 【請求項37】 患者における慢性関節リウマチの症状を抑制するための、またはそれを治療するための、 請 求 項 2 9 ~ 3 2 の い ず れ か 一 項 に 記 載 さ れ る 組 成 物 の 使 用 。 【請求項38】 患者における慢性関節リウマチの症状抑制薬、または治療薬を製造するための、請求項2 9~32のいずれか一項に記載される組成物の使用。 【請求項39】 患者における皮膚ガンを防止するための予防薬組成物であって、予防効果的な量のオキア ミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オキア 50

ミ油および/または海洋生物油が、以下の工程: a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し; b)これを液体内容物と固体内容物に分離し; c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し; d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたは t-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 10 脂質画分を抽出し; e)これを液体内容物と固体内容物に分離し; f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および g) 固体内容物を回収する工程 からなるプロセスから得られる予防薬組成物。 【請求項40】 患者における皮膚ガンを防止するための予防薬組成物であって、予防効果的な量のオキア ミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オキア ミ油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキサンエン酸、ホス ファチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスファチジ 20 ルエタノールアミン、スフィンゴミエリン、α-トコフェロール、アスタキサンチンおよ びフラボノイドを含む予防薬組成物。 【請求項41】 リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、β-カロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項40に記載 の組成物。 【請求項42】 皮膚ガンを防止する方法であって、請求項39~41のいずれか一項に記載される組成物 30 の療治効果的または予防効果的な量を患者に投与することを含む方法。 【請求項43】 前記投与が経口的に行われる、請求項42に記載の方法。 【請求項44】 前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範 囲の量で投与される、請求項42に記載の方法。 【請求項45】 前記量が4.8グラムである、請求項44に記載の方法。 【請求項46】 患者における皮膚ガンを防止するための、請求項39~41のいずれか一項に記載される 40 組成物の使用。 【請求項47】 患者における皮膚ガンを防止する医薬品を製造するための、請求項39~41のいずれか 一項に記載される組成物の使用。 【請求項48】 患者に皮膚に局所的に塗布する療治薬の経皮輸送を増強するための組成物であって、増強 効果的な量のオキアミ油および/または海洋生物油を薬学的に受容可能なキャリアととも に含み、前記オキアミ油および/または海洋生物油が、、以下の工程: a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ 50 て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し;

b)これを液体内容物と固体内容物に分離し;

c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し;

d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 脂質画分を抽出し;

e)これを液体内容物と固体内容物に分離し;

f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および

g) 固体内容物を回収する工程

からなるプロセスから得られる組成物。

【請求項49】

患者における皮膚に局所的に塗布する療治薬の経皮輸送を増強するための組成物であって、増強効果的な量のオキアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記オキアミ油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキサンエン酸、ホスファチジルコリン、ホスファチジルイノシトール、ホスファチ ジルセリン、ホスファチジルエタノールアミン、スフィンゴミエリン、 αートコフェロール、アスタキサンチンおよびフラボノイドを含む組成物。

【請求項50】

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リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、β-カロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項49に記載 の組成物。

【請求項51】

患者における皮膚に局所的に塗布する療治薬の経皮輸送を増強するための方法であって、 請求項48~50のいずれか一項に記載される組成物の増強効果的な量を前記患者に投与 することを含む方法。

【請求項52】

前記投与が経口的および/または局所的に行われる、請求項51に記載の方法。

【請求項53】

前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範囲の量で投与される、請求項51に記載の方法。

【請求項54】

前記量が4.8グラムである、請求項53に記載の方法。 【請求項55】

患者における皮膚に局所的に塗布する療治薬の経皮輸送を増強するための、請求項48~ 50のいずれか一項に記載される組成物の使用。

【請求項56】

患者における皮膚に局所的に塗布する療治薬の経皮輸送を増強する医薬品を製造するための、請求項48~50のいずれか一項に記載される組成物の使用。

【請求項57】

患者における皮膚に局所的に塗布する化粧品の経皮輸送を増強するための組成物であって、 増強効果的な量のオキアミ油および/または海洋生物油を薬学的に受容可能なキャリア とともに含み、前記オキアミ油および/または海洋生物油が、、以下の工程: a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し;

b)これを液体内容物と固体内容物に分離し;

c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 50

質富含有画分を回収し;

d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 脂質画分を抽出し;

e)これを液体内容物と固体内容物に分離し;

f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および

g) 固体内容物を回収する工程

からなるプロセスから得られる組成物。

【請求項58】

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患者における皮膚に局所的に塗布する化粧品の経皮輸送を増強するための組成物であって 、増強効果的な量のオキアミ油および/または海洋生物油を薬学的に受容可能なキャリア とともに含み、前記オキアミ油および/または海洋生物油が、エイコサペンタエン酸、ド コサヘキサンエン酸、ホスファチジルコリン、ホスファチジルイノシトール、ホスファチ ジルセリン、ホスファチジルエタノールアミン、スフィンゴミエリン、α-トコフェロー ル、アスタキサンチンおよびフラボノイドを含む組成物。 【請求項59】 リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ 20 チノール、カンタキサンチン、β-カロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項58に記載 の組成物。 【請求項60】 皮膚に局所的に塗布する化粧品の経皮輸送を増強するための方法であって、請求項57~ 59のいずれか一項に記載される組成物の増強効果的な量を患者に投与することを含む方 法。 【請求項61】 前記投与が経口的および/または局所的に行われる、請求項60に記載の方法。 30 【請求項62】 前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範 囲の量で投与される、請求項60に記載の方法。 【請求項63】 前記量が4.8グラムである、請求項62に記載の方法。 【請求項64】 患者における皮膚に局所的に塗布する化粧品経皮輸送を増強するための、請求項57~5 9のいずれか一項に記載される組成物の使用。 【請求項65】 患者における皮膚に局所的に塗布する化粧品の経皮輸送を増強する医薬品を製造するため の、請求項57~59のいずれか一項に記載される組成物の使用。 40 【請求項66】

患者における月経前症候群の症状を軽減させるための組成物であって、増強効果的な量の オキアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記 オキアミ油および/または海洋生物油が、、以下の工程:

a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し; b)これを液体内容物と固体内容物に分離し;

c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し;

d) 前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-50

ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 脂質画分を抽出し; e) これを液体内容物と固体内容物に分離し; f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し:および g) 固体内容物を回収する工程 からなるプロセスから得られる組成物。 【請求項67】 患者における月経前症候群の症状を軽減させるための組成物であって、増強効果的な量の 10 オキアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記 オキアミ油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキサンエン酸 、ホスファチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスフ アチジルエタノールアミン、スフィンゴミエリン、αートコフェロール、アスタキサンチ ンおよびフラボノイドを含む組成物。 【請求項68】 リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、β-カロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項67に記載 20 の組成物。 【請求項69】 患者における月経前症候群の症状を軽減させるための方法であって、請求項66~68の いずれか一項に記載される組成物の増強効果的な量を前記患者に投与することを含む方法 【請求項70】 前記投与が経口的に行われる、請求項69に記載の方法。 【請求項71】 前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範 囲の量で投与される、請求項69に記載の方法。 30 【請求項72】 前記量が4.8グラムである、請求項71に記載の方法。 【請求項73】 患者における月経前症候群の症状を軽減させるための、請求項66~68のいずれか一項 に記載される組成物の使用。 【請求項74】 患者における月経前症候群の症状を軽減させる医薬品を製造するための、請求項66~6 8のいずれか一項に記載される組成物の使用。 【請求項75】 患者における血中グルコースレベルを制御するための組成物であって、増強効果的な量の 40 オキアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記 オキアミ油および/または海洋生物油が、以下の工程: a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し; b) これを液体内容物と固体内容物に分離し; c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し; d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 50

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脂質画分を抽出し; e)これを液体内容物と固体内容物に分離し; f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および g) 固体内容物を回収する工程 からなるプロセスから得られる組成物。 【請求項76】 患者における血中グルコースレベルを制御するための組成物であって、増強効果的な量の オキアミ油および/または海洋生物油を薬学的に受容可能なキャリアとともに含み、前記 オキアミ油および/または海洋生物油が、エイコサペンタエン酸、ドコサヘキサンエン酸 、ホスファチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスフ r + i = j +ンおよびフラボノイドを含む組成物。 【請求項77】 リノール酸、α-リノール酸、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイ ン酸、ステアリン酸、コレステロール、トリグリセリド、モノグリセリド、全トランスレ チノール、カンタキサンチン、βーカロテン、亜鉛、セレン、ネルボン酸、ナトリウム、 カリウムおよびカルシウムからなる群の少なくとも1つをさらに含む、請求項76に記載 の組成物。 【請求項78】 患者における血中グルコースレベルを制御するための方法であって、請求項75~77の いずれか一項に記載される組成物の増強効果的な量を前記患者に投与することを含む方法 【請求項79】 前記投与が経口的に行われる、請求項78に記載の方法。 【請求項80】 前記オキアミ抽出物および/または海洋生物抽出物が1日あたり1グラム~4.8グラムの範 囲の量で投与される、請求項78に記載の方法。 【請求項81】 前記量が4.8グラムである、請求項80に記載の方法。 【請求項82】 患者における血中グルコースレベルを制御するための、請求項75~77のいずれか一項 に記載される組成物の使用。 【請求項83】 患者における血中グルコースレベルを制御する医薬品を製造するための、請求項75~7 7のいずれか一項に記載される組成物の使用。 【発明の詳細な説明】 【技術分野】 [0001]本発明は、いくつかの疾患を予防および/または治療することができる、オキアミおよび /または海洋生物に由来する多用途な療治用の抽出物に関する。 【背景技術】 [0002]オキアミは、特に南極水域において密集した群で群がる小さいエビ様甲殻類(しかしなが らエビとは異なる)に対する一般名である。オキアミは、重要なタンパク質源として、魚 類、ある種の鳥類、そして特にヒゲクジラには最も重要な食物源の1つである。オキアミ はまた、その健康上の利点がよく知られているω-3脂肪酸の良好な供給源でもある。 [0003]オキアミおよび/または海洋生物の酵素を、感染症、炎症、ガン、HIV/AIDS、痛み、ポリ プ、いぼ、痔、プラーク、しわ、薄毛、アレルギー性のかゆみ、接着不全、眼病、座瘡、

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囊胞性線維症、および免疫障害(自己免疫疾患およびガンを含む)などのヒトおよび動物 における非常に様々な疾患の治療のための使用が、この分野では知られている。 [0004]オキアミ油および/または海洋生物油は、自己免疫性マウス狼瘡および他の自己免疫疾患 の治療に使用できること、また心臓血管疾患の治療に使用できることが、この分野では知 られている。 [0005]しかし、これらの治療に使用されるオキアミ油および/または海洋生物油は、オキアミお よび/または海洋生物自身が持つ数ある有効成分の中のほんのひとつにしか過ぎないω-3 脂肪酸を有効成分として含有しているだけである。そのため、これらの疾患に対する治療 10 薬としてのオキアミ油および/または海洋生物油の潜在能力は弱められている。 [0006]天然資源を由来とする産生物を使用した治療に対する要望が大きくなっている。それゆえ 、疾患の予防および/または治療および/または疾病管理に対してより効果の高いオキア ミ抽出物および/または海洋生物抽出物を提供することが非常に望ましい。 【発明の開示】 【発明が解決しようとする課題】 [0007]本発明により、いくつかの疾患を予防および/または療治および/または治療する方法で 、療治効果的な量のオキアミ油および/または海洋生物油を患者に投与することを含む方 20 法が提供される。 【課題を解決するための手段】 [0008]本発明の好ましい実施形態において、オキアミ油および/または海洋生物油は、、以下の 工程: a)オキアミおよび/または海洋生物の材料をケトン溶媒(好ましくは、アセトン)に入れ て、前記海洋動物材料および/または水生動物材料からの可溶性脂質画分を抽出し; b) これを液体内容物と固体内容物に分離し; c)分離した液体内容物に存在する溶媒を蒸発させることによって液体内容物から第1の脂 質富含有画分を回収し; 30 d)前記固体内容物を、アルコール(好ましくは、エタノール、イソプロパノールまたはt-ブタノール)および酢酸エステル(好ましくは、酢酸エチル)からなる溶媒群から選択さ れる有機溶媒に入れて、前記海洋動物材料および/または水生動物材料からの残留可溶性 脂質画分を抽出し; e) これを液体内容物と固体内容物に分離し; f)分離した液体内容物から溶媒を蒸発させることによって第2の脂質富含有画分を回収 し;および g) 固体内容物を回収する工程 からなるプロセスから得られる。 [0009]40 本発明の好ましい実施形態において、オキアミ油および/または海洋生物油は、エイコサ ペンタエン酸、ドコサヘキサンエン酸、ホスファチジルコリン、ホスファチジルイノシト ール、ホスファチジルセリン、ホスファチジルエタノールアミン、スフィンゴミエリン、 α-トコフェロール、全トランスレチノール、アスタキサンチンおよびフラボノイドを含 む。 [0010]本発明の別の実施形態において、オキアミ油および/または海洋生物油は、エイコサペン タエン酸、ドコサヘキサンエン酸、リノレイン酸、α-リノレイン酸、リノール酸、アラ キドン酸、オレイン酸、パルミチン酸、パルミトレイン酸、ステアリン酸、ネルボン酸、 ホスファチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスファ 50

(11)

チジルエタノールアミン、スフィンゴミエリン、コレステロール、トリグリセリド、モノ グリセリド、αートコフェロール、全トランスレチノール、アスタキサンチン、カンタキ サンチン、βーカロテン、フラボノイド、亜鉛、セレン、ナトリウム、カリウムおよびカ ルシウムを含む。

(12)

[0011]

本発明のさらに別の実施形態において、オキアミ油および/または海洋生物油は、エイコ サペンタエン酸、ドコサヘキサンエン酸、リノレイン酸、α-リノレイン酸、リノール酸 、アラキドン酸、オレイン酸、パルミチン酸、パルミトレイン酸、ステアリン酸、ホスフ ァチジルコリン、ホスファチジルイノシトール、ホスファチジルセリン、ホスファチジル エタノールアミン、スフィンゴミエリン、コレステロール、トリグリセリド、モノグリセ リド、α-トコフェロール、全トランスレチノール、アスタキサンチン、カンタキサンチ ン、β-カロテン、亜鉛およびセレンを含む。

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本発明の方法によって治療および/または予防され得る疾患には、心臓血管疾患、関節炎、皮膚ガン、糖尿病、月経前症候群および経皮輸送増強がある。

[0013]

[0012]

本発明により、前記に記載された疾患を治療および/または予防および/または療治する ための組成物で、療治効果的な量のオキアミ油および/または海洋生物油を薬学的に受容 可能なキャリアとともに含む組成物もまた提供される。

[0014]

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本発明により、前記に記載された疾患を治療および/または予防および/または療治する ための、オキアミ油および/または海洋生物油の使用がさらに提供される。

[0015]

本発明により、前記に記載された疾患を治療および/または予防および/または療治する ための医薬品を製造するための、オキアミ油および/または海洋生物油の使用もまた提供 される。

【発明を実施するための最良の形態】

[0016]

本発明により、いくつかの疾患を防止および/または治療および/または療治するための 、オキアミおよび/または海洋生物の抽出物が提供される。 30

[0017]

酵素を含まない多用途な療治用の抽出油は、例えば、南極海(ナンキョクオキアミ(eupha sia superba))、太平洋(ツノナシオキアミ(euphasia pacifica))、大西洋、インド洋 (特に、モーリシャス島および/またはマダガスカルのレユニオン島の沿岸領域)、カナ ダ西海岸、日本沿岸、セントローレンス湾およびファンディ湾など、世界中のどこにでも 海洋環境にも生息するオキアミおよび/または海洋生物に由来する。この抽出油は遊離脂 肪酸脂質画分である。

[0018]

抽出プロセスは下記のように記載することができる:

(a)海洋性および/または水生のオキアミおよび/または海洋生物をケトン溶媒(好まし 40 くは、アセトン)に入れて、オキアミおよび/または海洋生物からの油脂を抽出する; (b)液相および固相を分離すること;

(c)液相に存在する溶媒を蒸発させることによって、工程(b)で得られた液相から脂質富含 有画分を回収する;

(d) 固相を有機溶媒に入れる。その際、有機溶媒としてはアルコール(好ましくは、エタ ノール、イソプロパノールまたはt-ブタノール)または酢酸エステル(好ましくは、酢酸 エチル)が使用できる。なお、これは、残留する可溶性脂質画分を固相から抽出するため に行われる。;

(e)液相および固相を分離すること;および

(f)液相に存在する溶媒を蒸発させることによって、工程(e)で得られた液相から脂質富含 50

有画分を回収すること。 [0019]酵素を含まないオキアミおよび/または海洋生物の抽出油の有効成分は下記の通りである 〔脂質〕 i)ω-3: i. エイコサペンタエン酸:>8g/100g ii. ドコサヘキサンエン酸:>2g/100g iii. リノレイン酸:>0.10g/100g iv. $\alpha - \mathcal{I} / \mathcal{I} / \mathcal{I}$ 酸: > 0.3g/100g 10 本発明においてω-3は30g/100gよりも多いことが好ましい実施形態である。 ii) $\omega - 6$: i. リノール酸: > 0.9g/100gii. アラキドン酸: <0.45g/100g、好ましくは<0.6g/100g iii) ω.−9 : i. オレイン酸: > 5g/100giv)パルミチン酸:>10g/100g v)パルミトレイン酸: 0.08g/100g vi)ステアリン酸: >0.5g/100g 〔リン脂質〕 20 ホスファチジルコリン:>4.5g/100g ホスファチジルイノシトール:>107mg/100g ホスファチジルセリン:>75mg/100g ホスファチジルエタノールアミン:>0.5g/100g スフィンゴミエリン:>107mg/100g 〔中性脂質〕 コレステロール: < 3g/100gトリグリセリド: < 55g/100g モノグリセリド:>0.5g/100g 本発明の別の実施形態において、オキアミ抽出物および/または海洋生物抽出物の中性脂 30 質は以下を含む: ジグリセリド:>0.5g/100g 〔抗酸化物質〕 $\alpha - h \exists \forall x \Box - \mu$ ($\forall \varphi \equiv \nu E$) : >1.01U/100g 全トランスレチノール (ビタミンA) : > 1500 IU/100g 〔色素〕 アスタキサンチン:>20mg/100g カンタキサンチン:>2mg/100g 〔金属〕 40 亜鉛:>0.1 mg/100 gセレン: > 0.1mg/100g 本発明の別の実施形態において、オキアミ抽出物および/または海洋生物の抽出物にはま た、下記が含まれる: フラボノイド:>0.5mg/100g ナトリウム:<500mg/100g カルシウム:>0.1mg/100g カリウム: > 50 mg/100 g $P N \ge - \phi L \ge < 8.5 mg/100 g$ タンパク質:>4g/100g 50

(13)

水分および揮発性成分: <0.8%。 [0020]オキアミ油抽出物および/または海洋生物抽出油の特性、抽出物が、抽出プロセスに由来 する溶媒残渣を25ppm未満で含有するものした。 該抽出油は下記の安定度指数を有している: 過酸化物価: < 0.1(mEq/kg) 油安定度指数: <0.1 (97.8℃で50時間後の数値) けん化指数:70~180 ヨウ素価: 60~130% 本発明をよりわかりやすく説明するために、以下に本発明の実施例を示して本発明に詳細 10 な説明するが、これは本発明の範囲を限定するものではない。 【実施例1】 [0021]心臓血管疾患の防止および/または治療 オキアミ抽出油および/または海洋生物抽出油は、コレステロールを生体内で低下させる ことが示されている。この抽出油はまた、血小板の接着およびプラークの形成を阻害し、 患者における血管内皮炎症を軽減させる。この抽出油は高血圧を予防できる。この抽出油 は低密度リポタンパク質の酸化を妨げる。この抽出油は、アポB-100の増大した細胞内分 解によってVLDLの分泌に対する阻害作用を有し得る。この抽出油はまた、CIIIアポリポタ ンパク質BおよびCIII非アポリポタンパク質Bリポタンパク質を減少させ、かつアンチトロ 20 ンビンIIIレベルを増大させる作用を示すことから、心筋梗塞後症候群を予防できる。オ キアミ抽出油および/または海洋生物抽出油は、冠状動脈疾患、高脂血症、高血圧、虚血 性疾患(すなわち狭心症、心筋梗塞、脳虚血、虚血の医学的や分析学的な証拠を伴わない 発作、不整脈)に関連するヒトでの心臓血管疾患に対する予防的使用に好適である。 [0022]動脈硬化性冠状動脈疾患および高脂血症の経過に対するオキアミ油および/または海洋生 物油の効果を評価するために、高脂血症が知られている患者で試験を行った(前向き臨床 試験、統計学的有意性p<0.05)。 [0023]

(14)

13名の患者グループにオキアミ油および/または海洋生物油の高濃度カプセル剤を投与し 30 た。魚油と、オキアミ油および/または海洋生物油とはともに、等しい量のω-3脂肪酸を 含有していた。推奨される投薬量は、800mgの油をそれぞれが含有するカプセルで1日あた り1錠~6錠である。この試験では、それぞれの患者に1日あたり6錠のカプセルを投与した

[0024]

該患者より、投薬前および2ヶ月の投薬後に、LDL、HDL、トリグリセリド、バイタルサイン(血圧や心拍数)、CBC、SGOT/SGPT、 y - GT、ALP、尿素、クレアチン、グルコース、K⁺、Na⁺、Ca²⁺および総間接的ビリルビンコレステロールのデータを採取した。

【0025】

表1には、前記の試験結果が示されている:

[0026]

【表1】

ベアードサンプル検定

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Ц |
| コレステロール | . 4854 | . 55800 | . 15476 | . 1582 | . 6320 | 3. 201 | 12 | . 008 |
| トリグリセリド | . 3538 | . 54543 | . 15127 | . 0242 | . 6834 | 2. 339 | 12 | . 037 |
| HDL | 2108 | —. 28859 | . 08281 | —. 3912 | 0303 | -2. 545 | 12 | . 026 |
| ГDГ | . 2846 | . 47333 | . 13128 | 0014 | . 5708 | 2. 168 | 12 | . 051 |
| Ch∘I∕HD∟ | . 3000 | . 53446 | . 14523 | . 0370 | . 5000 | 2. 420 | 12 | . 032 |
| | | | | | | | | |

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以上より、1g~4.8gのオキアミ抽出物を毎日の摂取することにより、被験患者に対して15%の範囲でコレステロールが低下し、15%の範囲でトリグリセリドが低下し、8%の範囲でHDLが増大し、13%の範囲でLDLが低下し、またコレステロール/HDL比が14%低下するという効果が認められた。

[0027]

このことは、オキアミ抽出物の摂取が、アテローム性動脈硬化の主要な原因因子であるこ 50

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とが知られている高脂血症に苦しんでいる患者に対して優れた効果を有することを示して いる。

(16)

【実施例2】

[0028]

関節炎の治療

オキアミ油および/または海洋生物油は、ヒト患者におけるインターロイキン-8およびイ ンターロイキン-1の産生を低下させることで、痛みを発する関節の数および日々使用する 鎮痛剤量を少なくするという臨床的な改善効果をもたらすことより、成人の関節炎、ステ ィル病、多関節性または少数関節性の若年性関節リウマチ、慢性関節リウマチ、変形性関 節症に伴う関節炎の症状を緩和する。なお、出血傾向のある患者または重度の精神病患者 はこの調査では除外した。

[0029]

クラスI、IIまたはIIIの進行である変形性関節症と診断され治療を受けており、試験前少なくとも3ヶ月間は非ステロイド性抗炎症剤(NSAIDs)および/または鎮痛剤の投与を受けている患者に関して、変形性関節症の臨床的経過に対するオキアミ油および/または海洋生物油の投与効果を評価するために、試験を行った(前向き臨床試験、統計学的有意性p<0.05)。

[0030]

13名の患者グループに、オキアミ油および/または海洋生物油の高濃度カプセルを、オキアミ油がカプセルあたり800mgであるカプセルの1日あたり6錠の割合で投与した。推奨さ 20れる投薬量は、純度の高いオキアミ抽出物で1日あたり1グラム~4.8グラムの範囲内である。被験患者には20%の脂肪(動物脂肪は10%未満)、40%のタンパク質、および40%の炭水化物からなる通常の健康食療法に従うことを依頼した。

[0031]

本試験において被験者の要件は、年齢が50歳~65歳の間であること(性別は男女どちらで もよい)、試験6ヶ月前~12ヶ月前に痛みとこりがあって原発性の変形性関節症の臨床的 診断(軽度~中程度)を受けた患者で、試験に先立ちX線撮影で病変が確認されているこ とである。あわせて、少なくとも試験前の3ヶ月間にわたって、アセトアミノフェン、抗 炎症剤、もしくはオピオイド鎮痛剤の使用を必要とする変形性関節症(0A)のある程度の症 状が認められていることも上記要件に含まれる。患者には、ウォッシュアウト目的のため 30 に試験開始前の1週間はあらゆる鎮痛薬の使用を止めるように依頼した。

[0032]

被験者より除外する基準は、重度の変形性関節症である場合、NSAIDs、アスピリン、もし くは他の抗炎症剤を継続的に使用し止めることが出来ない場合、無作為訪問で4週間以内 に局所的鎮痛剤の使用したと認められる場合、過去3ヶ月以内に膝へのステロイド注射を 行っている場合、3ヶ月以内に理学療法または筋肉調整を開始した場合、海産食物アレル ギー、抗凝血剤またはサリチラートを使用している場合、1日あたりカクテル3杯を越える アルコールを摂取している場合、痛みの評価に影響をきたすような医学的疾患や関節炎疾 患を併発している場合、どちらかの膝が手術前(関節鏡検査を含む)である場合、公知の 「二次性」変形性関節症の要因が認められる場合である。

[0033]

評価は、NSAIDsおよび/または鎮痛剤および/またはSAARDsの1日量、痛みを有する関節の数、腫れた関節の数、朝のこわばりの持続時間、目視によるアナログスケール(0~100)でのWOMACスケール、ならびにSF36に基づいた。予備的な結果が2ヶ月後に得られた。日常活動のために要求されるNSAIDおよび/または鎮痛剤および/またはSAARDの数が開始時および開始後2ヶ月で記録されている。

[0034]

表2に示した結果より、関節炎の緩和に対するオキアミ抽出物の摂取の効果が明らかにと なった。

[0035]

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【表2】

| | | 頻度 | % | 有効% | 累積% |
|-------|------------|----|--------|--------|-------|
| 変化 | なし | 3 | 23. 1 | 23. 1 | 23. 1 |
| 痛みの緩和 | | 10 | 76. 9 | 76. 9 | 100.0 |
| 合 | ā † | 13 | 100. 0 | 100. 0 | |

これは、13名の内の10名(76.9%)が痛みの著しい緩和および大きい関節(下部背骨、膝、肩)の柔軟性の著しい改善を報告したことを示している。

(17)

【実 施 例 3 】

【0036】

皮膚ガンの予防

オキアミ油および/または海洋生物油は、そのレチノールの抗ガン性作用、アスタキサン チンの抗ガン性作用、およびそのリン脂質の抗ガン性作用のために皮膚ガンの予防剤であ ることが示されている。

[0037]

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UVBにより誘導される皮膚ガンに対するオキアミ油および/または海洋生物油の潜在的な 光保護能力を評価するために、研究を、皮膚ガンに対するその感受性が証明されているの で、ヌードマウス、好ましくは、C57BL6ヌード類遺伝子系マウスであるB6NU-T(ヘテロ接 合体)に対して行った。

【0038】

被験マウスを下記のグループとした:魚油を投与するグループ(全48匹)、うち経口補給 (po)による投与(16匹)、局所適用による投与(16匹)、poおよび局所適用による投与 (16匹);オキアミ油および/または海洋生物油を投与するグループ(全48匹)、;poに よる投与(16匹)、局所適用による投与(16匹)、poおよび局所適用による投与(16匹) 。皮膚ガンの防止に対するオキアミ油および/または海洋生物油の効力を明らかにするた めに、無作為化盲検対照方式で試験を行った(統計学的有意性p<0.05)。半数のマウス は100重量%のオキアミ油および/または海洋生物油を含有する油を経口的にまたは局所 的にまたはそれらの両方の方法で投与し、残る半数についても魚油で同様に投与を行った

。 【0039】

栄養摂取は、最初の1週間については無脂肪餌であり、その後の2週間~20週間については、1日あたり1mlの油の量で下記に記載するグループに従って変更された。

[0040]

マウスは下記のように6群に分けられた:

グループA:魚油の補給(総カロリーの20%)を伴う無脂肪餌

グループB: 無脂肪餌(100%のカロリー)+1日に2回の魚油の局所適用

グループC: 魚油の補給(総カロリーの20%)を伴う無脂肪餌+1日に2回のダイズ油の局 所適用

グループD:オキアミ油および/または海洋生物油の補給(総カロリーの20%)を伴う無 脂肪餌

グループE:無脂肪餌(100%のカロリー)+1日に2回のオキアミ油および/または海洋生物油の局所適用

グループF:オキアミ油および/または海洋生物油の補給(総カロリーの20%)を伴う無 脂肪餌+1日に2回のオキアミ油および/または海洋生物油の局所適用。 【0041】 マウスは、第2週~第20週の期間中、蛍光試験ランプ(放射スペクトル:270~400nm)を 使用してUVB線に暴露した。該試験ランプをマウスから30cmの距離に配置し、1日あたり30 分間UVBに暴露して試験を行った。試験開始20週間終了後、または悪性の腫瘍が生じた時 点で、被験マウスをエーテルで麻酔し屠殺した。皮膚のガン発生の徴候を病理学者によっ て盲検的に調べられた。

[0042]

下記の表(表3~表8)には、マウスの皮膚に紫外線照射試験を5週間の行った時のガンの発生について得られた結果が示されている。

【0043】

【表3】

オキアミ抽出物の経口摂取

| | | | | 頻度 | 割合(%) | 有効% | 累積% |
|---|---|---|---|----|-------|--------|--------|
| 有 | 効 | 良 | 性 | 14 | 87. 5 | 87. 5 | 87. 5 |
| | | ガ | ン | 2 | 12. 5 | 12. 5 | 100. 0 |
| | | 合 | 計 | 16 | 100.0 | 100. 0 | |

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[0044]

【表4】

コントロールの経口摂取

| | | | 頻度 | 割合(%) | 有効% | 累積% |
|---|---|----|----|-------|--------|-------|
| 有 | 効 | 良性 | 14 | 87.5 | 87. 5 | 87. 5 |
| | | ガン | 2 | 12. 5 | 12. 5 | 100.0 |
| | | 合計 | 16 | 100.0 | 100. 0 | |

【0045】

【表5】

オキアミ抽出物の局所的摂取

| | | 頻度 | 割合(%) | 有効% | 累積% |
|----|----|----|--------|--------|--------|
| 有効 | 良性 | 16 | 100. 0 | 100. 0 | 100. 0 |

【0046】 【表6】

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| | | | 頻度 | 割合(%) | 有効% | 累積% |
|---|---|----|----|-------|--------|--------|
| 有 | 効 | 良性 | 5 | 31.3 | · 31.3 | 31.3 |
| | | ガン | 11 | 85. 8 | 85. 8 | 100. 0 |
| | | 合計 | 16 | 100.0 | 100. 0 | |

コントロールの局所的摂取

(19)

[0047]

【表7】

オキアミ抽出物の経口摂取および局所的摂取

| | 頻度 | 割合(%) | 有効% | 累積% |
|------|----|-------|--------|--------|
| 有效良性 | 16 | 100.0 | 100. 0 | 100. 0 |

[0048]

【表8】

コントロールの経口摂取および局所的摂取

| | | | | 頻度 | 割合(%) | 有効% | 累積% |
|---|---|---|---|----|-------|--------|--------|
| 有 | 効 | 良 | 性 | 10 | 62. 5 | 62. 5 | 62. 5 |
| | | ガ | ン | 5 | 37. 5 | 37.5 | 100. 0 |
| | | 合 | 計 | 16 | 100.0 | 100. 0 | |

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これらの結果より、オキアミ油の経口使用および局所的使用はともに、皮膚ガンを誘導するUVBの有害な作用から皮膚を保護する効果的があることが示された。

【実施例4】

[0049]

塗布療治薬における経皮輸送

オキアミ油および/または海洋生物油は、皮膚科学的な局所的に用いる塗布療治薬に対す る基質として経皮輸送を増強する。オキアミ油および/または海洋生物油は、クリーム、 軟膏、ゲル、ローションおよびオイルによって皮膚科学的処置において使用することがで きる。オキアミ油および/または海洋生物油はまた、麻酔剤、コルチコステロイド、抗炎 症剤、抗生物質およびケトン分解機能に関連することなどの様々な治療適用において使用 することができる。

[0050]

オキアミ油および/または海洋生物油を局所的治療のための基質として用いた場合の効力 、またオキアミ油および/または海洋生物油を単独であるいは基質として用いた場合の経 50

皮吸収速度を評価するために、C57BL6ヌード類遺伝子系マウスであるB6NU-T(ヘテロ接合体)に対する無作為化盲検対照方式で試験を行った。

[0 0 5 1 **]**

表5および表6に表される結果は、オキアミ油による局所的治療により、真皮を介したレ チノールおよび他の抗酸化物質の吸収が促進され、これにより強力な光保護能力がもたら され、その結果、UVBによって誘導される皮膚ガンからの100%保護されたのである。対照 的に、魚油を全トランスレチノールとともに塗布した場合は、ガン発生率が68.8%であっ た。

【実施例5】

[0052]

皮膚に局所的に塗布する化粧品の経皮輸送を

オキアミ油および/または海洋生物油は、クリーム、軟膏、ゲル、ローションまたはオイ ルによる皮膚の水和、しわ防止、角質溶解剤、剥皮および美顔用パックに関連する皮膚科 学的な局所的化粧適用のための基質としての経皮送達を増強するために使用することがで きる。

[0053]

老化によるしわ、および顔面のしわにおけるオキアミ油および/または海洋生物油の効果 を評価するために、顔の乾燥肌およびしわに悩む被験者に対する前向き臨床試験により研 究を行った。これら被験者は、他の皮膚科学的もしくは非皮膚科学的な病気によって重度 に限定された予後診断を受けた者ではなく、また出血傾向のある患者および重度の精神病 患者はこの調査では除外した。顔の乾燥肌またはしわを有する13名の健康な白人女性がこ の研究には含まれている。被験女性は、800mgのオキアミ油を含有するカプセルを、1日あ たり6カプセル摂取することが依頼されている。推奨される1日投薬量は約1g~4.8gのオ キアミ抽出物である。

[0054]

表9には、前記に記載された方法に従って皮膚の水和について得られた結果が示されている。

【 O O 5 5】

【表9】

皮膚の保湿効果の変化

| | 頻度 | % | 有効% | 累積% |
|--------|----|--------|-------|--------|
| 変化なし | 4 | 30. 8 | 30.8 | 30. 8 |
| 保湿効果あり | 9 | 69. 2 | 69.2 | 100. 0 |
| 合計 | 13 | 100. 0 | 100.0 | |

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2ヶ月にわたる試験的研究の結果では、13名の内の9名(69.2%)がヒト被験者における皮 膚(顔、手および腕)の保湿効果の上昇、肌のきめと弾力性の著しい改善が見られた。 【0056】

さらに、これらの結果はまた、オキアミ抽出物がしわを取る方法として有用であることを 示している。オキアミ油に含まれる全トランスレチノールのしわ取り剤としての仕組みは 次のとおりである:

再生作用および特有な抗炎症作用

・血液循環の改善

・細胞分裂および代謝回転の速度を増大させて表皮再生を活性化させること

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・ケラチンの分化を促進させること

・コラーゲンを再生すること

・新陳代謝される皮膚の最表層の細胞について、処置を行わなかった日光による損傷細胞より正常に成熟できること

(21)

・皮膚を構造的に支持するタンパク質コラーゲンおよびエラスチンを分解する酵素の活性 化を低減させること。

[0057]

患者の皮膚に施されたオキアミ抽出物を用いて得られた結果は、オキアミ抽出物が、水和 を増大させること、および上記に記載される仕組みによるしわ取り効果を有していること を示している。

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【実施例6】 【0058】

月経前症候群

表10には、女性における月経前症候群に関連する痛みおよび気分変化を軽減させるためのオキアミ油の使用から得られた結果が示される。オキアミ油の抽出物が、2ヶ月間にわたり7名の女性に投与した。被験女性には、800mgのオキアミ油を含有するカプセルを、1日あたり6カプセルを投与した。推奨される1日投薬量は約1g~4.8gのオキアミ抽出物である。被験者はすべて、自身の普段の食習慣を続けること、また食事について何らかの制限を開始しないように指示された。重大な副作用は何ら報告されなかった。

【0059】

被験者の女性はすべて、顕著な情緒的および/または身体的な不快を月経の7日前~10日 前に訴えている人である。月経前症候群の評価は、既に確立している0(症状なし)~10 (耐えられない)で表される自己評価の視覚的アナログ尺度を用いて行い、これによって 月経前の不快症状に対するオキアミ抽出物の効果を評価する予備的なデータとして使用した。

[0060]

2ヶ月間の投薬治療をおこなった本研究の被験女性の60%にあたるデータ分析を行った。 大多数の女性(73.3%)は、月経前の情緒的苦悩および身体的苦悩の両方について、臨床 的に著しい軽減が認められた(表10参照)。

【0061】

【表10】

| PMS症状 | 頻度 | 有効% | 累積% |
|-------|--------|--------|--------|
| 変化なし | 26. 7 | 26. 7 | 26. 7 |
| 前向き | 73. 3 | 73. 3 | 100. 0 |
| 合計 | 100. 0 | 100. 0 | |

月経前症候群の症候学に対するオキアミ抽出物の効果の頻度分布

【実施例7】

[0062]

糖尿病

ヒト患者8名に対し、800mgのオキアミ油を含有するカプセルを、1日あたり6カプセル、2 ケ月間に渡り投与した。推奨される1日投薬量は約1g~4.8gのオキアミ抽出物である。 表11には、2ヶ月後の患者について試験されたグルコースの変化が示されている。 【0063】

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【表11】

患者におけるグルコースの変化

| ベ ア ー ド 差 | | | | | | | |
|-----------|--------|---------|-------------|----------------|--------|-----|-------|
| 試験パラメーター | 平均值 | SD. | 標準誤差
平 均 | 差の95%信頼性区間 | t 值 | d f | 自由度 |
| グルコース | . 5778 | . 60369 | . 20123 | . 1137—1. 0418 | 2. 871 | 8 | . 021 |

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血中グルコースの20%の低下が、オキアミ抽出物を摂取した患者について認められた。こ のことは、オキアミ抽出物の摂取が血中グルコース含有量を抑制し、従って、ヒト患者に おいて糖尿病を抑制することを示している。

【0064】

本発明をその具体的な実施形態に結び付けて記載したが、さらなる改変が可能であり、付記される請求項の範囲に従うような、そして本明細書に記載した本質的特徴に当てはめる ことが可能であるような、そして本発明に関連する技術分野における公知のあるいは慣例 的な方法の範囲内にあるような、本発明が開示する内容からの逸脱を含み、また本発明の 原理に一般的に従うような本発明のあらゆる適応、利用もしくは改変を含むものとして、 この出願は意図されるものであると理解されるであろう。

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| EAS | Thies KRILL AND/OR MARINE EXTRACTS FOR PRIP
US, ANTHRITIN, SKIN CANCER, DIABITTS, PRIMIP
ADD/REI: The present invention rolates to a method of treat | ISTRUAL | SYNDROME AND TRANSDERMAL TRANSPORT |

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PCT/CA02/09843

Krill and/or marine extracts for prevention and/or treatment of cardiovascular diseases, arthritis, skin cancer, diabetes, premenstrual syndrome and transdermal transport.

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BACKGROUND OF THE INVENTION

Field of the Invention

This invention relates to multi-therapeutic extracts derived from krill and/or marine, which can prevent and/or treat several diseases.

Description of Prior Art

Krill is the common name for small, shrimp-like crustaceans, however not shrimp, that swarm in dense shoals, especially in Antarctic waters. It is one of the most important food source for fish, some kind of birds and especially for baleen whales as being an important source of protein. Krill is also a good source of omega-3 fatty acid, which are well known for their health benefits.

It is known in the art to use krill and/or marine enzymes for the treatment of a great variety of diseases in human and animals such as infections. Inflammations, cancers, HIV/AIDS, pain, polyps, warts, hemorrholds, plaque, wrinkles, thin hairs, allergic tich, anti-adhesion, eye disease, acne, cystic fibrosis and immune disorders including autoimmune disease and cancer.

It is also known in the art that krill and/or marine oil may be used for the treatment of autoimmune murine lupus and other autoimmune diseases and can also be used for treating cardiovascular diseases.

However, the krill and/or marine oil used for these treatments has only conserved its ornega-3 fatty acids as active ingredients, which is a very small part of all the active ingredients of the krill and/or marine itself. This fact reduces the potential of the krill and/or marine oil as a treatment for these diseases.

There is an increasing demand for treatments using products derived from a natural source, therefore, it would be highly desirable to be provided with a krill and/or marine extract having an enhanced potential for prevention and/or treatment and/or management of disease.

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SUMMARY OF THE INVENTION

In accordance with the present invention there is provided a method of prevention, therapy and/or treatment of several disease, the method comprising the administration of a therapeutically effective amount of knil and/or marine oil to a patient.

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In a preferred embodiment of the present invention the krill and/or marine oll is obtained from a process comprising the steps of:

(a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from the marine and/or aquatic animal material;

(b) separating the liquid and solid contents;

(c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

(d) placing the solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from the marine and/or aquatic material;

(e) separating the liquid and solid contents;

(f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

(g) recovering the solid contents.

In a preferred embodiment of the present invention, the krill and/or marine oil comprises Eicosapentanoic acid, Docosahexanoic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolaimine, Sphingomyelin, a-tocopherol, all-trans retinol, Astaxanthin and flavonoid.

In another embodiment of the present invention, the krill and/or marine oil comprises Elcosapentanoic acid, Docosahexanoic acid, Linolenic acid, Alpha-linolenic acid, Linoleic acid, Arachidonic acid, Oleic

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acid, palmitic acid, palmitoleic acid, stearic acid, nervonic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, Cholesterol, Triglycerides, Monoglycerides, a-tocopherol, all-trans retinol, Astaxanthin, Canthaxanthin, β-carotene, flavonoid, Zinc, Selenium, sodium, potassium and calcium.

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In another embodiment of the present invention, the krill and/or marine oil comprises Eicosapentanoic acid, Docosahexanoic acid, Linolenic acid, Alpha-Ilnolenic acid, Linoleic acid, Arachidonic acid, Oleic acid, palmitic acid, palmitoleic acid, stearic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, Cholesterol, Triglycerides, Monoglycerides, a-tocopherol, all-trans retinol, Astaxanthin, Canthaxanthin, β-carotene, Zinc and Selenium.

The diseases that can be treated and/or prevented by the method of the present invention are cardiovascular diseases, arthritis, skin cancer, diabetes, premenstrual syndrome and transdermal transport enhancement.

In accordance with the present invention there is also provided a composition for the treatment and/or prevention and/or therapy of the previously mentioned diseases, the composition comprising a therapeutically effective amount of kill and/or marine oil in association with a pharmaceutically acceptable carrier.

In accordance with the present invention, it is further provided the use of krill and/or marine oil for the treatment and/or prevention and/or therapy of the previously mentioned diseases.

In accordance with the present invention, it is also provided the use of knill and/or marine oil for the manufacture of a medicament for the treatment and/or prevention and/or therapy of the previously mentioned diseases.

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DETAILED DESCRIPTION OF THE INVENTION

In accordance with the present invention, there is provided krill and/or marine extract for prevention and/or treatment and/or therapy of several diseases.

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A multi-therapeutic oil extract tree of enzyme is derived from krill and/or marine, found in any marine environment around the world, for example, the Antarctic ocean (euphasia superba), the Pacific ocean (euphasia pacifica), the Atlantic ocean, the Indian ocean, in particular coastal regions of Mauritius Island and/or Reunion Island of Madagascar, Canadian West Coast, Japanese Coast, St-Lawrence Gulf and Fundy Bay, and this oil extract is a free fatty acid lipid fraction.

The extraction process can be described as the following:

(a) Placing marine and/or aquatic krill and/or marine in a ketone solvent, preferably acetone, to achieve the extraction of grease from the krill and/or marine;

(b) Separating the liquid and the solid phases;

(c) Recovering a lipid rich fraction from the liquid phase obtained at step (b) by evaporation of the solvent present in the liquid phase;

(d) Placing the solid phase in an organic solvent, which can be alcohol, preferably ethanol, isopropanol or t-butanol, or esters of acetic acid, preferably ethyl acetate. This in order to extract the remaining soluble lipid fraction from the solid phase;

(e) Separating the liquid and the solid phases; and

(f) Recovering a lipid rich fraction from the liquid phase obtained at step (e) by evaporation of the solvent present in the liquid phase.

The active components of the enzyme-free krill and/or marine oil extract are:

lipids

i) Omega-3:

- i, Eicosapentanoic acld: >8g/100g
- ii. Docosahexanoic acid: >2g/100g

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ili, Linolenic acid: >0.10g/100g

iv. Alpha-linolenic acid; >0.3g/100g

In the preferred embodiment of the present invention, the Omega-3 are

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found in more than 30g/100g.

ii) Omega-6: i, Linoleic acid: >0.9g/100g

ii. Arachidonic acid: <0.45g/100g, preferably < 0.6g/100g

iii) Omega-9: i. Oleic acid: >5g/100g

iv) palmitic acid: >10g/100g

v) palmitoleic acid; 0.08g/100g

vi) stearic acid: > 0.5g/100g

Phospholipids

Phosphalidylcholine: >4.5g/100g

Phosphatidylinositol: >107mg/100g

Phosphatidylserine: >75 mg/100g

Phosphatidylethanolamine: >0.5g/100g

Sphingomyelin: >107mg/100g

Neutral lipids

Cholesterol: <3g/100g

Triglycerides: <55g/100g

Monoglycerides: >0.5g/100g

in another embodiment of the present invention, the neutral lipids of the

krill and/or marine extract also comprises:

Diglycerides: >0.5g/100g

Antioxydants

 α -tocopherol (vitamin E): >1.0 IU/100g

all-trans retinol (vitamin A): >1500 IU/100g

β-carotene: > 3000 μg/100 ml

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Pigments

Astaxanthin: >20 mg/100g Canthaxanthin: > 2 mg/100g Metals Zinc: >0.1 mg/100g Selenium: >0.1 mg/100g In another embodiment of the present invention, the krill and/or marine extract also comprises: Flavonoids: >0.5mg/100g Sodium: < 500mg/100g Calcium; >0.1mg/100g Potassium; > 50mg/100g Aluminum: < 8.5mg/100g Protein: > 4g/100g Moisture and volatile matter. <0.8% After characterization of the krill and/or marine oil extract, it was determined that the extract contains less than 25 ppm of solvent residue from the extraction process. The oil has the following stability indexes: Peroxide value: < 0.1(mEq/kg) Oil Stability index: < 0.1 after 50 hours at 97.8°C Saponification index: 70-180 Iodine value:60-130% The present invention will be more readily understood by

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referring to the following examples which are given to illustrate the invention rather than to limit its scope.

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Example 1

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Cardiovascular disease prevention and/or treatment

Krill and/or marine oil has been shown to decrease cholesterol in vivo. It also inhibits platelet adhesion and plaque formation and reduces vascular endothelial inflemmation in a patient. It can offer hypertension prophylaxis. It prevents oxidation of low-density lipoprotein. It may have an inhibitory effect on the secretion of VLDL due to increased intracellular degradation of apo 8-100. It also offers a post-myocardial infarction prophylaxis because of Its ability to decrease CIII apolipoprotein B, to decrease CIII non-apolipoprotein B lipoproteins and to increase antithrombin III levels. Krill and/or marine oil Is suitable for prophylactic usage against cardiovascular disease in human where cardiovascular disease relates to coronary artery disease, hypertipidemia, hypertension, ischemic disease (relating to angina, myocardial infarction, carebral ischemia, shock without clinical or laboratory evidence of ischemia, armythmia)

To evaluate the effects of krill and/or marine oil on the course of arterioscierotic coronary artery disease and hyperlipidemia, a study was performed (prospective clinical trial, statistical significance p<0.05) with patients with known hyperlipidemia.

A group of 13 patients took krill and/or marine oil concentrate gelules. Both fish oil and krill and/or marine oil contained equal amounts of omega-3 fatty acids. Recommended dosage is of 1 to 6 capsules per day, each capsule containing 800 mg of oil. In this study, each patient took 6 capsules per day.

The patients were tested for LDL, HDL, Triglycerides, vital signs, CBC, SGOT/SGPT, γ -GT, ALP, Urea, Creatine, Glucose, K*, Na*, Ca²⁺ and total indirect bilinubin cholesterol before treatment and also at 2 months.

Table 1 is showing the results obtained from the previously described tests:

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Table 1 Paired Samples Test

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| Paired | Samples | 1621 |
|--------|---------|------|
| | | |

| Parameter
tested | | | 95% Confidence
Interval of the
Difference | | t-value | đi | Sig. (2-
tailed) | |
|---------------------|-------|--------|---|-------|---------|--------|---------------------|------|
| | | | | Lower | Upper | | | _ |
| Cholesterol | .4954 | .55800 | .15476 | .1582 | .8326 | 3.201 | 12 | 800. |
| Triglycericies | .3538 | .54543 | .15127 | .0242 | .6834 | 2.339 | 12 | .037 |
| HOL | 2108 | .29859 | .08281 | 3912 | - 0303 | -2.545 | 12 | .026 |
| LDL | .2846 | .47333 | .13128 | 0014 | .5706 | 2.168 | 12 | .051 |
| Chol / HDL | .3600 | .53446 | .14823 | .0370 | .6830 | 2.429 | 12 | .032 |

From the above, it was shown that a daily uptake of 1 to 4.8 g of krill extract was providing to the patients a cholesterol decrease in the range of 15%, a triglycerides decrease in the range of 15%, a HDL increase in the range of 8%, a LDL decrease in the range of 13% and a Cholesterol/HDL ratio decrease of 14%.

This shows that an uptake of krill extract has a beneficial effect on patient suffering from hyperlipidemia, which is known to be the primary causative factor of atherosclerosis.

Example 2 Arthritis treatment

Krill and/or marine oil offers symptomatic relief for Arthritis where arthritis relates to adult arthritis, Still's disease, polyarticular or psuciarticular juvenile rheumatoid arthritis, rheumatoid arthritis, osteoarthritis because it has been shown that it provides a clinical improvement in decreasing the number of tender joints and of analgesics consumed daily by decreasing the production of Interleukin-8 and Interleukin-1 in human patients. Patients with a bleeding tendency or severe psychiatric disease were excluded from the study.

To evaluate the effects of krill and/or marine oil supplementation on the clinical course of osteoarthritis, a study was performed (prospective clinical trial, statistical significance p<0.05) with patients diagnosed with and treated for osteoarthritis which is Active class I, II or III and having

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treatment with NSAIDs and/or analgesics for at least 3 months before enrollment.

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A group of 13 patients took krill and/or marine oil concentrate capsules at a daily rate of 6 capsules of 800mg kril oil per capsule. The recommended dosage varies between 1 and 4.8 grams of pure krill extract per day. Patients were asked to follow a nonnal healthy diet consisting of 20% fat (less than 10% animal fat), 40% protein and 40% carbohydrates.

The inclusion criteria for the study are being aged between 50 and 65 years, both genders being admissible, having a clinical diagnosis of primary osteoarthritis (mild to moderate) 6 to 12 months prior to study enrollment including pein and stiffness, radiographic confirmation of illness prior to enrollment. It also include evidence of measurable symptoms of OA for at least 3 months prior to study enrollment requiring the use of acetaminophen, anti-inflammatory agents or opioid analgesics. Patients were asked to stop the use of all "pain-killers" the week prior to initiation of the trial for wash-out purposes.

The Exclusion criteria were a severe osteoarthnitis, unavoidable sustained use of NSAID's, aspirin or other medicines for anti-inflammatory use, use of topical analgesics within 4 weeks of randomization visit, steroid injection into either knee within past 3 months, initiation of physical therapy or muscle conditioning within 3 months, initiation of physical therapy or muscle conditioning within 3 months, iseafood allergies, use of anticoagulants or salicylates, alcohol consumption exceeding 3 mixed drinks per day, concurrent medical/arthnitic disease that could confound or interfere with the evaluation of pain, prior surgery (including arthroscopy) of either knee, a known "secondary" cause of osteoarthnitis.

Evaluation was based on daily dose of NSAIDs and/or analgesics and/or SAARDs, number of painful joints, number or swollen joints, duration of moming stiffness, visual analog scale (0-100) WOMACscale and SF36. Preliminary results have been obtained after 2 months. The number of NSAIDs and/or analgesics and/or SAARDs required for daily functioning has been recorded at initiation and at 2 months after initiation.

Results shown at Table 2 demonstrate the effect of an uptake of krill extract on the relief of arthritis.

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| | Table | 2 | | |
|-------------|-----------|-------|---------|------------|
| | Frequency | % | Valld % | Cumulative |
| No change | 3 | 23.1 | 23.1 | 23.1 |
| Pain relief | 10 | 76.9 | 76.9 | 100.0 |
| Total | 13 | 100.0 | 100.0 | Γ |

This shows that ten out of 13 (76.9%) people reported a significant pain relief and improvement of flexibility of large joints (lower back, knees, shoulders)

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Example 3

Skin Cancer Prophylaxis

Krill and/or marine oil has been shown to be a skin cancer prophylactic because of its retinol anti-carcinogenic effect, Astaxanthin anti-carcinogenic effect and its phopholipid anti-carcinogenic effect.

To evaluate the photoprotective potential of krill and/or marine oil against UVB-induced skin cancer, a study was performed on nude mice, preferably on C57BL6 Nude Congenic Mice - B6NU-T (heterozygotes) because of their proven susceptibility to skin cancer.

Groups were formed as follows: 48 fish oil: 16 with oral supplementation (po) 16 with local application, 16 with po and local application, 48 kill and/or marine oil: 16 with po, 16 with local application, 16 with po and local application. In order to establish efficacy of kill and/or marine oil for the prevention of skin cancer, the test was conducted as a randomized blind controlled trial (statistical significance p<0.05). Half of the mice have been treated orally or topically or both with oil containing 100% by weight kill and/or marine oil and the other half have been treated the same way with fish oil.

Nutrition was fat-free chow for the first week and was modified accordingly with the assigned group as described below for the following 2-20 weeks in the quantity of 1 ml of oil per day.

The mice were divided in six groups as follows:

Group A: fat-free chow with supplementation of fish oil (20% of total calories)

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Group B: fat-free chow (100% of calories) + local application of fish oil 2 times per day

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Group C: fat free chow with supplementation of fish oil (20% of total calories) + local application of soy oil 2 times per day

Group D: fat-free chow with supplementation of krill and/or marine oil (20% of total calories)

Group E: fat free chow (100% of calories) + local application of krill and/or marine oil 2 times per day

Group F: fat-free chow with supplementation or krill and/or marine oil (20% of total calories) + local application of krill and/or marine oil 2 times per day

The mice had been submitted to UVB radiation using a fluorescent test lamp, emission spectrum 270-400 nm during weeks 2-20. The essay were performed during 30 minutes of UVB exposure per day and the test lamp was at a distance of 30 cm from the mice. At the end of the 20 weeks, or when malignant tumors had formed, mice were anesthetized with ether and sacrificed. Skin was examined blind by pathologists for signs of carcinogenesis.

The following tables (Tables 3-8) are showing the results obtained about the incidence of cancer when ultra-violet radiations are administered to mice's skin during 5 weeks.

Table 3

| Krill extract Oral uptake | | | | | | |
|---------------------------|--------|-----------|---------|---------------|--------------------|--|
| | | Frequency | Percent | Valid Percent | Cumulative Percent | |
| Valid | Benign | 14 | 87.5 | 87.5 | 87.5 | |
| | Cancer | 2 | 12.5 | 12.5 | 100.0 | |
| | Total | 16 | 100.0 | 100.0 | | |

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Table 4

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| | | Frequency | Percent | Valid Percent | Cumulative Percent |
|-------|--------|-----------|---------|---------------|--------------------|
| Valid | Benign | 14 | 87.5 | 87.5 | 87.5 |
| | Cancer | 2 | 12.5 | 12.5 | 100.0 |
| | Total | 16 | 100.0 | 100.0 | |

Table 5

| Krill extract topical uptake | | | | | | |
|------------------------------|--------|-----------|---------|---------------|--------------------|--|
| | | Frequency | Percent | Valid Percent | Cumulative Percent | |
| Valid | BENIGN | 16 | 100.0 | 100.0 | 100.0 | |

Table 6

| Control | topical | uptake |
|---------|---------|--------|
| | | |

| | | Frequency | Percent | Valid Percent | Cumulative Percent |
|-------|--------|-----------|---------|---------------|--------------------|
| Valid | BENIGN | 5 | 31.3 | 31.3 | 31.3 |
| | Cancer | 11 | 58.8 | 68.8 | 100.0 |
| | Total | 16 | 100.0 | 100.0 | |

Table 7 Krill extract topical and oral untake

| | _ | Frequency | Percent | Valid Percent | Cumulative Percent |
|-------|--------|-----------|---------|---------------|--------------------|
| Valid | BENIGN | 16 | 100.0 | 100.0 | 100.0 |

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Table 8

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Control topical and oral uptake

| | | Frequency | Percent | Valid Percent | Cumulative Percent |
|-------|--------|-----------|---------|---------------|--------------------|
| Valid | BENIGN | 10 | 62.5 | 62.5 | 62.5 |
| | Cancer | 6 | 37.5 | 37.5 | 100.0 |
| | Total | 16 | 100.0 | 100.0 | |

The results obtained shows that both oral and topical use of krill oil is effective for the protection of the skin against the harmful effects fo UVB radiation induced skin cancer.

Example 4

Transdermal transport in therapeutic applications

Krlli and/or marine oil enhances transdermal transportation as a substrate for dermatological topical therapeutic applications. It may be used in dermatological treatments via creams, orintments, gels, totions and oils. It may also be used in various therapeutic applications such as relating to anesthesic, corticosteroids, anti-inflammatory, antibiotic and ketolytic functions.

To evaluate the efficacy of krill and/or marine oil as a substrate for topical treatments and the speed of transdermal absorption of krill and/or marine alone or as a substrate, a study was performed as a randomized blind controlled trial on C57BL6 nude Congenic Mice - B6NU-T (heterozygotes).

The results appearing in tables 5 and 6 are showing that topical treatment with krill oil faciliate the absorption of retinol and other antioxydants through the dermis which in turn result in significant photoprotective potential which in turn results in 100% protection from UVB induced skin cancer. In contrast, fish oil application with all-trans retinol resulted in 68.8% incidence of cancer.

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Example 5

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Transdermal Transport for dermatological topical cosmetic applications

Krill and/or marine oil can be used to enhance transdermal transportation as a substrate for dermatological topical cosmetic applications where cosmetic applications relate to skin hydration, antiwrinkle, keratolytics, peeling and mask via creams, ointments, gels, lotions or oils.

To evaluate the effects of Krill and/or marine oil in aging and facial wrinkles, a study was conducted as a prospective clinical trial on patients concerned about facial dryness and wrinkles. Those patients had no prognosis severely limited by other dermatological or nondermatological condition, bleeding tendency or severe psychiatric disease.

13 Healthy caucasian women with facial dryness or winkles have been included in this study. Women have been asked to take 6 capsules a day, each capsule containing 800 mg of krill extract. The recommended daily dosage is of about 1 to 4.8 g of krill extract.

Table 9 shows results obtained on skin hydration following the method previously described.

| <u>Table 9</u>
Changes in skin hydration | | | | | | |
|---|-----------|-------|---------|-----------------|--|--|
| | Frequency | % | Valid % | Cumulative
% | | |
| No change | 4 | 30.8 | 30.8 | 30.8 | | |
| Hydration | 9 | 69.2 | 69.2 | 100.0 | | |
| Total | 13 | 100.0 | 100.0 | 1 | | |

The results of the pilot study after 2 months indicate that nine out of 13 (69.2%) people reported a significant improvement of the hydration, texture and elasticity of the skin (face, hands and arms) in human patients.

Moreover, these results are also indicative that krill extract is useful for anti-wrinkle treatment. The mechanism of ell-trans retinol, which is included in the krill oil, as an anti-wrinkle works as follows:

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- Regeneration and distinctive anti-inflammatory effects

- Improve blood irrigation

- Increases the epidermis regeneration by increasing the rate of cell division and turnover

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- Accelerates the differentiation of keratin

- Regenerates the collager

- Allows cells in the top layer of the skin, which are always being replaces, to mature more normally than untreated sun-damaged cells

- Reduces the activation of enzymes that break down the proteins collagen and elastin that provide structural support for the skin.

The results obtained with krill extract administered on a patent's skin show that the krill extract is having an anti-wrinkle effect by increasing the hydration and the mechanism above described.

<u>Example 6</u> Premenstrual syndrome

Table 10 shows results obtained from the use of krill oil to reduce the pain and mood changes associated with premenstrual syndrome in women. Krill oil extract was administered to 7 women during 2 months. The women were taking 6 capsules of krill extract per day, each capsule containing 800 mg of krill oil. A recommended daily intake of krill oil is of about 1 to 4.8 grams. All participants were advised to continue with their usual nutrition habits and to refrain from initiating any restrictions in their diet. No serious side effects were reported.

All woman enrolled reported noticeable emotional and/or physical discomfort 7 to 10 days prior to menstruation. A self-assessment visual analogue scale validated for the assessment of the premenstrual syndrome, ranging from 0 (no symptoms) to 10 (unbearable) was used as a primary outcome in order to evaluate the effect of krill extract on premenstrual discomfort.

Data analysis has been reported on 60% of the women participating in the study who have completed a two months regimen. The

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majority of the women (73.3%) showed a clinically significant reduction in both emotional and physical distress prior to menstruation (see Table 10).

Table 10

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Frequency distribution of the effect of krill extract on premenstrual syndrome symptomatology

| PMS symptoms | Frequency % | Valid % | Cumulative % |
|--------------|-------------|---------|--------------|
| No change | 26.7 | 26.7 | 26.7 |
| Positive | 73.3 | 73.3 | 100.0 |
| Total | 100.0 | 100.0 | |

Example 7 Diabetes

8 human patients were taking krill extract at the dosage of 6 capsules a day, each capsule containing 800 mg of krill extract, during 2 months. A recommended daily intake of krill oil is of about 1 to 4.8 grams. The Table 11 is showing the variation in the glucose tested for the patients after 2 months.

<u>Table 11</u> Variation in glucose in patients

| | | | Paired | Differences | | | |
|---------------------|-------|--------|-----------------------|---|-------|----|-----------------------|
| Parameter
tested | Mean | SD. | Std.
Error
Mean | 95% Confidence
Interval of the
Difference | | đť | Sig.
(2-
tailed |
| Glucose | .5778 | .60369 | .20123 | .1137 - 1.0418 | 2.871 | 8 | .021 |

A blood glucose decrease of 20% was obtained for the patients taking krill extract, which shows that an uptake of krill extract is controlling blood glucose content and therefore controlling diabetes in human patients.

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While the invention has been described in connection with specific embodiments thereof, it will be understood that it is capable of further modifications and this application is intended to cover any variations, usas, or adaptations of the invention following, in general, the principles of the invention and including such departures from the present disclosure as come within known or customary practice within the art to which the invention pertains and as may be applied to the essential features hereinbefore set forth, and as follows in the scope of the appended claims.

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WHAT IS CLAIMED IS:

 A composition for decreasing cholesterol in a patient comprising an effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil is obtained from a process comprising the steps of:

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 a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

b) separating the liquid and solid contents;

c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanoi, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or aquatic animal material;

e) separating the liquid and solid contents;

f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

2. A composition for decreasing cholesterol in a patient comprising an effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Eicosapentanoic acid, Docosahexanoic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylethanolarmine, Sphingomyelin, α -tocopherol, Astaxanthin, and flavonoid.

 The composition of claim 2, further comprising at least one of the group consisting of Linolsic acid, Alpha-linoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol,

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triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β -carotene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

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4. A method of decreasing cholesterol in a patient, said method comprising administering an effective amount of the composition of any one of claims 1-3 to said patient.

5. The method of claim 4, wherein said administering is effected orally.

6. The method of claim 4, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

7. The method of claim 6, wherein said quantity is 4.8 grams.

8. Use of the composition of any one of claims 1-3 for decreasing cholesterol in a patient.

9. Use of the composition of any one of claims 1-3 for the production of a medicament for decreasing cholesterol in a patient.

10. A composition for inhibiting platelet adhesion and plaque formation in arterles of a patient comprising an effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said knill and/or marine oil is obtained from a process comprising the steps of:

 a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

b) separating the liquid and solid contents;

c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or aquatic animal material;

e) separating the liquid and solid contents;

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f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

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g) recovering the solid contents.

11. A composition for inhibiting platelet adhesion and plaque formation in arteries of a patient comprising an effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Elcosapentanoic acid, Docosahexanoic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, α-tcopherol, Astaxanthin, and flavonoid.

12. The composition of claim 11, further comprising at least one of the group consisting of Linoleic acid, Alpha-linoleic acid, arachidonic acid, oleic acid, palmittle acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β-carotene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

13. A method of inhibiting platelet adhesion and plaque formation in arteries of a patient, said method comprising administering an effective amount of the composition of any one of claims 11-12 to said patient.

14. The method of claim 13, wherein said administering is effected orally.

15. The method of claim 13, wherein said knill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

16. The method of claim 15, wherein said quantity is 4.8 grams.

17. Use of the composition of any one of claims 11-13 for inhibiting platelet adhesion and plaque formation in arteries of a patient.

18. Use of the composition of any one of claims 11-13 for the production of a medicament for inhibiting platelet adhesion and plaque formation in arteries of a patient.

19. A prophylactic composition for preventing hypertension in a patient comprising a prophylactic effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil is obtained from a process comprising the steps of:

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 a) placing krill and/or marine material in a ketone solvent, preferably acelone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

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b) separating the liquid and solid contents;

c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or aquatic animal material;

e) separating the liquid and solid contents;

 f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

20. A prophylactic composition for prevention of hypertension in a patient comprising a prophylactic effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Eicosapentanoic acid, Docosahexanoic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, a-tocopherol, Astaxanthin, and flavonoid.

21. The composition of claim 20, further comprising at least one of the group consisting of Linoleic acid, Alpha-Inoleic acid, arachidonic acid, oleic acid, palmitoleic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β -carotene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

22. A method of preventing hypertension in a patient, said method comprising administering a prophylactic effective amount of the composition of any one of claims 19-21 to said patient.

23. The method of claim 22, wherein said administering is effected orally.

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24. The method of claim 22, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

25. The method of claim 24, wherein said quantity is 4.8 grams

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26. Use of the composition of any one of claims 19-21 for preventing hypertension in a patient.

27. Use of the composition of any one of claims 19-21 for the production of a medicament for preventing hypertension in a patient.

28. A therapeutical composition for symptomatic controlling or treating arthritis comprising a therapeutically effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil is obtained from a process comprising the steps of:

 a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

b) separating the liquid and solid contents;

c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or acuatic animal material;

e) separating the liquid and solid contents;

f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

29. The composition of claim 28, wherein said arthritis is selected from the group consisting of rheumatoid arthritis and osteoarthritis.

30. A therapeutical composition for symptomatic controlling or treating arthritis comprising a therapeutically effective amount of krill and/or

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 marine oil in association with a pharmaceutically acceptable carrier,

 wherein said krill and/or marine oil comprises Eicosapentanoic acid,

 Docosahexanoic acid, Phosphatidylcholine,

 Phosphatidylserine, Phosphatidylethanolamine,

 Sphingomyelin, α

 tocopherol, Astaxanthin, and flavnold.

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31. The composition of claim 30, further comprising at least one of the group consisting of Linoleic acid, Alpha-linoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β -carotene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

32. The composition of claim 30, wherein said arthritis is selected from the group consisting of rheumatoid arthritis and osteoarthritis.

33. A method for symptomatic controlling or treating arthritis in a patient, said method comprising administering a therapeutically effective amount of the composition of any one of claims 29-32 to said patient.

34. The method of claim 33, wherein said administering is effected orally.

35. The method of claim 33, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

36. The method of claim 35, wherein said quantity is 4.8 grams.

 Use of the composition of any one of claims 29-32 for symptomatic controlling or treating rheumatoid arthritis in a patient.

38. Use of the composition of any one of claims 29-32 for the production of a medicament for symptomatic controlling or treating rheumatoid arthritis in a patient.

39. A prophylactic composition for prevention of skin cancer in a patient comprising a prophylactic effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil is obtained from a process comprising the steps of:

 a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

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b) separating the liquid and solid contents;

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c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanot, isopropanoi or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or aquatic animal material;

e) separating the liquid and solid contents;

f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

40. A prophylactic composition for prevention of skin cancer in a patient comprising a prophylactic effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Eicosapentanoic acid, Docosahexanoic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, α -tocopherol, Astaxanthin, and flavonoid.

41. The composition of claim 40, further comprising at least one of the group consisting of Linoleic acid, Alpha-linoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β-carotene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

42. A method of prevention of skin cancer, said method comprising administering a therapeutically or a prophylactic effective amount of the composition of any one of claims 39-41 to a patient.

43. The method of claim 42, wherein said administering is effected orally.

44. The method of claim 42, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

45. The method of claim 44, wherein said quantity is 4.8 grams.

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46. Use of the composition of any one of claims 39-41 for preventing skin cancer in a patient.

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47. Use of the composition of any one of claims 39-41 for the production of a medicament for preventing skin cancer in a patient.

48. A composition for enhancing transdermal transportation for dermatological topical therapeutic applications in a patient comprising an enhancing effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil is obtained from a process comprising the steps of:

 a) placing knll and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

b) separating the liquid and solid contents;

c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from seid merine and/or aquatic animal meterial;

e) separating the liquid and solid contents;

f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

49. A composition for enhancing transdermal transportation for dermatological topical therapeutic applications in a patient comprising an enhancing effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Eicosapentanoic acid, Docosahexanoic acid, Phosphatidyleoline, Phosphatidylisostol, Phosphatidylserine,

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Phosphatidylethanolamine, Sphingomyelin, α -tocopherol, Astaxanthin, and flavonoid.

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50. The composition of claim 49, further comprising at least one of the group consisting of Linoleic acid, Alpha-linoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β-carolene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

51. A method for enhancing transdermal transportation for dermatological topical therapeutic applications in a patient, said method comprising administering an enhancing effective amount of the composition of any one of claims 48-50 to said patient.

52. The method of claim 51, wherein said administering is effected orally and/or topically.

53. The method of claim 51, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

54. The method of claim 53, wherein said quantity is 4.8 grams.

55. Use of the composition of any one of claims 48-50 for enhancing transdermal transportation for dermatological topical therapeutic applications in a patient.

56. Use of the composition of any one of claims 48-50 for the production of a medicament for enhancing transdermal transportation for dermatological topical therapeutic applications in a patient.

57. A composition for enhancing transdermal transportation for dermatological cosmetic applications in a patient comprising an enhancing effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil is obtained from a process comprising the steps of:

 a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

b) separating the liquid and solid contents;

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 c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

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d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or aquatic animal material;

e) separating the liquid and solid contents;

f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

58. A composition for enhancing transdermal transportation for dermatological cosmetic applications in a patient comprising an enhancing effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Eicosapentanoic acid, Docosahexanoic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, α-tocopherol, Astaxanthin, and flavonoid.

59. The composition of claim 58, further comprising at least one of the group consisting of Linoleic acid, Alpha-linoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, ali-trans retinol, canthaxanthin, β-carotene, zinc, selenium, nervoric acid, sodium, potassium and calcium.

60. A method for enhancing transdermal transportation for dermatological cosmetic applications, said method comprising administering an enhancing effective amount of the composition of any one of claims 57-59 to a patient.

61. The method of claim 60, wherein said administering is effected orally and/or topically.

62. The method of claim 60, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

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63. The method of claim 62, wherein said quantity is 4.8 grams.

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64. Use of the composition of any one of claims 57-59 for enhancing transdermal transportation for dermatological cosmetic applications in a patient.

65. Use of the composition of any one of claims 57-59 for the production of a medicament for enhancing transdermal transportation for dermatological cosmetic applications in a patient.

66. A composition for reducing premenstrual syndrome's symptoms in a patient comprising an enhancing effective amount of knill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said knill and/or marine oil is obtained from a process comprising the steps of:

 a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

b) separating the liquid and solid contents;

c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or aquatic animal material;

e) separating the liquid and solid contents;

f) recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

67. A composition for reducing premenstrual syndrome's symptoms in a patient comprising an enhancing effective amount of knil and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Elcosapentanolc acid, Doccsahexanoic acid, Phosphatidylcholine, Phosphatidylinositel.

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Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, αtocopherol. Astaxanthin, and flavonoid.

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68. The composition of claim 67, further comprising at least one of the group consisting of Linoleic acid, Alpha-inoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β-carotene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

69. A method for reducing premenstrual syndrome's symptoms in a patient, said method comprising administering an enhancing effective amount of the composition of any one of claims 66-68 to said patient.

70. The method of claim 69, wherein said administering is effected orally.

71. The method of claim 69, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

72. The method of claim 71, wherein said quantity is 4.8 grams.

73. Use of the composition of any one of claims 66-58 for reducing premenstrual syndrome's symptoms in a patient.

74. Use of the composition of any one of claims 66-68 for the production of a medicament for reducing premenstrual syndrome's symptoms in a patient.

75. A composition for controlling blood glucose level in a patient comprising an enhancing effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil is obtained from a process comprising the steps of:

 a) placing krill and/or marine material in a ketone solvent, preferably acetone to achieve extraction of the soluble lipid fraction from said marine and/or aquatic animal material;

b) separating the liquid and solid contents;

c) recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents;

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d) placing said solid contents in an organic solvent selected from the group of solvents consisting of alcohol, preferably ethanol, isopropanol or t-butanol and esters of acelic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from said marine and/or aquatic animal material;

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e) separating the liquid and solid contents;

 recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and

g) recovering the solid contents.

76. A composition for controlling blood glucose level in a patient comprising an enhancing effective amount of krill and/or marine oil in association with a pharmaceutically acceptable carrier, wherein said krill and/or marine oil comprises Elcosapertanoic acid, Docosahexanoic acid, Phosphatidylcholine, Phosphatidylinositol, Phosphatidylserine, Phosphatidylethanolamine, Sphingomyelin, α-tocopherol, Astaxanthin, and flavonoid.

77. The composition of claim 76, further comprising at least one of the group consisting of Linoleic acid, Alpha-linoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinol, canthaxanthin, β-carotene, zinc, selenium, nervonic acid, sodium, potassium and calcium.

78. A method for controlling blood glucose level in a patient, said method comprising administering an enhancing effective amount of the composition of any one of claims 75-77 to said patient.

79. The method of claim 78, wherein said administering is effected orally.

80. The method of claim 78, wherein said krill and/or marine extract is administered in a quantity in a range of 1 to 4.8 grams per day.

81. The method of claim 80, wherein said quantity is 4.8 grams.

82. Use of the composition of any one of claims 75-77 for controlling blood glucose level in a patient.

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83. Use of the composition of any one of claims 75-77 for the production of a medicament for controlling blood glucose level in a patient.

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| This International Search Report has not been established in respect of certain claims under A | uticle 17(2)(a) for the following reasons: |
| 1. X Claims Nos.: because they relate to subject matter not required to be saarched by this Authority, n | |
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| This international Searching Authority found multiple Inventions in this international application | , aa foliowa: |
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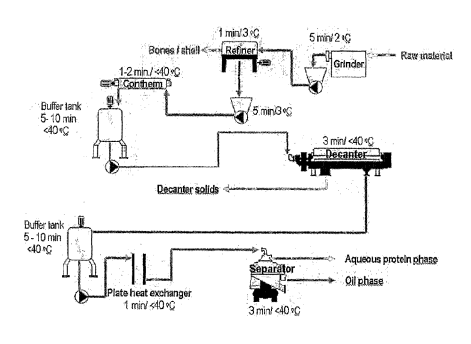
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[Continued on next page]



(57) Abstract: In accordance with the present disclosure there is provided a novel marine lipid extract obtainable by a process wherein processing temperature below 60 0C; mechanical and physical disruption of the lipid cell membrane to facilitate low temperature extraction; processing takes place under inert gas to prevent oxidation or denaturation of fat and proteins; intermediate processing tanks kept at a minimum level to reduce residence time; and the oil is frozen immediately after recovery to stabilize it.

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THROMBOSIS PREVENTING KRILL EXTRACT

FIELD OF THE INVENTION

This invention relates to novel extracts derived from krill, which can prevent and/or treat thrombosis. This invention also relates to a method for the extraction of lipid fractions from krill in order to obtain the novel extracts of the present invention. More specifically, the invention relates to an improved method of extracting lipid fractions without using high temperatures and/or organic solvents.

BACKGROUND OF THE INVENTION

Krill is the common name for small, shrimp-like crustaceans that swarm in dense shoals, especially in Antarctic waters. It is one of the most important food sources (especially protein) for fish, some kind of birds and especially for baleen. Krill is also a good source of omega-3 fatty acids, which are well known for their beneficial effects on human health.

It is known in the art to use krill and/or marine enzymes for the treatment of a great variety of diseases in human and animals such as infections, inflammations, cancers, HIV/AIDS, pain, polyps, warts, hemorrhoids, plaque, wrinkles, thin hair, allergic itch, anti-adhesion, eye disease, acne, cystic fibrosis and immune disorders including autoimmune diseases and cancer.

It is also known in the art that krill and/or marine oils may be used for the treatment of autoimmune murine lupus and other autoimmune diseases and can also be used for treating cardiovascular diseases.

However, most of the krill oil extracts used for these treatments has only conserved its omega-3 fatty acids as active ingredients, which is a very small part of all the active ingredients of the krill itself. This fact dramatically reduces the potential of the krill and/or marine oil as a treatment for these diseases.

There is an increasing demand for treatments using products derived from a natural source, therefore, it would be highly desirable to be provided with a krill and/or marine extract having an enhanced potential for prevention and/or treatment and/or management of disease. US Patent 6,800,299 discloses a method for extracting lipid fractions from marine and aquatic animal material by acetone extraction. The resulting non-soluble and particulate fraction is preferably subjected to an additional solvent extraction with an alcohol, preferably ethanol,

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isopropanol or t-butanol or an ester of acetic acid, preferably ethyl acetate to achieve extraction of the remaining soluble lipid fraction from the marine and aquatic animal material. The remaining non-soluble particulate content is also recovered since it is enriched in proteins and contains a useful amount of active enzymes. Also provided herein is a krill extract. It is reported that these marine and aquatic animal oils have anti-inflammatory properties. Marine and aquatic animal oils are also reported as helpful in reducing the incidence of cardiovascular disease. As a further example the patent mentions that krill may be used as a source of enzymes for debridement of ulcers and wounds or to facilitate food digestion.

WO02102394A2 discloses a process for the preparation of a krill oil extract, which process includes the steps of placing krill and/or marine material in a ketone solvent to achieve extraction of the soluble lipid fraction from the krill; then separating the liquid and solid contents; then recovering a first lipid rich fraction from the liquid contents by evaporation of the solvent present in the liquid contents; then placing the solid contents in an organic solvent to achieve extraction of the remaining soluble lipid fraction from the krill material; then separating the liquid and solid contents; then recovering a second lipid rich fraction by evaporation of the solvent from the liquid contents; and finally recovering the solid contents. Diseases that can be treated and/or prevented by using the krill oil extract are *inter alia* cardiovascular diseases. In this respect it is mentioned that the Krill oil has been shown to decrease cholesterol *in vivo*, inhibit platelet adhesion and plaque formation and reduce vascular endothelial inflammation in a patient.

Canadian Patent 1,098,900 describes a method for extracting oils and producing proteins from krill comprising emulsification of lipids of krill in an aqueous medium, separation of the emulsion of lipids from the krill mass, alkaline extraction of proteins from the krill mass, separation of the protein extract produced from chitin integuments, and finally separation of protein from the protein extract. The document mentions that krill is a prospective source of food and other practically useful products such as chitin and lipids which find wide application in different branches, such as food industry, textile, and medicine.

WO03011873A2 discloses a phospholipid extract from *inter alia* krill, with therapeutic properties, such as those essential for the maintenance of a healthy cardiovascular system. The phospholipid extract comprises a variety of phospholipids, fatty acid, metals and a novel flavonoid. The method for the preparation of this extract is generally carried out by a method similar to the one described in US Patent 6,800,299 (see above; includes organic solvents),

which procedure produces two successive lipid fractions and a dry residue enriched in protein, including active enzymes.

WO8401715A1 and WO09533471A1 disclose various aspects of so-called krill enzymes, which are water-soluble. It is mentioned that in krill a mixture of different enzymes exists, such as e.g. proteinases (with acidic and neutral-to-alkaline pH-optima), peptidases (exo- and endopeptidases), lipases, phospholipases, amylases and other carbohydrate degrading enzymes, phosphatases nucleases, nucleotidases and esterases. The proteolytic (trypsin-like) activity existing in a water extract from krill has been studied and described. WO09533471A1 disclose the use of one or more krill enzymes for the manufacture of an intravasal pharmaceutical composition for thrombolysis in a mammal host.

The potential of krill oil to prevent thrombosis has been disclosed in the prior art; however such a preventive effect has so far only been ascribed to the presence of powerful antioxidants and the special composition of poly-unsaturated fatty acids. The present inventors have surprisingly found that krill oil prepared by a novel process, which is from a physical-chemical point of view very gentle to the krill material due to relatively low temperature and no use of organic solvents, comprises other therapeutically valuable components than known from conventional krill oil extracts as well as other known fish oil; such components include inter alia high molecular (MWt > 200 kDa) hydrophobic proteins.

SUMMARY OF THE INVENTION

In accordance with the present invention there is provided a novel krill oil extract for the prevention and/or treatment of thrombosis.

The general extraction method of the present invention will now be described. The starting material, consisting of freshly harvested and preferably finely divided krill material, is subjected to extraction, for about two hours and preferably overnight. However, extraction time is not critical to the yield of lipid extraction. To facilitate extraction, it is preferable to use particles of less than 0.5 mm in diameter. Extraction is preferably conducted under inert atmosphere and at a temperature in the order of about 5° C or less. The inventors have also envisaged that the present invention may be carried out by applying supercritical CO2 extraction.

Preferably, the beginning of the extraction will be conducted under agitation for about 10 to 40

minutes, preferably 20 minutes. The solubilized lipid fractions are separated from the solid material by standard techniques including, for example, filtration, centrifugation or sedimentation. Filtration is preferably used.

DETAILED DESCRIPTION OF THE INVENTION

In accordance with the present invention, there is provided a novel krill extract for prevention and/or treatment and/or therapy of thrombosis.

The novel oil extract is derived from krill found in any marine environment around the world, for example, the Antarctic ocean (euphasia superba), the Pacific ocean (euphasia pacifica), the Atlantic ocean, the Indian ocean, in particular coastal regions of Mauritius Island and/or Reunion Island of Madagascar, Canadian West Coast, Japanese Coast, St-Lawrence Gulf and Fundy Bay, and this oil extract is a lipid fraction.

According to a first aspect of the present invention there is provided a method for extracting lipid fractions from krill, said method comprising the steps of:

- placing the krill material in a blender to mechanically disrupt fat cell membranes;
- separating the liquid and solid components;
- recovering a lipid rich fraction from the liquid component;

wherein the extraction is performed quickly at a temperature below 60 °C and does not involve the use of organic solvents.

According to another aspect of the present invention there is provided a method for extracting lipid fractions from krill, said method comprising the steps of:

- Feeding freshly captured krill into a grinder to produce a slurry
- Heating the slurry gently to a temperature below 90°C for less than 45 minutes
- Separating the solid material from the liquid
- Separating the liquid into an aqueous phase and a krill oil phase

wherein the extraction does not involve the use of organic solvents.

According to the invention there is also provided a pharmaceutical composition for the treatment of thrombosis in a patient comprising an effective amount of a krill oil extract obtainable by a method according to the present invention.

EXAMPLES

As used herein, the term omega-3 fatty acid refers to polyunsaturated fatty acids that have the final double bond in the hydrocarbon chain between the third and fourth carbon atoms from the methyl end of the molecule. Non-limiting examples of omega-3 fatty acids include, but are not limited to 5,8,11,14,17-eicosapentaenoic acid (EPA), 4,7,10,13,16,19-docosahexanoic acid (DHA) and 7,10,13,16,19-docosapentanoic acid (DPA).

Example 1

Preparation of the krill oil extract of the present invention (see also Fig 1).

Preparation of the krill oil

The method of preparation is a continuous flow process and so the times given represent the average time that the material is in each stage of the process and the temperatures are typical (and may vary by $\pm 3^{\circ}$ C).

- 1. The freshly captured krill are fed into the grinder together with process water and shredded at 2°C for 5 minutes.
- This is then fed into a refiner which separates the chitin shell from the slurry (1 minute, 3°C).
- 3. The slurry is then passed into a heat exchanger and warmed gently up to a temperature about 35°C (max below 40°C) (1-2 minutes) and then stored in a buffer tank for 5 to 10 minutes. All subsequent processes occur at temperatures below 40°C.
- 4. A centrifugal decanter is then used to separate the solid material from the liquid (3 minutes).
- 5. The liquid fraction is then stored in a buffer tank for 5 to 10 minutes.
- 6. The temperature of the liquid is adjusted to 35 °C using a countercurrent plate heat exchanger (1 minute).
- 7. The liquid is then separated into an aqueous phase and a krill oil phase.

Preparation of the stock solution of krill oil

- 1. Autoclave glycerol (analytical quality) and leave to cool to room temperature
- 2. Mix 100 µL of krill oil with 1000 µL autoclaved glycerol
- 3. Shake mixture for 6 min. on a minibead beater (Biospec. Products, USA) at room temperature
- Add 900 μL diluted CPD solution (Compoflex[®], Fresenius HemoCare 61348 Bad Hamburg, Germany containing: citric acid monohydrate 3.27g, Sodium citrate dihydrate 26.3g, sodium dihydrogen phosphate dehydrate 2.51g, glucose monohydrate 25.3g made up to 1L).
- 5. Shake mixture for 6 min. on minibead beater 4 times at room temperature

Preparation of the dilute solutions of krill oil

- 1. The stock is diluted sequentially (1:10), shaking for 6 minutes at each dilution.
- 2. Immediately before use, shake mixture for 6 min. on a minibead beater at room temperature

Preparation of the stock solution of CPD Glycerol control solution

- 1. Mix 100 µL of diluted CPD solution with 1000 µL autoclaved glycerol
- 2. Shake mixture for 6 min. on minibead beater at room temperature
- 3. Add 900 µL diluted CPD solution
- 4. Shake mixture for 6 min. on minibead beater 2 times at room temperature
- 5. Repeat 4. immediately before use

Example 2

Effect of the krill oil extract of the present invention on the aggregation time of thrombocytes.

Preparation of human blood

Blood samples were taken from normal subjects. 3.8 mm plastic tubes containing 0.38 ml 0.129M sodium citrate buffer (CPD buffer, pH 5.5) were used to store the blood. The buffered blood was then mixed with the krill or fish oil to achieve a final oil concentration varying from $5x10^{-2}$ to $5x10^{-18}$ Vol%. The blood cells were treated with krill or fish oil for 60 minutes before aggregation tests were performed.

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Blood aggregation time

The thrombocyte aggregation tests were performed with a PFA 100 aggregometer (Dade Bering), which is a microprocessor controlled apparatus with single test vials. The unit comprises a small reservoir, a capillary and a membrane, which is covered with 2 mg genuine, type 1 collagen and 50 mg adenosin-5'-diphosphate (ADP). The blood is pipetted directly into the reservoir and aspirated through a capillary with a diameter of 200 μ m with a constant negative pressure resulting in high shear stress. The capillary ends with a membrane having an aperture with a diameter of 150 μ m. The thrombocytes are then activated by collagen and ADP. Upon aggregation the blood flow is stopped due to clogging, which is referred to as closing time. The test automatically stops after 300 seconds. The normal value is between 62.5 – 120.5 seconds for ADP.

Determination of anti-aggregation effect

Dilute krill oil solutions were added to whole human blood samples and allowed to react in accordance with the following steps:

- 1. Serial dilutions of the krill oil or other oils under investigation were added to the human blood samples and gently shaken for 1 hour at room temperature on a "HETO-blood turner" at a rotational speed of 10 rpm.
- Exactly 800μL of the blood-oil sample were placed in the reaction cartridge (DADE PFA collagen/epitest cartridge containing 4μg epinephrine bitartrate and 2μg type 1 equine collagen). The blood was then allowed to clot at 37°C for up to 300 seconds (preset instrument maximum).
- 3. Measurements were read from the display and printed out recorded

Figure 2 shows the effects of various oils on the rate of aggregation of human whole blood.

Samples of human whole blood are aggregated at the start and end of every experiment ("Start blood" and "End blood" on the abscissa) to determine the rate of blood aggregation for the donor. As an additional control carried out just after and just before the start and end whole blood aggregation determinations, an aliquot of the vehicle is added and the aggregation determination is repeated ("Start Glycerol/CPD" and "End Glycerol/CPD"). These controls are performed to ensure that the ability of the blood to aggregate does not change during the experimentation (see the trend line for the glycerol/CPD points). Between these control experiments, the blood is treated (as described in the text) with various concentrations of the different oils for 1hr before its ability to aggregate is determined. Dotted line - fish oil; dashed

line a commercially available krill oil; solid line krill oil prepared in the manner disclosed here. The graph shows typical data from a single patient.

As can be seen in Figure 2, the fish oil can be diluted to a concentration of only about 1×10^{-4} before it looses its effect. A commercially available krill oil can be diluted to about 5×10^{-6} before it looses its effect (i.e. it is about 500 times more effective than fish oil). Krill oil prepared in the manner described here can be diluted to a concentration of about 5×10^{-12} before it looses its effect. This is a million times more effective than the existing krill oil preparations and five hundred million times better than fish oil (note that the abscissa is a logarithmic scale).

Figure 3 (graph 1) shows the inhibiting effect of krill oil on the aggregation of thrombocytes in blood samples from 6 subjects. It also appears that the effect varies from subject to subject; and furthermore blood from one of the subjects was not influenced at all by the presence of krill oil.

In Figure 4 (graph 2) the effect of krill oil C on blood from the same subject was analysed twice with a 21 day interval. The effect of krill oil C on the aggregation of thrombocytes is significant; however it must be concluded that the difference in the concentration required to achieve a significant inhibition varies with more than 10^{-3} Vol%.

Example 3

Comparison of krill oils and fish oils with respect to the effect on blood aggregation

These experiments serve to demonstrate that the krill oil obtainable by the process of the present invention prevents formation of thrombosis (based on the same experimental procedure as laid down in Examples 1 and 2) to a higher degree than known krill oils and other fish oils.

The experiments include 2 fish oils as well as 3 different krill oils:

| Krill oil A: | Krill caught in large nets and subjected to a long process |
|--------------|--|
| | time |
| Krill oil B: | Krill caught in smaller nets and subjected to a short process |
| | time |
| Krill oil C: | Krill sucked up and processed very rapidly (in accordance with the |
| | present invention) |
| Fish oil A: | Newly cold pressed cod fish oil |
| Fish oil B: | Pikasol (OTC registered natural pharmaceutical containing |
| | concentrated Omega-3 rich fish oil; contains 62% omega-3 fatty |
| | acids, mainly EPA and DHA; Pikasol is produced from highly |
| | refined fish oil from the cleanest oceans in the world) |

The oils are dissolved in a 1:1 mixture with glycerol and CPD (Gly/CPD-mixture). Every single dillution is performed with the Gly/CPD-mixture to ensure that the glycerol concentration remains constant about $5x10^{-3}$ Vol%.

It is known that the quality of krill oil may vary considerably due to the way the krill material has been "caught". As discussed above the prior envisages that the amount of phosholipids, omega-3 and omega-6 polyunsaturated fatty acids and various antioxidants is responsible for the therapeutic effects attributable to krill oil. As appears from Fig 5 (graph 3) the three krill oils have very different effects on the aggregation. Surprisingly, the different effects could not be ascribed to differences in the amount of e.g. polyunsaturated fatty acids. On the contrary it appeared (based on 2D gel electrophoresis) that 5 proteins were present in Krill oil C (according to the present invention) but only in minute amounts in krill oil B and not traceable in krill oil A. This observation stems with the fact that many proteins in krill are extremely sensible for proteolytic degradation, which starts right after the krill has been caught.

As already mentioned the therapeutic effect of antioxidants and polyunsaturated fatty acids from fish oil on cardiovascular diseases is well known. Accordingly, the present inventors have compared the effect the effect of Krill oil C and fish oils A and B with respect to their ability to prevent thrombose formation (verified with the above described aggregation test). Fig 6 (graph 4) demonstrates that Krill oil C (according to the present invention) has a far more pronounced inhibitory effect on the thrombocyte aggregation than is the case with the fish oils.

Conclusions drawn from Examples 1-3

Based on the experimental evidence provided so far the following conclusions may be drawn:

- Krill oil prepared by the process according to the present invention has a strong inhibitory effect on human thrombocyte aggregation in blood samples,
- The difference between the intensity of the effect may possibly be ascribed to certain proteins of the krill oil, and
- There is a substantial difference between how blood from different subjects responds to the krill oil with respect to aggregation time, however it may validly said that the krill oil obtained with the process of the present invention is far more effective that krill oils and fish oils obtained by traditional high temperature/solvent extraction methods.

Example 4

Phospholipids are to be extracted from the solid fraction obtained in example 1 (step 4) using ethanol. After removal of the ethanol, the phospholipids are to be mixed with the krill oil phase obtained from the liquid fraction in example 1 (step 7) into a krill oil composition. The anti-thrombotic effects of this krill oil composition are to be compared with other krill oil products extracted with organic solvents by investigating the effect on the aggregation time of thrombocytes in-vitro. The krill oil products (mixtures of krill triglycerides and krill phospholipids) for this comparison are to be extracted from krill or krill meal using organic solvents as described in US 6,800,299. It is to be observed that the anti-thrombotic effects of the krill oil product extracted with organic solvents such as acetone.

Example 5

The krill oil compositions tested in example 4, the krill oil extracted obtained in example 1 (step 7), krill oil obtained using organic solvents and a control are to be administered in humans (invivo) for a period of 5 weeks. Diets are to contain approximately 38% of energy as fat excluding the lipid in the supplement. Around 2 g of each product are to be administered in a way that preserves the biological effect of the krill oil. Non-limiting examples of administration are oral, sublingual or transdermal. After termination of the experiment, ex vivo and in vitro platelet aggregation, and variables of coagulation, fibrinolysis, and hematology are to be evaluated. Ex vivo platelet aggregation time are to be measured by filtragometry and in vitro platelet aggregation induced by collagen and ADP measured by PFA 100 aggregometer. Variables of coagulation (factor VII amidolytic activity and concentrations of fibrinogen and prothrombin fragment 1 and 2) and fibrinolysis [plasminogen activator inhibitor (PAI) activity

and concentrations of tissue plasminogen activator (tPA)/PAI-1 complexes] are to be determined by standard methods. It is to be observed that the subjects treated with the krill oil composition described in example 4 and the krill lipid extract obtained in example 1 (step 7) show superior anti-thrombotic activity than subjects treated with krill oil compositions obtained using organic solvents and control. Prevention of thrombosis is linked to prevention of myocardial infarction and stroke. Hence, the krill oil composition described in example 4 and example 1 (Step 7) can be used to prevent these pathologies.

CLAIMS

1. A method for extracting lipid fractions from krill, said method comprising the steps of:

- placing the krill material in a grinder or blender to mechanically disrupt cell membranes;
- separating the liquid and solid components;
- recovering a lipid rich fraction from the liquid component;

wherein the extraction is performed at a temperature below 60 °C and does not involve the use of organic solvents.

2. A method as in claim 1, wherein separating the liquid and solid components is effected by techniques selected from the group consisting of mechanical pressing, filtration, centrifugation and sedimentation.

3. A method as in claim 1, wherein the extraction is performed at a temperature below 27°C, preferably below 15°C, more preferably below 5°C.

4. A krill oil extract obtainable by a method according to any one of claims 1 to 3.

5. A krill oil extract according to claim 4 for use as a medicament.

6. A pharmaceutical composition comprising the krill oil extract of claim 4.

7. A pharmaceutical composition for the treatment of thrombosis in a patient comprising an effective amount of a krill oil extract obtainable by a method according to any one of claims 1 to 3 in association with a pharmaceutically acceptable carrier.

8. The composition of claim 7, further comprising at least one of compounds selected from the group consisting of glycerol, dimethyl-sulphoxide (DMSO), linoleic acid, alpha-linoleic acid, arachidonic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, cholesterol, triglycerides, monoglycerides, all-trans retinal, canthexanthin, carotene, zinc, selenium, sodium, potassium and calcium.

9. Use of the krill oil extract obtainable by the method of any one of claims 1-3 for the production of a medicament for decreasing development of thrombosis in a patient.

10. A composition for inhibiting platelet adhesion and plaque formation in arteries of a patient comprising an effective amount of krill oil extract in association with a pharmaceutically acceptable carrier, wherein said krill oil extract is obtainable from a method according to any one of claims 1 to 3.

11. A krill oil extract obtainable by a method comprising the steps of:

- Feeding freshly captured krill into a grinder to produce a slurry
- Heating the slurry gently to a temperature below 90°C in less than 45 minutes
- Separating the solid material from the liquid
- Separating the liquid into an aqueous phase and a krill oil phase

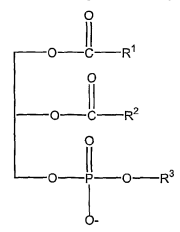
wherein the extraction does not involve the use of organic solvents.

12. A food product comprising the krill oil extract of claim 4.

13. An animal feed comprising the krill oil extract of claim 4.

14. A food supplement comprising the krill oil extract of claim 4.

15. A composition comprising the krill oil extract of claim 4 and phospholipids, said phospholipids having the following structure:



wherein R1 is a fatty acid, R2 is a fatty acid, and R3 is selected from the group consisting of H or choline, ethanolamine, inositol or serine.

16. The composition in claim 15, wherein at least 1% (w/w) of the said fatty acids are unsaturated fatty acids.

17. The composition in claim 15, wherein at least 1% (w/w) of the said fatty acids are omega-3 fatty acids.

18. A food product comprising the composition in any of the claims 15 to 17.

19. An animal feed comprising the composition in any of the claims 15 to 17

20. A food supplement comprising the composition in any of the claims 15 to 17.

21. A pharmaceutical comprising the composition in any of the claims 15 to 17.

22. A method of preventing platelet adhesion in a patient comprising administering to said patient a therapeutically effective amount of the composition in any of the claims 15 to 21.

23. A method for preventing stroke or heart attack in a patient comprising administering to said patient a therapeutically effective amount of the composition in any of the claims 15 to 21.

24. A method of preventing platelet adhesion and plaque formation in a patient comprising administering to said patient a therapeutically effective amount of krill oil, wherein said krill oil is obtained without organic solvent extraction.

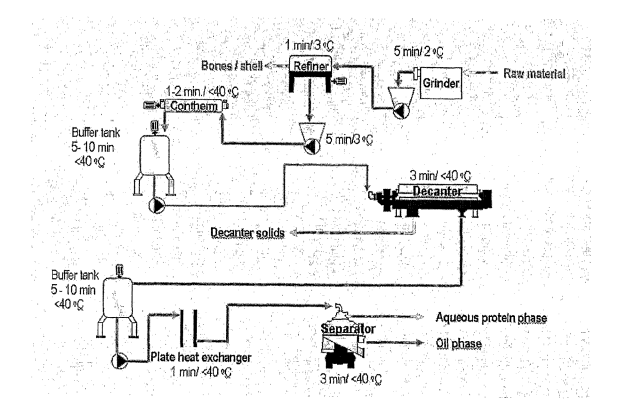
25. A method of preventing platelet adhesion and plaque formation in a patient comprising administering to said patient a therapeutically effective amount of krill oil composition, wherein said krill oil composition comprises triglyceride, phospholipid and protein fractions.

26. The method of claim 25, wherein said protein fraction comprises high molecular weight hydrophobic proteins.

27. A composition comprising a krill oil extract isolated from krill comprising triglyceride, phospholipid and protein fractions.

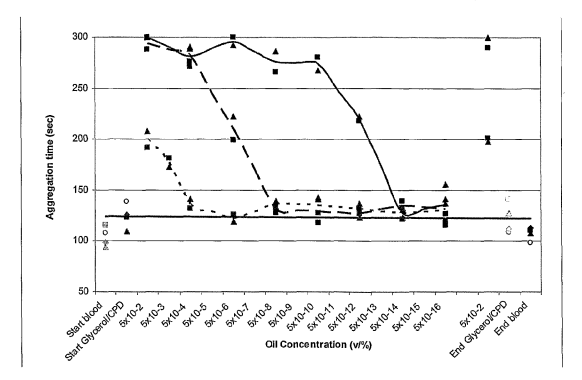
28. A pharmaceutical comprising the composition of claim 27.

Figure 1



RIMFROST EXHIBIT 1024 page 0862

Figure 2



RIMFROST EXHIBIT 1024 page 0863



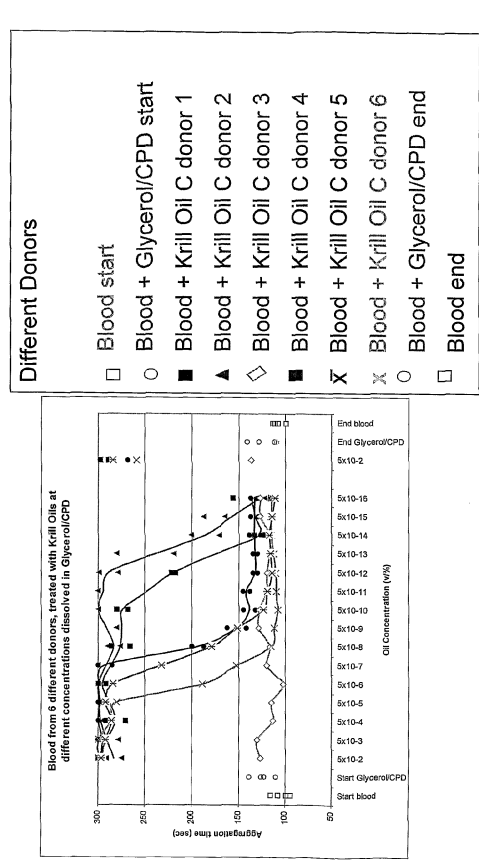
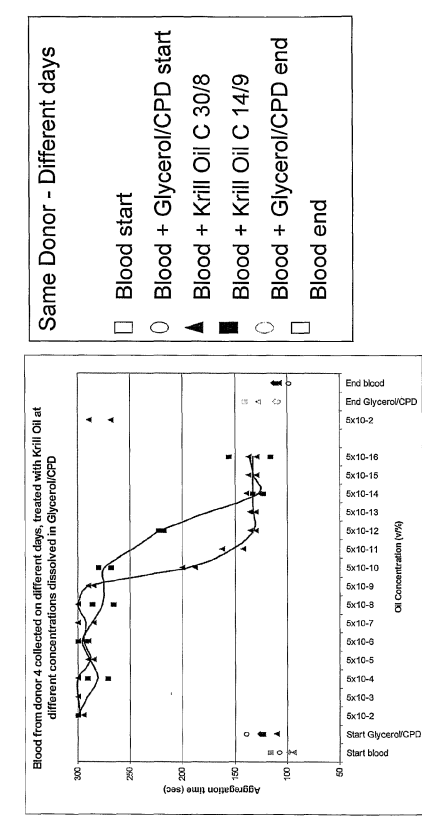


Figure 4



WO 2007/080515

4/6

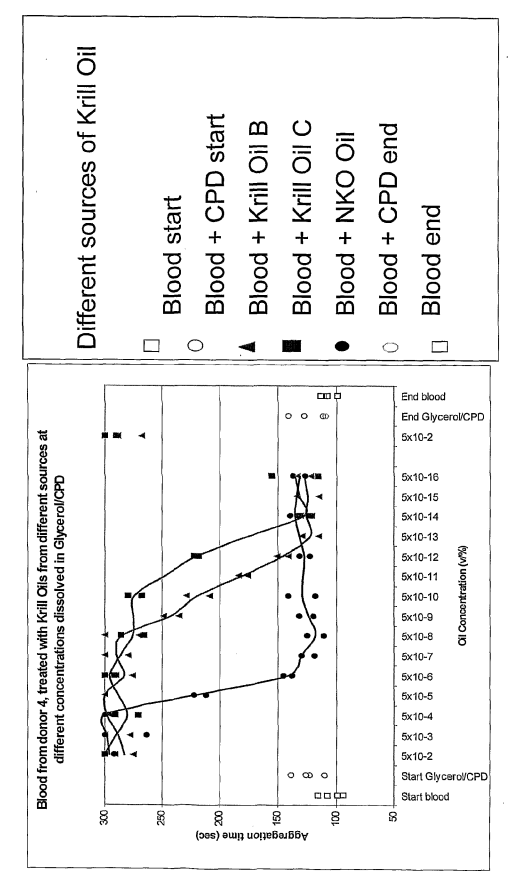


Figure 5

Blood + CPD start Blood + Krill Oil C Blood + CPD end Blood + Fish Oil Blood + Fish Oil Different treatment Blood start Blood end (Pikasol) \bigcirc \bigcirc шоф End blood End Glycerol/CPD 0 0 Ø 5x10-2 Blood from donor 4, treated with Krill and Fish Oils at different 5x10-16 5x10-15 concentrations dissolved in Glycerol/CPD 5x10-14 5x10-13 Oil Concentration (v/%) 5x10-12 5x10-11 5x10-10 5x10-9 5x10-8 5x10-7 5x10-6 5x10-5 5x10-4 5x10-3 5x10-2 00 0 Start Glycerol/CPD 0 Start blood 300 250 100 22 200 150 (cee) emit noitspergeA

Figure 6

6/6

RIMFROST EXHIBIT 1024 page 0867

| | INTERNATIONAL SEARC | H REPORT | International application No
PCT/IB2007/000099 |
|--|---|--|--|
| A. CLASS | AG1P7/02 AG1P9/10 AG1P9/
A23L1/30 A23D9/00 | /14 C11B1, | /02 C11B1/14 |
| According | to International Patent Classification (IPC) or to both national clas | sification and IPC | |
| | SEARCHED | | |
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A23L A61P A23D | lication symbols) | |
| Documenta | ation searched other than minimum documentation to the extent t | hat such documents are inc | cluded in the fields searched |
| | data base consulted during the international search (name of dat
Iternal, CHEM ABS Data, WPI Data, | | al, search terms used) |
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| X Furth | ner documents are listed in the continuation of Box C. | X See patent fai | i mily annex. |
| A' docume
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P' docume | Int which may throw doubts on priority claim(s) or
is cited to establish the publication date of another
n or other special reason (as specified)
ent referring to an oral disclosure, use, exhibition or
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ent published prior to the international filing date but | or priority date an
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Ind not in conflict with the application but
nd the principle or theory underlying the
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International application No
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INTERNATIONAL SEARCH REPORT

International application No. PCT/IB2007/000099

| Box II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet) |
|---|
| This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons: |
| 1. X Claims Nos.:
because they relate to subject matter not required to be searched by this Authority, namely:
Although claims 22-26 are directed to a method of treatment of the |
| human/animal body, the search has been carried out and based on the alleged effects of the compound/composition. |
| 2. Claims Nos.:
because they relate to parts of the International Application that do not comply with the prescribed requirements to such
an extent that no meaningful International Search can be carried out, specifically: |
| 3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). |
| Box III Observations where unity of invention is lacking (Continuation of item 3 of first sheet) |
| This International Searching Authority found multiple inventions in this international application, as follows: |
| 1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims. |
| 2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. |
| 3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.: |
| 4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: |
| Remark on Protest |
| No protest accompanied the payment of additional search fees. |

Form PCT/ISA/210 (continuation of first sheet (2)) (January 2004)

RIMFROST EXHIBIT 1024 page 0871

| | | ATIONAL SEAR(
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| APPLICATION NO. | FILING DATE | FIRST NAMED INVENTOR | ATTORNEY DOCKET NO. | CONFIRMATION NO. |
| 12/057,775 | 03/28/2008 | Inge Bruheim | NATNUT-14409/US-5/ORD | 1945 |
| 72960
Casimir Jones, | 7590 01/06/2012
S.C. | | EXAM | INER |
| 2275 DEMING | WAY, SUITE 310 | WAY, SUITE 310 | | BORAH K |
| MIDDLETON, | WI 53562 | | ART UNIT | PAPER NUMBER |
| | | | 1651 | |
| | | | | |
| | | | MAIL DATE | DELIVERY MODE |
| | | | 01/06/2012 | PAPER |

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

| | Application No. | Applicant(s) |
|--|--|---|
| | 12/057,775 | BRUHEIM ET AL. |
| Office Action Summary | Examiner | Art Unit |
| | DEBBIE K. WARE | 1651 |
| The MAILING DATE of this communication app
Period for Reply | ears on the cover sheet with the c | correspondence address |
| A SHORTENED STATUTORY PERIOD FOR REPLY
WHICHEVER IS LONGER, FROM THE MAILING D/ Extensions of time may be available under the provisions of 37 CFR 1.13
after SIX (6) MONTHS from the mailing date of this communication. If NO period for reply is specified above, the maximum statutory period w Failure to reply within the set or extended period for reply will, by statute,
Any reply received by the Office later than three months after the mailing
earned patent term adjustment. See 37 CFR 1.704(b). | ATE OF THIS COMMUNICATION
36(a). In no event, however, may a reply be tin
vill apply and will expire SIX (6) MONTHS from
, cause the application to become ABANDONE | N.
nely filed
the mailing date of this communication.
D (35 U.S.C. § 133). |
| Status | | |
| 1) Responsive to communication(s) filed on <u>31 O</u> 2a) This action is FINAL. 2b) This 3) An election was made by the applicant in responsive in the restriction requirement and election 4) Since this application is in condition for allowar closed in accordance with the practice under E | action is non-final.
onse to a restriction requirement
have been incorporated into this
nce except for formal matters, pro | s action.
Disecution as to the merits is |
| Disposition of Claims | | |
| 5) Claim(s) <u>1-90</u> is/are pending in the application. 5a) Of the above claim(s) <u>1-54 and 56-90</u> is/are 6) Claim(s) is/are allowed. 7) Claim(s) <u>50-55</u> is/are rejected. 8) Claim(s) is/are objected to. 9) Claim(s) are subject to restriction and/or | e withdrawn from consideration. | |
| Application Papers | | |
| 10) The specification is objected to by the Examine 11) The drawing(s) filed on <u>7/16/08 and 3/28/08</u> is/a Applicant may not request that any objection to the a Replacement drawing sheet(s) including the correct 12) The oath or declaration is objected to by the Example of the second se | are: a) accepted or b) \Box object
drawing(s) be held in abeyance. See
ion is required if the drawing(s) is ob | e 37 CFR 1.85(a).
jected to. See 37 CFR 1.121(d). |
| Priority under 35 U.S.C. § 119 | | |
| 13) Acknowledgment is made of a claim for foreign a) All b) Some * c) None of: 1. Certified copies of the priority documents 2. Certified copies of the priority documents 3. Copies of the certified copies of the priority application from the International Bureau * See the attached detailed Office action for a list | s have been received.
s have been received in Applicati
rity documents have been receive
a (PCT Rule 17.2(a)). | ion No
ed in this National Stage |
| Attachment(s) 1) | 4) Interview Summary
Paper No(s)/Mail D
5) Notice of Informal F
6) Other: | ate |

Office Action Summary RIMFROST EXHIBIT 1024 page 0874

DETAILED ACTION

Claims 1-90 are pending.

Information Disclosure Statement

The information disclosure statements (IDSs) submitted on March 8, 2011, were received. The submission are in compliance with the provisions of 37 CFR 1.97. Accordingly, the information disclosure statements are being considered by the examiner.

Election/Restrictions

Applicant's election without traverse of Group VIII, claims 50-55, in the reply filed on October 31, 2011, is acknowledged.

Claims 1-49 and 56-90 are hereby withdrawn from further consideration pursuant to 37 CFR 1.142(b) as being drawn to a nonelected inventions, there being no allowable generic or linking claim. Election was made **without** traverse in the reply filed on October 31, 2011.

Claim Rejections - 35 USC § 112

The following is a quotation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

Claims 50-55 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

Claim 50 recites the limitation "oil" in line 3. There is insufficient antecedent

basis for this limitation in the claim.

Claim 54 recites the limitation "said co-solvent" in line 4. There is insufficient

antecedent basis for this limitation in the claim. It is unclear that the recitation refers to

"a cosolvent" in first extraction step and second extraction. Also "a second extraction"

lacks antecedent basis for step b) because it does not refer to step b) in terms of

second extraction step. Also "An oil" as recited in claim 55, lacks antecedent basis.

Claim Rejections - 35 USC § 102

The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

A person shall be entitled to a patent unless –

(b) the invention was patented or described in a printed publication in this or a foreign country or in public use or on sale in this country, more than one year prior to the date of application for patent in the United States.

Claims 50-53 and 55 are rejected under 35 U.S.C. 102(b) as being clearly

anticipated by Patent Abstract of Japan 04-057853, dated Feb. 25, 1992, cited on

enclosed PTO-1449 Form.

Claims drawn to method for producing oil and an oil produced thereby.

Abstract 04-057853 teaches method for extracting krill oil comprising a) providing

krill meal; and extracting oil from the krill meal (powdered form of krill parts). The meal

(powdered form of krill parts) can be provided from heat-treated krill parts and is

storable. The extracting is carried out by supercritical extraction. An oil is produced by

the method.

The claims are identical to the abstract as discussed above and are considered to be clearly anticipated by the teachings therein. Krill shells are part of krill and oil is obtained from the krill parts. The krill parts are dried and hence subjected to heating to

provide for the krill meal which is subjected to supercritical extraction in two steps to

obtain the oil.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all

obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

The factual inquiries set forth in Graham v. John Deere Co., 383 U.S. 1, 148

USPQ 459 (1966), that are applied for establishing a background for determining

obviousness under 35 U.S.C. 103(a) are summarized as follows:

- 1. Determining the scope and contents of the prior art.
- 2. Ascertaining the differences between the prior art and the claims at issue.
- 3. Resolving the level of ordinary skill in the pertinent art.
- 4. Considering objective evidence present in the application indicating obviousness or nonobviousness.

This application currently names joint inventors. In considering patentability of

the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of

the various claims was commonly owned at the time any inventions covered therein

were made absent any evidence to the contrary. Applicant is advised of the obligation

under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was

not commonly owned at the time a later invention was made in order for the examiner to

consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g)

prior art under 35 U.S.C. 103(a).

Claim 54 is rejected under 35 U.S.C. 103(a) as being unpatentable over JP as cited and discussed above, in view of Kamiya et al (US 20060193962A1), cited on enclosed PTO-892 Form.

Claims are discussed above as if the JP abstract.

Kamiya et al, US 20060193962A1, teach extraction with supercritical fluid and solvent [0043], and the solvent can be a monohydric alcohol [0049], ranging from 1 to 20% [0059].

Claim differs from JP in that monohydric alcohol is not disclosed.

It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to carry out supercritical extraction of JP using a co-solvent monohydric alcohol as disclosed by Kamiya et al to produce oil from krill.

Each of the claim feature are disclosed and one of skill would have been motivated to carry out the process steps to provide oil with the expectation of successful results. Clearly the claim is prima facie obvious over the cited prior art.

All claims fail to be patentably distinguishable over the state of the art discussed above and cited on the enclosed PTO-892 and/or PTO-1449. Therefore, the claims are properly rejected.

The remaining references listed on the enclosed PTO-892 and/or PTO-1449 are cited to further show the state of the art.

No claims are allowed.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to DEBBIE K. WARE whose telephone number is (571)272-0924. The examiner can normally be reached on 9:30-6:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Mike Wityshyn can be reached on 571-272-0926. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

> /Deborah K. Ware/ Deborah K. Ware Primary Examiner Art Unit 1651

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12/057,775 | Applicant(s)/Pater
Reexamination
BRUHEIM ET AL. | | |
|----------------------------|---------------------------------------|---|-------------|--|
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| | DEBBIE K. WARE | 1651 | Page 1 of 1 | |
| | | | | |

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Fealey, Terence, Marietta, GA, UNITED STATES Bailes, Julian E., Morgantown, WV, UNITED STATES ΡT US 20110257267 A1 20111020 ΑT US 2010-904049 A1 20101013 (12) PRAI US 2009-251230P 20091013 (61) DT Utility FS APPLICATION LN.CNT 2397 INCL INCLM: 514/547.000 INCLS: 514/560.000; 514/549.000 514/547.000 NCL NCLM: NCLS: 514/549.000; 514/560.000 IPC IPCI A61K0031-232 [I,A]; A61P0025-00 [I,A]; A61K0031-202 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 3 OF 27 USPATFULL on STN L4AN 2011:251469 USPATFULL SOLVENT-FREE PROCESS FOR OBTAINING PHOSPHOLIPIDS AND NEUTRAL ENRICHED ТΤ KRILL OILS ΙN Sclabos Katevas, Dimitri, Santiage, CHILE Toro Guerra, Raul R., Santiage, CHILE Chiong Lay, Mario M., Santiage, CHILE PA THAROS LTD., Santiago, CHILE (non-U.S. corporation) LONZA LTD., Basel, SWITZERLAND (non-U.S. corporation) ΡI US 20110224450 A1 20110915 AI US 2011-96644 A1 20110428 (13) RLI Continuation-in-part of Ser. No. WO 2009-IB7269, filed on 30 Oct 2009, PENDING Utility DT APPLICATION FS LN.CNT 2021 INCLM: 554/023.000 INCL INCLS: 554/008.000; 554/078.000 NCL NCLM: 554/023.000 554/008.000; 554/078.000 NCLS: IPC IPCI C11B0001-00 [I,A]; C07F0009-10 [I,A] IPCR C11B0001-00 [I,A]; C07F0009-10 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 4 OF 27 USPATFULL on STN T.4 AN 2011:212256 USPATFULL ΤI METHOD FOR PRODUCING LIPIDS ΙN Yoshikawa, Kazuhiro, Tokyo, JAPAN Mikajiri, Akihiro, Tokyo, JAPAN ΡA NIPPON SUISAN KAISHA, LTD., Tokyo, JAPAN (non-U.S. corporation) US 20110189760 ΡT A1 20110804 US 2009-120842 AI 20090924 (13) A1 WO 2009-JP66530 20090924 PCT 371 date 20110425 JP 2008-248986 PRAI 20080926 DT Utility FS APPLICATION LN.CNT 1345 INCLM: 435/271.000 INCL INCLS: 554/020.000 NCL NCLM: 435/271.000 NCLS: 554/020.000 IPCI IPC C11C0001-00 [I,A]; C11B0001-00 [I,A] C11C0001-00 [I,A]; C11B0001-00 [I,A] IPCR CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 5 OF 27 USPATFULL on STN T.4

2011:211870 USPATFULL ΑN ТΤ METHOD FOR CONCENTRATING LIPIDS ΤN Yoshikawa, Kazuhiro, Tokyo, JAPAN PA NIPPON SUISAN KAISHA, LTD., Tokyo, JAPAN (non-U.S. corporation) ΡT US 20110189374 A1 20110804 ΑI US 2009-120875 A1 20090924 (13) WO 2009-JP66529 20090924 20110425 PCT 371 date PRAI JP 2008-248986 20080926 DT Utility APPLICATION FS LN.CNT 961 INCL INCLM: 426/601.000 INCLS: 554/008.000 NCL 426/601.000 NCLM: NCLS: 554/008.000 IPCI TPC A23D0009-00 [I,A]; C11B0001-06 [I,A] IPCR A23D0009-00 [I,A]; C11B0001-06 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. L4ANSWER 6 OF 27 USPATFULL on STN AN 2011:198158 USPATFULL ΤI METHODS OF TREATING AND PREVENTING NEUROLOGICAL DISORDERS USING DOCOSAHEXAENOIC ACID ΙN AISEN, Paul S., Solana Beach, CA, UNITED STATES Quinn, Joseph F., Portland, OR, UNITED STATES Yurko-Mauro, Karin, Silver Spring, MD, UNITED STATES ΡA MARTEK BIOSCIENCES CORPORATION, Columbia, MD, UNITED STATES (U.S. corporation) A1 20110721 ΡT US 20110177061 A1 20100709 (12) US 2010-833913 ΑT US 2009-224836P PRAI 20090710 (61) US 2010-359792P 20100629 (61) DT Utility FS APPLICATION LN.CNT 2653 INCL INCLM: 424/133.100 INCLS: 514/560.000; 514/120.000; 514/547.000; 514/549.000; 514/297.000; 514/319.000; 514/479.000; 514/215.000; 424/184.100; 424/172.100; 424/152.100; 514/458.000 NCL NCLM: 424/133.100 NCLS: 424/152.100; 424/172.100; 424/184.100; 514/120.000; 514/215.000; 514/297.000; 514/319.000; 514/458.000; 514/479.000; 514/547.000; 514/549.000; 514/560.000 A61K0031-202 [I,A]; A61K0031-661 [I,A]; A61K0031-232 [I,A]; TPC IPCI A61K0031-473 [I,A]; A61K0031-445 [I,A]; A61K0031-27 [I,A]; A61K0031-55 [I,A]; A61K0039-00 [I,A]; A61K0039-395 [I,A]; A61K0031-355 [I,A]; A61P0025-28 [I,A]; A61P0025-00 [I,A] A61K0031-202 [I,A]; A61K0031-232 [I,A]; A61K0031-27 [I,A]; IPCR A61K0031-355 [I,A]; A61K0031-445 [I,A]; A61K0031-473 [I,A]; A61K0031-55 [I,A]; A61K0031-661 [I,A]; A61K0039-00 [I,A]; A61K0039-395 [I,A]; A61P0025-00 [I,A]; A61P0025-28 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 7 OF 27 USPATFULL on STN T.4 2011:146375 USPATFULL AN ΤI KRILL OIL PROCESS IN Breivik, Harald, Porsgrunn, NORWAY Thorstad, Olav, Porsgrunn, NORWAY ΡA PRONOVA BIOPHARMA NORGE AS, Lysaker, NORWAY (non-U.S. corporation) РT US 20110130458 A1 20110602 ΑI US 2009-992365 A1 20090515 (12)

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WO 2009-NO184 20090515 20110211 PCT 371 date 20080515 (61) PRAT US 2008-53455P Utility DT APPLICATION FS LN.CNT 688 INCL INCLM: 514/560.000 INCLS: 426/608.000; 426/417.000 NCL 514/560.000 NCLM: 426/417.000; 426/608.000 NCLS: A61K0031-202 [I,A]; A61P0003-06 [I,A]; A61P0003-00 [I,A]; IPC IPCI A61P0009-00 [I,A]; A61P0009-04 [I,A]; A61P0009-10 [I,A]; A23D0007-00 [I,A]; A23D0009-00 [I,A] TPCR A61K0031-202 [I,A]; A23D0007-00 [I,A]; A23D0009-00 [I,A]; A61P0003-00 [I,A]; A61P0003-06 [I,A]; A61P0009-00 [I,A]; A61P0009-04 [I,A]; A61P0009-10 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 8 OF 27 USPATFULL on STN T.4 2011:117434 USPATFULL AN ТΤ POWDERED COMPOSITION CONTAINING OIL-SOLUBLE COMPONENT, FUNCTIONAL FOOD USING THE SAME, AND PACKAGED PRODUCT THEREOF ΙN Suzuki, Keiichi, Kanagawa, JAPAN Sasaki, Hidemi, Kanagawa, JAPAN Serizawa, Shinichiro, Kanagawa, JAPAN Arakawa, Jun, Kanagawa, JAPAN PA FUJIFILM CORPORATION, Minato-ku, Tokyo, JAPAN (non-U.S. corporation) ΡI US 20110104340 A1 20110505 ΑT US 2008-673977 A1 20080819 (12) WO 2008-JP65061 20080819 20100218 PCT 371 date PRAI JP 2007-213712 20070820 JP 2007-230582 20070905 DT Utility FS APPLICATION LN.CNT 2345 INCL INCLM: 426/096.000 INCLS: 426/654.000; 426/590.000 426/096.000 NCL NCLM: NCLS: 426/590.000; 426/654.000 IPC IPCI A21D0002-16 [I,A]; A23L0002-52 [I,A] IPCR A21D0002-16 [I,A]; A23L0002-52 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 9 OF 27 USPATFULL on STN T.4 AN 2011:117391 USPATFULL METHODS OF USING KRILL OIL TO TREAT RISK FACTORS FOR ТΤ CARDIOVASCULAR, METABOLIC, AND INFLAMMATORY DISORDERS BRUHEIM, Inge, Volda, NORWAY ΙN Tilseth, Snorre, Bergen, NORWAY Cohn, Jeffery, Sydney, AUSTRALIA Griinari, Mikko, Espoo, FINLAND Mancinelli, Daniele, Orsta, NORWAY Hoem, Nils, Oslo, NORWAY Vik, Hogne, Eiksmarka, NORWAY Banni, Sebastiano, Calgliari, ITALY Aker BioMarine A.S.A., Oslo, NORWAY (non-U.S. corporation) PA ΡI US 20110104297 A1 20110505 AI US 2010-790575 A1 20100528 (12) Continuation-in-part of Ser. No. US 2008-57775, filed on 28 Mar 2008, RLI PENDING PRAI US 2007-975058P 20070925 (60)

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US 2009-155758P 20090226 (61) PRAT Utility DT FS APPLICATION LN.CNT 3112 INCLM: 514 2 INCL NCL NCLM: 514/005.500 NCLS: 514/691.000 A61K0038-02 [I,A] IPC IPCI IPCR A61K0038-02 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 13 OF 27 USPATFULL on STN T.4 AN 2010:255355 USPATFULL ΤТ LOW VISCOSITY PHOSPHOLIPID COMPOSITIONS ΤN Tilseth, Snorre, Bergen, NORWAY PA AKER BIOMARINE ASA, Oslo, NORWAY (non-U.S. corporation) ΡT US 20100226977 A1 20100909 US 2010-711553 20100224 (12) AI A1 Continuation-in-part of Ser. No. US 2008-201325, filed on 29 Aug 2008, RLI PENDING PRAI US 2009-155767P 20090226 (61) US 2007-968765P 20070829 (60) Utility DT APPLICATION FS LN.CNT 2394 INCL INCLM: 424/456.000 INCLS: 426/601.000; 426/417.000; 514/078.000 NCL NCLM: 424/456.000 NCLS: 426/417.000; 426/601.000; 514/078.000 TPC IPCI A61K0031-685 [I,A]; A23D0009-00 [I,A]; A23D0009-02 [I,A]; A61K0009-48 [I,A]; A61P0009-00 [I,A]; A61P0019-00 [I,A]; A61P0029-00 [I,A] IPCR A61K0031-685 [I,A]; A23D0009-00 [I,A]; A23D0009-02 [I,A]; A61K0009-48 [I,A]; A61P0009-00 [I,A]; A61P0019-00 [I,A]; A61P0029-00 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 14 OF 27 USPATFULL on STN L4 2010:228249 USPATFULL AN METHODS FOR IMPROVING COGNITIVE FUNCTION AND DECREASING HEART RATE ТΤ ΙN YURKO-MAURO, Karin, Silver Spring, MD, UNITED STATES MARTEK BIOSCIENCES CORPORATION, Columbia, MD, UNITED STATES (U.S. PA corporation) ΡT US 20100203123 Α1 20100812 ΑT US 2010-699009 Α1 20100202 (12) US 2009-149310P 20090202 (61) PRAT US 2009-183548P 20090602 (61) DT Utility APPLICATION FS LN.CNT 2358 INCL INCLM: 424/456.000 INCLS: 514/560.000; 514/549.000; 514/458.000 NCL NCLM: 424/456.000 514/458.000; 514/549.000; 514/560.000 NCLS: TPC IPCI A61K0009-64 [I,A]; A61K0031-20 [I,A]; A61K0031-22 [I,A]; A61K0031-355 [I,A]; A61P0025-00 [I,A]; A61P0009-00 [I,A] IPCR A61K0009-64 [I,A]; A61K0031-20 [I,A]; A61K0031-22 [I,A]; A61K0031-355 [I,A]; A61P0009-00 [I,A]; A61P0025-00 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 15 OF 27 USPATFULL on STN T.4

ΤТ PROCESS FOR PRODUCTION OF OMEGA-3 RICH MARINE PHOSPHOLIPIDS FROM KRILL ΤN Breivik, Harald, Porsgrunn, NORWAY ΡТ US 20100143571 A1 20100610 ΑI US 2007-515098 A1 20071115 (12) WO 2007-NO402 20071115 20100217 PCT 371 date PRAI US 2006-859289P 20061116 (60) DT Utility FS APPLICATION LN.CNT 537 INCLM: 426/643.000 INCL INCLS: 426/417.000; 554/021.000; 568/366.000; 536/020.000 NCL 426/643.000 NCLM: NCLS: 426/417.000; 536/020.000; 554/021.000; 568/366.000 IPCI A23L0001-325 [I,A]; A23K0001-10 [I,A]; A23K0001-18 [I,A]; TPC C11B0001-10 [I,A]; C07C0045-78 [I,A]; C08B0037-08 [I,A] IPCR A23L0001-325 [I,A]; A23K0001-10 [I,A]; A23K0001-18 [I,A]; C07C0045-78 [I,A]; C08B0037-08 [I,A]; C11B0001-10 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. T.4 ANSWER 16 OF 27 USPATFULL on STN AN 2009:109974 USPATFULL ΤI Polyunsaturated Fatty Acid-Containing Solid Fat Compositions and Uses and Production Thereof ΙN Namal Senanayake, S.P. Janaka, Lexington, KY, UNITED STATES Ahmed, Naseer, Lexington, KY, UNITED STATES Fichtali, Jaouad, Lexington, KY, UNITED STATES PA Martek Biosciences Corporation, Columbia, MD, UNITED STATES (U.S. corporation) A1 20090416 ΡT US 20090099260 US 2008-201728 A1 20080829 (12) ΑI US 2007-969536P PRAI 20070831 (60) Utility DT APPLICATION FS LN.CNT 2660 INCL INCLM: 514/560.000 INCLS: 426/601.000; 426/072.000 514/560.000 NCL NCLM: 426/072.000; 426/601.000 NCLS: IPC TPCT A61K0031-20 [I,A]; A23D0007-005 [I,A]; A23L0001-30 [I,A] IPCR A61K0031-20 [I,A]; A23D0007-005 [I,A]; A23L0001-30 [I,A] CAS INDEXING IS AVAILABLE FOR THIS PATENT. T.4 ANSWER 17 OF 27 USPATFULL on STN AN 2009:67318 USPATFULL METHOD FOR MAKING KRILL MEAL ТΤ ΙN Tilseth, Snorre, Bergen, NORWAY Hostmark, Oistein, Loddefjord, NORWAY PA Aker BioMarine ASA, Oslo, NORWAY (non-U.S. corporation) ΡI US 20090061067 A1 20090305 AI US 2008-201325 A1 20080829 (12) PRAI US 2007-968765P 20070829 (60) DT Utility FS APPLICATION LN.CNT 2307 INCL INCLM: 426/602.000 INCLS: 426/417.000; 210/149.000; 426/480.000; 426/609.000; 426/648.000; 426/608.000; 366/145.000; 366/147.000 NCL NCLM: 426/602.000 210/149.000; 366/145.000; 366/147.000; 426/417.000; 426/480.000; NCLS: 426/608.000; 426/609.000; 426/648.000 IPC IPCI A23D0007-005 [I,A]; A23D0007-02 [I,A]; A23D0007-04 [I,A];