## UNITED STATES PATENT AND TRADEMARK OFFICE

# **CERTIFICATE OF CORRECTION**

PATENT NO. : 9,644,169 B2 Page 1 of 1

APPLICATION NO. : 15/180431 DATED : May 9, 2017

INVENTOR(S) : Inge Bruheim, Snorre Tilseth and Daniele Mancinelli

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

In the Claims

Column 36, Lines 31-32 should be deleted and replaced with:

a) obtaining a denatured krill product produced by treating freshly harvested krill to denature lipases and

Signed and Sealed this Twentieth Day of June, 2017

Joseph Matal

Doseph

Performing the Functions and Duties of the Under Secretary of Commerce for Intellectual Property and Director of the United States Patent and Trademark Office

## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent of: Inge Bruheim et al. Confirmation No.: 2763

Patent No.: 9,644,169
Application No.: 15/180,431
Issued: 09-May-2017

Entitled: BIOEFFECTIVE KRILL OIL COMPOSITIONS

# REQUEST FOR CERTIFICATE OF CORRECTION OF PATENT

Certificate of Correction Branch Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir or Madam:

Pursuant to 35 U.S.C. §255 and 37 C.F.R. §1.323, patentee respectfully request that the Director issue a Certificate of Correction in the above-referenced patent to correct a typographical error in claim 12.

Please correct the Letters Patent at Column 36, lines 31-32, as follows:

 a) obtaining a denatured krill product produced by treating freshly harvested krill krill to denature lipases and

The above-noted correction does not involve such changes in the patent as would constitute new matter or would require reexamination.

A completed Form PTO/SB/44 accompanies this request, with the above-noted correction printed thereon. Accordingly, a Certificate of Correction is believed proper and issuance thereof is respectfully requested.

The director is hereby authorized to charge the \$100 fee due with the filing of this Request to deposit account number 50-4302 referencing attorney docket number AKBM-14409/US-12/CON.

Patent No. 9,644,169 Request for Certificate of Correction

Respectfully submitted,

CASIMIR JONES, S.C.

Dated: May 15, 2017

/J. Mitchell Jones/
J. Mitchell Jones
Reg. No. 44,174
2275 Deming Way, Suite 310
Middleton, WI 53562
608 662 1277

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(Also Form PTO-1050)

# UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

	Page <u>1</u> of <u>1</u>
PATENT NO. : 9,644,169	
APPLICATION NO.: 15/180,431	
ISSUE DATE : 09-May-2017	
INVENTOR(S) : Inge Bruheim, Snorre Tilseth, Daniele Mancinelli	
It is certified that an error appears or errors appear in the above-identified patent and t is hereby corrected as shown below:	hat said Letters Patent
In the claims, column 36, lines 31-32 should be deleted and replaced with:	
a) obtaining a denatured krill product produced by treat- ing freshly harvested krill to denature lipases and	
	ļ

MAILING ADDRESS OF SENDER (Please do not use customer number below):

Casimir Jones SC 2275 Deming Way, Suite 310 Middleton, WI 53562

This collection of information is required by 37 CFR 1.322, 1.323, and 1.324. 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 1.0 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: Attention Certificate of Corrections Branch, Commissioner for Patents, 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.

## 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:

- 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.
- 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.
- 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 inspection or an issued patent.
- 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:	15	180431						
Filing Date:	13-	Jun-2016						
Title of Invention:	BIC	DEFFECTIVE KRILL O	IL COMPOSITIC	DNS				
First Named Inventor/Applicant Name:	Ing	je Bruheim						
Filer:	John Mitchell Jones							
Attorney Docket Number:	AKBM-14409/US-12/CON							
Filed as Large Entity								
Filing Fees for Utility under 35 USC 111(a)								
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:								
Certificate of correction		1811	1	100	100			

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Extension-of-Time:				
Miscellaneous:				
	Tot	al in USD	(\$)	100

Electronic Ack	knowledgement Receipt
EFS ID:	29205478
Application Number:	15180431
International Application Number:	
Confirmation Number:	2763
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS
First Named Inventor/Applicant Name:	Inge Bruheim
Customer Number:	72960
Filer:	John Mitchell Jones/Mallory Checkett
Filer Authorized By:	John Mitchell Jones
Attorney Docket Number:	AKBM-14409/US-12/CON
Receipt Date:	15-MAY-2017
Filing Date:	13-JUN-2016
Time Stamp:	15:26:01
Application Type:	Utility under 35 USC 111(a)

# **Payment information:**

Submitted with Payment	yes
Payment Type	DA
Payment was successfully received in RAM	\$100
RAM confirmation Number	051617INTEFSW00002232504302
Deposit Account	
Authorized User	

The Director of the USPTO is hereby authorized to charge indicated fees and credit any overpayment as follows:

File Listing	<b>j</b> :				
Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
			78110		
1	Request for Certificate of Correction	14409US12CON_RequestforCer tCorrection.pdf	fc60e456ce73ef1082aac788aac87e0b3b71 2def	no	2
Warnings:					
Information:					
			164787		
2	Request for Certificate of Correction	14409US12CON_CertCorrectio nForm.pdf	a788c9542fee0e4991d3b56f571aed1a0c9e 3b77	no	2
Warnings:					
Information:					
			30251		
3	Fee Worksheet (SB06)	fee-info.pdf	16f13bb5916a91fb64292b9140c6f467f0e7f 31e	no	2
Warnings:					
Information:					
		Total Files Size (in bytes)	2	73148	

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

#### 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.



UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450

P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	ISSUE DATE	PATENT NO.	ATTORNEY DOCKET NO.	CONFIRMATION NO.
15/180,431	05/09/2017	9644169	AKBM-14409/US-12/CON	2763

72960 7596

04/19/2017

Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON, WI 53562

## **ISSUE NOTIFICATION**

The projected patent number and issue date are specified above.

# **Determination of Patent Term Adjustment under 35 U.S.C. 154 (b)**

(application filed on or after May 29, 2000)

The Patent Term Adjustment is 0 day(s). Any patent to issue from the above-identified application will include an indication of the adjustment on the front page.

If a Continued Prosecution Application (CPA) was filed in the above-identified application, the filing date that determines Patent Term Adjustment is the filing date of the most recent CPA.

Applicant will be able to obtain more detailed information by accessing the Patent Application Information Retrieval (PAIR) WEB site (http://pair.uspto.gov).

Any questions regarding the Patent Term Extension or Adjustment determination should be directed to the Office of Patent Legal Administration at (571)-272-7702. Questions relating to issue and publication fee payments should be directed to the Application Assistance Unit (AAU) of the Office of Data Management (ODM) at (571)-272-4200.

APPLICANT(s) (Please see PAIR WEB site http://pair.uspto.gov for additional applicants):

Inge Bruheim, Volda, NORWAY; AKER BIOMARINE ANTARCTIC AS, Stamsund, NORWAY; Snorre Tilseth, Bergen, NORWAY; Daniele Mancinelli, Orsta, NORWAY;

The United States represents the largest, most dynamic marketplace in the world and is an unparalleled location for business investment, innovation, and commercialization of new technologies. The USA offers tremendous resources and advantages for those who invest and manufacture goods here. Through SelectUSA, our nation works to encourage and facilitate business investment. To learn more about why the USA is the best country in the world to develop technology, manufacture products, and grow your business, visit <u>SelectUSA.gov</u>.

# INFORMATION DISCLOSURE STATEMENT BY APPLICANT

( Not for submission under 37 CFR 1.99)

Application Number		15180431 - GAU: 1672
Filing Date		2016-06-13
First Named Inventor	Inge E	Bruheim
Art Unit		
Examiner Name		
Attorney Docket Number		AKBM-14409/US-12/CON

	11	20140107072	2014-04	2014-04-17 S		et al.				
hange(s) a	12	20090061067	2009-03	2009-03-05		Snorre Tilseth et al.				
to document Q.N./ 2/31/2016	13	20140010888	2014-0	1-09	Bruheim et AKER BIOMAI AS	al. RINE ANTAR <del>CTIC</del>				
If you wis	h to ac	dd additional U.S. Pub	lished Application	n citation	n information p	lease click the Add	d buttor	ı. Add		
-			FOREIG	GN PAT	ENT DOCUM	ENTS		Remove		
Examiner Initial*	Cite No	Foreign Document Number <sup>3</sup>	Country Code <sup>2</sup> i	Kind Code <sup>4</sup>	Publication Date	Name of Patentee Applicant of cited Document	; 	where Rel	or Relevant	T5
	1									
If you wis	h to ac	ld additional Foreign F	Patent Document	citation	information pl	ease click the Add	button	Add		
			NON-PATE	NT LITE	RATURE DO	CUMENTS		Remove		
Examiner Initials*	Cite No	Include name of the a (book, magazine, jou publisher, city and/or	rnal, serial, symp	osium,	catalog, etc), c					T5
	1	Takahashi et al., Predid High Performance Liqui							erolipid on	
	2		anaka, Biosynthesis of 1,2-dieicosapentaenoyl-sn-glycero-3-phosphocholine in Caenorhabditis elegans, Eur. J. Biochem. 263, 189±194 (1999)							
	3	Tocher, Chapter 6, Glyo and Mommsen (eds.)(1		etabolis	m, Biochemistry	and molecular biolog	gy of fisl	hes, vol. 4,	Hochachka	

ALL REFERENCES CONSIDERED EXCEPT WHERE LINED THROUGH. /Y.K.C/

# INFORMATION DISCLOSURE STATEMENT BY APPLICANT

( Not for submission under 37 CFR 1.99)

Application Number		15180431 - GAU: 1672
Filing Date		2016-06-13
First Named Inventor	Inge E	Bruheim
Art Unit		
Examiner Name		
Attorney Docket Number		AKBM-14409/US-12/CON

	xaminer iitial*	Cite No	Publication Number	Kind Code <sup>1</sup>	Publication Date	Name of Patentee or Applicant of cited Document	Pages,Columns,Lines where Relevant Passages or Relevant Figures Appear
	nge(s) a ocument		20140274968		2014-09-18	Berge et al.  AKER BIOMARINE ANTARCTIC  AS	
(Q.  2/3	N./ 51/2016	<sup>5</sup> 2	20150030718		2015-01-29	Saebo AKER BIOMARINE ANTARCTIC AS	
		3	20140088043		2014-03-27	Hoem et al.  AKER BIOMARINE ANTARCTIC  AS	
		4	20140088047		2014-03-27	Hoem et al.  AKER BIOMARINE ANTARCTIC  AS	
		5	20140080791		2014-03-20	Berge et al.  AKER BIOMARINE ANTARCTIC  AS	
		6	20150164841		2015-06-18	Hoem et al.  AKER BIOMARINE ANTARCTIC  AS	
		7	20140363517		2014-12-11	Bruheim et al.  AKER BIOMARINE ANTARCTIC  AS	
		8	20100226977		2010-09-09	Snorre Tilseth et al.	
		9	20150050403		2015-02-19	Tilseth et al.  AKER BIOMARINE ANTARCTIC  AS	
		10	20140005421		2014-01-02	Bruheim et al.  AKER BIOMARINE ANTARCTIC  AS	

Doc code: IDS Doc description: Information Disclosure Statement (IDS) Filed

PTO/SB/08a (03-15) Approved for use through 07/31/2016. OMB 0651-0031 U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

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INFORMATION DISCLOSURE	Application Number			
	Filing Date		2016-06-13	
	First Named Inventor Inge Br		Bruheim	
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit			
(Not for Submission under or STR 1.55)	Examiner Name			
	Attorney Docket Number	er	AKBM-14409/US-12/CON	

	U.S.PATENTS Remove					
Examiner Initial*	No	Patent Number	Kind Code <sup>1</sup>	Issue Date	Name of Patentee or Applicant of cited Document	Pages,Columns,Lines where Relevant Passages or Relevant Figures Appear
Change(s) a to documen Q.N./ 12/31/201	t, 1	9119864		2015-09-01	Bruheim, et al.  AKER BIOMARINE ANTARCTIC  AS	
	2	9072752		2015-07-07	Bruheim, et al.  AKER BIOMARINE ANTARCTIC  AS	
	3	9034388		2015-05-19	Inge Bruheim et al.	
	4	9028877		2015-05-12	Bruheim, et al.  AKER BIOMARINE ANTARCTIC  AS	
	5	9078905		2015-07-14	Bruheim, et al.  AKER BIOMARINE ANTARCTIC  AS	
	6	B372812		2013-02-12	Snorre Tilseth et al.	
	7	8697138		2014-04-15	Inge Bruheim et al.	
If you wis	h to add	additional U.S. Paten	t citatio	n information pl	ease click the Add button.	Add
U.S.PATENT APPLICATION PUBLICATIONS Remove						

ALL REFERENCES CONSIDERED EXCEPT WHERE LINED THROUGH. /Y.K.C/

Doc code: IDS Doc description: Information Disclosure Statement (IDS) Filed

PTO/SB/08a (03-15) Approved for use through 07/31/2016. OMB 0651-0031 U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

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.

	Application Number		
INFORMATION PIGGI COURT	Filing Date		2016-06-13
INFORMATION DISCLOSURE	First Named Inventor	Inge E	Bruheim
STATEMENT BY APPLICANT ( Not for submission under 37 CFR 1.99)	Art Unit		
(Not for Submission under or of K 1.33)	Examiner Name		
	Attorney Docket Number		AKBM-14409/US-12/CON

	U.S.PATENTS						Remove
Examiner Initial*	Cite No	Patent Number	Kind Code <sup>1</sup>	Issue Date	ate of sited Document		Columns,Lines where nt Passages or Relevant Appear
	1	4119619		1978-10-10 ROGOZHIN SERGEI VASILIEVICH et al.			
hange(s) a	2 pplied	5434183		07/1995 <del>1997-07-18</del>			
Y.Y.S./ /5/2017	3	6537787		2003-03-25	BRETON		
	4	5800299		2004-10-05	BEAUDOIN & MARTIN		
If you wis	h to add	additional U.S. Pater	nt citatio	n information p	ease click the Add button.		Add
			U.S.P	ATENT APPLI	CATION PUBLICATIONS		Remove
Examiner Initial*	Cite No	Publication Number	Kind Code <sup>1</sup>	Publication Date	Name of Patentee or Applicant of cited Document	Relevar	Columns,Lines where nt Passages or Relevant Appear
	1	20030044495		2003-03-06	KAGAN and BRAUN		
	2	20040241249		2004-12-02	SAMPALIS		
If you wis	h to add	additional U.S. Publi	shed Ap	plication citatio	n information please click the Add	d button.	Add
	FOREIGN PATENT DOCUMENTS Remove						

# **INFORMATION DISCLOSURE STATEMENT BY APPLICANT**

( Not for submission under 37 CFR 1.99)

Application Number		15180431 - GAU: 1672
Filing Date		2016-06-13
First Named Inventor Inge E		Bruheim
Art Unit		
Examiner Name		
Attorney Docket Number		AKBM-14409/US-12/CON

Change(s) to documen	applied	2004-100943	wo	1 1/2004 <del>2006 05 31</del>	BTG INTERNATIONAL LIMITED
/Y.Y.S./ 1/5/2017		2005-018632	wo	2005-03-03	BTG INTERNATIONAL LIMITED
	23	2006/030552	wo	2006-03-23	SUNTORY LIMITED
	24	1706106	EP	2009-07-15	BRUZZESE
	25	1689413	EP	2006-08-16	ENZYMOTEC LTD
	26	2004-534800	JP	2004-11-18	Kohyo
	27	1660071	EP	2007-01-17	NIPPON SUISAN KAISHA, LTD.
	28	2002/083122	wo	2002-10-24	YEDA RESEARCH AND DEVELOPMENT CO. LTD
	29	05/004593	wo	2005-01-20	Norway Seafoods AS
	30	08/117062	wo	2008-10-02	Aker Biomarine ASA
	31	02/102394	wo	2002-12-27	Neptune Technologies & Bioress

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
15/180,431	06/13/2016	Inge Bruheim	AKBM-14409/US-12/CON	2763
72960 Casimir Jones, S	7590 03/28/201 S.C.	7	EXAM	IINER
	WAY, SUITE 310		CUTLIFF, YA	TE KAI RENE
			ART UNIT	PAPER NUMBER
			1672	
			NOTIFICATION DATE	DELIVERY MODE
			03/28/2017	ELECTRONIC

# 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.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

docketing@casimirjones.com pto.correspondence@casimirjones.com

	Application No.	Applicant(s)
	15/180,431	BRUHEIM ET AL.
Response to Rule 312 Communication	Examiner	Art Unit
	YATE' K. CUTLIFF	1672
The MAILING DATE of this communication ap	ppears on the cover sheet wit	h the correspondence address –
<ol> <li>The amendment filed on <u>15 March 2017</u> under 37 CFR 1</li> <li>a) ☑ entered.</li> </ol>	.312 has been considered, and	I has been:
b)  entered as directed to matters of form not affecting	the scope of the invention.	
c) disapproved because the amendment was filed after  Any amendment filed after the date the issue fee  the required fee to withdraw the application from	e is paid must be accompanied	
d) disapproved. See explanation below.		
e)   entered in part. See explanation below.		
The amendment to claims 8 and 18 embody merely the coof the claims.		
	/YATE' K. CUTLIFF Primary Examiner, A	

U.S. Patent and Trademark Office PTOL-271 (Rev. 04-01) OK TO ENTER: /Y.K.C/ 03/21/2017

#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:Inge Bruheim et al.Confirmation No.:2763Serial No.:15/180,431Group No.:1672Filed:13-Jun-2016Examiner:CUTLIFF

Entitled: BIOEFFECTIVE KRILL OIL COMPOSITIONS

# AMENDMENT AFTER NOTICE OF ALLOWANCE UNDER 37 C.F.R. § 1.312

Commissioner of Patents Mail Stop - Amendment P.O. Box 1450 Arlington, VA 22313-1450

### Examiner Cutliff:

Applicants respectfully request entry of this amendment under 37 C.F.R. §1.312.

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. **AKBM-14409/US-12/CON**. This paragraph is intended to be a CONSTRUCTIVE PETITION FOR EXTENSION OF TIME in accordance with 37 C.F.R. § 1.136(a)(3).

Amendments to the Claims begin at page 2.

**Remarks** begin at page 5.



# UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
15/180,431 06/13/2016		Inge Bruheim	AKBM-14409/US-12/CON 2763		
7:	590 03/22/2017		EXAM	NER	
Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310			CUTLIFF, YAT	E KAI RENE	
MIDDLETON, W			ART UNIT	PAPER NUMBER	
			1672		
			NOTIFICATION DATE	DELIVERY MODE	
			03/22/2017	ELECTRONIC	

## NOTICE OF NON-COMPLIANT INFORMATION DISCLOSURE STATEMENT

An Information Disclosure Statement (IDS) filed <u>03.17.17</u> in the above-identified application fails to meet the requirements of 37 CFR 1.97(d) for the reason(s) specified below. Accordingly, the IDS will be placed in the file, but the information referred to therein has not been considered.

The IDS is not compliant with 37 CFR 1.97(d) because:

- The IDS lacks a statement as specified in 37 CFR 1.97(e).
- ☐ The IDS lacks the fee set forth in 37 CFR 1.17(p).
- ☐ The IDS was filed after the issue fee was paid. Applicant may wish to consider filing a petition to withdraw the application from issue under 37 CFR 1.313(c) to have the IDS considered. See MPEP 1308.

571-272-4200 or 1-888-786-0101
Application Assistance Unit
Office of Data Management

#### PART B - FEE(S) TRANSMITTAL

## Complete and send this form, together with applicable fee(s), to: Mail Mail Stop ISSUE FEE

Commissioner for Patents P.O. Box 1450

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INSTRUCTIONS: This form should be used for transmitting the ISSUE FEE and PUBLICATION FEE (if required). Blocks 1 through 5 should be completed where appropriate. All further correspondence including the Patent, advance orders and notification of maintenance fees will be mailed to the current correspondence address as indicated unless corrected below or directed otherwise in Block 1, by (a) specifying a new correspondence address; and/or (b) indicating a separate "FEE ADDRESS" for

maintenance fee notifications. Note: A certificate of mailing can only be used for domestic mailings of the Fee(s) Transmittal. This certificate cannot be used for any other accompanying papers. Each additional paper, such as an assignment or formal drawing, must have its own certificate of mailing or transmission. CURRENT CORRESPONDENCE ADDRESS (Note: Use Block 1 for any change of address) Certificate of Mailing or Transmission I hereby certify that this Fee(s) Transmittal is being deposited with the United States Postal Service with sufficient postage for first class mail in an envelope addressed to the Mail Stop ISSUE FEE address above, or being facsimile transmitted to the USPTO (571) 273-2885, on the date indicated below. 72960 12/21/2016 Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON, WI 53562 (Depositor's name (Signature (Date APPLICATION NO. FILING DATE FIRST NAMED INVENTOR ATTORNEY DOCKET NO. CONFIRMATION NO. 15/180.431 06/13/2016 Inge Bruheim AKBM-14409/US-12/CON 2763 TITLE OF INVENTION: BIOEFFECTIVE KRILL OIL COMPOSITIONS APPLN. TYPE ISSUE FEE DUE PUBLICATION FEE DUE PREV. PAID ISSUE FEE **ENTITY STATUS** TOTAL FEE(S) DUE DATE DUE UNDISCOUNTED \$0 03/21/2017 \$960 \$0 \$960 nonprovisional **EXAMINER** ART UNIT CLASS-SUBCLASS CUTLIFF, YATE KAI RENE 1672 554-210000 1. Change of correspondence address or indication of "Fee Address" (37 CFR 1.363). 2. For printing on the patent front page, list Casimir Jones, S.C. (1) The names of up to 3 registered patent attorneys ☐ Change of correspondence address (or Change of Correspondence Address form PTO/SB/122) attached. or agents OR, alternatively, (2) The name of a single firm (having as a member a registered attorney or agent) and the names of up to 2 registered patent attorneys or agents. If no name is listed, no name will be printed. ☐ "Fee Address" indication (or "Fee Address" Indication form PTO/SB/47; Rev 03-02 or more recent) attached. Use of a Customer Number is required. 3. ASSIGNEE NAME AND RESIDENCE DATA TO BE PRINTED ON THE PATENT (print or type) PLEASE NOTE: Unless an assignee is identified below, no assignee data will appear on the patent. If an assignee is identified below, the document has been filed for recordation as set forth in 37 CFR 3.11. Completion of this form is NOT a substitute for filing an assignment. (B) RESIDENCE: (CITY and STATE OR COUNTRY) (A) NAME OF ASSIGNEE AKER BIOMARINE ANTARCTIC AS STAMSUND, NORWAY Please check the appropriate assignee category or categories (will not be printed on the patent): 🔲 Individual 📮 Corporation or other private group entity 🖵 Government 4b. Payment of Fee(s): (Please first reapply any previously paid issue fee shown above) 4a. The following fee(s) are submitted: Issue Fee A check is enclosed. Publication Fee (No small entity discount permitted) Payment by credit card. Form PTO-2038 is attached. Advance Order - # of Copies \_ The director is hereby authorized to charge the required fee(s), any deficiency, or credits any overpayment, to Deposit Account Number \_504302\_ 5. Change in Entity Status (from status indicated above) NOTE: Absent a valid certification of Micro Entity Status (see forms PTO/SB/15A and 15B), issue fee payment in the micro entity amount will not be accepted at the risk of application abandonment. Applicant certifying micro entity status. See 37 CFR 1.29 Applicant asserting small entity status. See 37 CFR 1.27  $\underline{NOTE}$ : If the application was previously under micro entity status, checking this box will be taken to be a notification of loss of entitlement to micro entity status. <u>NOTE:</u> Checking this box will be taken to be a notification of loss of entitlement to small or micro entity status, as applicable. ☐ Applicant changing to regular undiscounted fee status. NOTE: This form must be signed in accordance with 37 CFR 1.31 and 1.33. See 37 CFR 1.4 for signature requirements and certifications. Authorized Signature /J. Mitchell Jones/ March 21, 2017 Typed or printed name \_ J. Mitchell Jones 44,174 Registration No.

#### Page 2 of 3 **RIMFROST EXHIBIT 1111** Page 0020

Electronic Patent Application Fee Transmittal					
Application Number:	15	180431			
Filing Date:	13-	Jun-2016			
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS				
First Named Inventor/Applicant Name:	Inge Bruheim				
Filer:	John Mitchell Jones				
Attorney Docket Number:	AK	BM-14409/US-12/C	ON		
Filed as Large Entity					
Filing Fees for Utility under 35 USC 111(a)					
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:					
UTILITY APPL ISSUE FEE		1501	1	960	960

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Extension-of-Time:				
Miscellaneous:				
	Tot	al in USD	(\$)	960

Electronic Ack	Electronic Acknowledgement Receipt				
EFS ID:	28679646				
Application Number:	15180431				
International Application Number:					
Confirmation Number:	2763				
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS				
First Named Inventor/Applicant Name:	Inge Bruheim				
Customer Number:	72960				
Filer:	John Mitchell Jones/Mallory Checkett				
Filer Authorized By:	John Mitchell Jones				
Attorney Docket Number:	AKBM-14409/US-12/CON				
Receipt Date:	21-MAR-2017				
Filing Date:	13-JUN-2016				
Time Stamp:	10:00:00				
Application Type:	Utility under 35 USC 111(a)				

# **Payment information:**

Submitted with Payment	yes
Payment Type	DA
Payment was successfully received in RAM	\$960
RAM confirmation Number	032117INTEFSW00010749504302
Deposit Account	
Authorized User	

The Director of the USPTO is hereby authorized to charge indicated fees and credit any overpayment as follows:

File Listing:					
Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1	Issue Fee Payment (PTO-85B)		98510		1
		14409US12CON_IssueFeeTrans mittal.pdf	9a406fe698e55f0c48a6d04cc4da8698316e 4d2b	no	
Warnings:		-		l	
Information:					
			30484		
2 Fee Worksheet (SB06) fee-info.p		fee-info.pdf	61f157c91240ecefa1876097d6ed2e3de833 bd2a	no	2
Warnings:		-			
Information:					

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Total Files Size (in bytes):

#### 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.

128994

Electronic Acknowledgement Receipt				
EFS ID:	28665295			
Application Number:	15180431			
International Application Number:				
Confirmation Number:	2763			
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS			
First Named Inventor/Applicant Name:	Inge Bruheim			
Customer Number:	72960			
Filer:	John Mitchell Jones/Mallory Checkett			
Filer Authorized By:	John Mitchell Jones			
Attorney Docket Number:	AKBM-14409/US-12/CON			
Receipt Date:	17-MAR-2017			
Filing Date:	13-JUN-2016			
Time Stamp:	17:24:22			
Application Type:	Utility under 35 USC 111(a)			

# **Payment information:**

Submitted with Payment	no
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# File Listing:

Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
		IDDD047 00745 D. W. C. L.	1904523		
1	Other Reference-Patent/App/Search documents	IPR2017-00745_PetitionforInter PartesReview.pdf	e73e2b349d301fba71293a509c8e9400a7f b8c4a	no	82

Warnings: RIMFROST EXHIBIT 1111 Page 0025

Information:					
			1807020		
2	Other Reference-Patent/App/Search documents	IPR2017-00747_PetitionforInter PartesReview.pdf	8019fb462d1e065c45b8788de85f108f7652 8654	no	80
Warnings:					
Information:					
			2892891		
3	Other Reference-Patent/App/Search documents	2017-00746_877_IPR.pdf	0138d0266c9f0f0ba94b34e3753fa09a0c62 c02c	no	93
Warnings:					
Information:					
			4355272		
4	Other Reference-Patent/App/Search documents	2017-00748_877_IPR.pdf	no 65b11f71e2b1f111ace884443e1fcb9a13cc Sa6d		94
Warnings:			1		
Information:					
		ITC_337-	352309	no	
5	Other Reference-Patent/App/Search documents	TA-1019RespondentsNoticeofP riorArt_02_01_2017_uspto.pdf	2469901b244d42b3587a63a591631030b2 5bf1dd		49
Warnings:		-	1	·	
Information:					
			92621		
6	Other Reference-Patent/App/Search documents	AU2014256345_14409AU13_N oticeofOppositionFiledRimfrost .pdf		no	3
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Information:					
		AU2014256345_14409AU13_N	99805		
7	Other Reference-Patent/App/Search documents	oticeofOppositionFiledEnzymo tec.pdf		no	3
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Information:					
			308855		
8 Other Reference-Patent/App/Search documents ITC_337- TA-1019MotionforLeavetoAm nd_uspto.pdf		f466150cc0196cfdd1b840f454c679952677 2c7a	no	66	
Warnings:					
Information:					

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#### New Applications Under 35 U.S.C. 111

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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.

Doc code: IDS Doc description: Information Disclosure Statement (IDS) Filed PTO/SB/08a (03-15)
Approved for use through 07/31/2016. OMB 0651-0031
U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

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.

	Application Number		15180431	
	Filing Date		2016-06-13	
INFORMATION DISCLOSURE	First Named Inventor Inge Br		Bruheim	
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit		1672	
(Not for Submission ander or of K 1.00)	Examiner Name	CUTL	IFF, YATE KAI RENE	
	Attorney Docket Number	er	AKBM-14409/US-12/CON	

	U.S.PATENTS Remove									
Examiner Initial*	Cite No	Patent Number	Kind Code <sup>1</sup>	Issue D	sue Date Name of Patentee of Applicant Relevan		s,Columns,Lines where ant Passages or Relevant es Appear			
	1									
If you wis	h to add	additional U.S. Pater	nt citatio	n inform	ation pl	ease click the	Add button.		Add	
			U.S.P	ATENT	APPLI	CATION PUBL	LICATIONS		Remove	
Examiner Initial*	Cite N	o Publication Number	Kind Code <sup>1</sup>	Publica Date	Publication Name of Patentee of Applicant Releva		s,Columns,Lines where rant Passages or Relevant es Appear			
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				FOREIC	N PAT	ENT DOCUM	ENTS		Remove	
Examiner Initial*	l I	Foreign Document Number³	Country Code <sup>2</sup> i		Kind Code <sup>4</sup>	Publication Date	Name of Patented Applicant of cited Document	e or V	Pages,Columns,I where Relevant Passages or Rele Figures Appear	T5
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Examiner Initials*										

# INFORMATION DISCLOSURE STATEMENT BY APPLICANT

( Not for submission under 37 CFR 1.99)

Application Number		15180431	
Filing Date		2016-06-13	
First Named Inventor	Inge Bruheim		
Art Unit		1672	
Examiner Name CUTLI		IFF, YATE KAI RENE	
Attorney Docket Number		AKBM-14409/US-12/CON	

	1	Petition for Inter Partes Review, U.S. Patent No. 9,078,905, Case No.: IPR2017-00745, filed January 27, 2017					
	2	Petition for Inter Partes Review, U.S. Patent No. 9,078,905, Case No.: IPR2017-00747, filed January 27, 2017					
	3	Petition for Inter Partes Review, U.S. Patent No. 9,028,877, Case No.: IPR2017-00746, filed February 3, 2017					
	4	Petition for Inter Partes Review, U.S. Patent No. 9,028,877, Case No.: IPR2017-00748, filed February 3, 2017					
	5	Respondents' Notice of Prior Art, United States International Trade Commission, Investigation No. 337-TA-1019, dated February 1, 2017					
	6	Notice of Opposition, Rimfrost AS, AU Patent Application No. 2014256345, filed March 1, 2017					
	7	Notice of Opposition, Enzymotec Ltd., AU Patent Application No. 2014256345, filed March 1, 2017					
	8	Respondents' Motion for Leave to Amend Their Response to the Complaint and Notice of Investigation, United States International Trade Commission, Investigation No. 337-TA-1019, dated March 14, 2017					
If you wis	h to ad	d additional non-patent literature document citation information please click the Add button Add					
EXAMINER SIGNATURE							
Examiner	Signa	ture Date Considered					
		itial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through a conformance and not considered. Include copy of this form with next communication to applicant.					

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

RIMFROST EXHIBIT 1111 Page 0029

English language translation is attached.

# INFORMATION DISCLOSURE STATEMENT BY APPLICANT

( Not for submission under 37 CFR 1.99)

Application Number		15180431	
Filing Date		2016-06-13	
First Named Inventor	Inge E	Bruheim	
Art Unit		1672	
Examiner Name	CUTL	IFF, YATE KAI RENE	
Attorney Docket Number		AKBM-14409/US-12/CON	

#### **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)	2017-03-17
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.** 

# **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 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.
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Electronic Acknowledgement Receipt				
EFS ID:	28667645			
Application Number:	15180431			
International Application Number:				
Confirmation Number:	2763			
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS			
First Named Inventor/Applicant Name:	Inge Bruheim			
Customer Number:	72960			
Filer:	John Mitchell Jones			
Filer Authorized By:				
Attorney Docket Number:	AKBM-14409/US-12/CON			
Receipt Date:	17-MAR-2017			
Filing Date:	13-JUN-2016			
Time Stamp:	17:28:16			
Application Type:	Utility under 35 USC 111(a)			

# **Payment information:**

Submitted with Payment	no
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# File Listing:

Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1	Information Disclosure Statement (IDS) Form (SB08)	14409US12CON_IDS_3-17-17. pdf	1035601 e259241296879cd9881526fea7f252c72661 059a	no	4

Warnings: RIMFROST EXHIBIT 1111 Page 0032

#### Information:

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#### Total Files Size (in bytes):

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# New Applications Under 35 U.S.C. 111

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#### New International Application Filed with the USPTO as a Receiving Office

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#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: Inge Bruheim et al. Confirmation No.: 2763
Serial No.: 15/180,431 Group No.: 1672
Filed: 13-Jun-2016 Examiner: CUTLIFF

Entitled: BIOEFFECTIVE KRILL OIL COMPOSITIONS

# AMENDMENT AFTER NOTICE OF ALLOWANCE UNDER 37 C.F.R. § 1.312

Commissioner of Patents Mail Stop - Amendment P.O. Box 1450 Arlington, VA 22313-1450

#### **Examiner Cutliff:**

Applicants respectfully request entry of this amendment under 37 C.F.R. §1.312.

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. **AKBM-14409/US-12/CON**. This paragraph is intended to be a CONSTRUCTIVE PETITION FOR EXTENSION OF TIME in accordance with 37 C.F.R. § 1.136(a)(3).

Amendments to the Claims begin at page 2.

**Remarks** begin at page 5.

## **AMENDMENTS TO THE CLAIMS:**

This listing of the claims replaces all prior versions and listings of the claims. Please amend the claims as follows:

- 1. (Previously presented) A method of production of krill oil comprising:
  - a) providing krill;
- b) treating said krill to denature lipases and phospholipases in said krill to provide a denatured krill product;
  - c) storing said denatured krill product for a storage period of from 1 to 24 months;
- d) after said storage period, extracting oil from said denatured krill product with a polar solvent to provide a krill oil with from about 3% to about 15% ether phospholipids w/w of said krill oil astaxanthin esters in amount of greater than about 100 mg/kg of said krill oil.
- 2. (Original) The method of claim 1, wherein said steps a and b are performed on a ship.
- 3. (Original) The method of claim 1, wherein said treating comprises heating.
- 4. (Original) The method of claim 1, wherein said denatured krill product is a krill meal.
- 5. (Original) The method of claim 1, wherein said krill is freshly harvested.
- 6. (Original) The method of claim 1, further comprising encapsulating said krill oil.
- 7. (Original) The method of claim 1, wherein said krill is Antarctic krill.
- 8. (Currently amended) The method of <u>claim 7elaim 1</u>, wherein said Antarctic krill is Euphausia superba.
- 9. (Original) The method of claim 1, wherein said krill oil contains astaxanthin esters in an amount of greater than about 200 mg/kg of said krill oil.

- 10. (Original) The method of claim 1, wherein said krill oil comprises at least 30% total phospholipids w/w of said krill oil.
- 11. (Original) The method of claim 1, wherein said krill oil comprises at least 30% phosphatidylcholine w/w of said krill oil.
- 12. (Previously presented) A method of production of krill oil comprising:
- a) obtaining a denatured krill product produced by treating freshly harvested krill krill to denature lipases and phospholipases in said krill and that has been stored from 1 to 24 months; and
- b) extracting oil from said denatured krill product that has been stored from 1 to 24 months with a polar solvent to provide a krill oil with from about 3% to about 15% ether phospholipids w/w of said krill oil astaxanthin esters in amount of greater than about 100 mg/kg of said krill oil.
- 13. (Previously presented) The method of claim 12, wherein said treating comprises heating.
- 14. (Previously presented) The method of claim 12, wherein said denatured krill product is a krill meal.
- 15. (Previously presented) The method of claim 12, wherein said krill is freshly harvested.
- 16. (Previously presented) The method of claim 12, further comprising encapsulating said krill oil.
- 17. (Previously presented) The method of claim 12, wherein said krill is Antarctic krill.
- 18. (Currently amended) The method of <u>claim 17</u> elaim 10, wherein said Antarctic krill is Euphausia superba.

- 19. (Previously presented) The method of claim 12, wherein said krill oil contains astaxanthin esters in an amount of greater than about 200 mg/kg of said krill oil.
- 20. (Previously presented) The method of claim 12, wherein said krill oil comprises at least 30% total phospholipids w/w of said krill oil.

# **REMARKS**

Applicants now amend claims 8 and 18 to correct typographical errors. The amendments add no new matter.

Applicants respectfully request entry of the present amendment, pursuant to 37 C.F.R. §1.312. In the event the Examiner believes the amendment should not be entered, applicants respectfully request that she telephone the undersigned to set up an interview.

Please charge any fee required for entry of this Amendment, or credit any overpayment, to Deposit Account No. 50-4302.

Respectfully submitted,

Dated: March 14, 2017

/J. Mitchell Jones/
J. Mitchell Jones
Reg. No. 44,174
Casimir Jones, S.C.
2275 Deming Way, Suite 310
Middleton, Wisconsin 53562
608.662.1277

Electronic Acknowledgement Receipt					
EFS ID:	28631570				
Application Number:	15180431				
International Application Number:					
Confirmation Number:	2763				
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS				
First Named Inventor/Applicant Name:	Inge Bruheim				
Customer Number:	72960				
Filer:	John Mitchell Jones/Mallory Checkett				
Filer Authorized By:	John Mitchell Jones				
Attorney Docket Number:	AKBM-14409/US-12/CON				
Receipt Date:	15-MAR-2017				
Filing Date:	13-JUN-2016				
Time Stamp:	14:00:22				
Application Type:	Utility under 35 USC 111(a)				

# **Payment information:**

Submitted with Payment	no
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# File Listing:

Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
			83285		
1		14409US12CON_312Amendme nt.pdf	1a47b366751694062ee4dc14582c89443be 43e0e	yes	5

	Multipart Description/PDF files in .zip description								
	Document Description	Start	End						
	Amendment after Notice of Allowance (Rule 312)	1	1						
	Claims	2	4						
	Applicant Arguments/Remarks Made in an Amendment	5	5						
Warnings:									

Information:

Total Files Size (in bytes): 83285

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

#### 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.



# United States Patent and Trademark Office

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.			
15/180,431	7/180,431 06/13/2016 Inge Bruheim		AKBM-14409/US-12/CON 2763				
72960 Casimir Jones	7590 02/14/2017 S.C.		EXAM	INER			
	G WAY, SUITE 310		CUTLIFF, YAT	TE KAI RENE			
	,		ART UNIT	PAPER NUMBER			
			1672	***************************************			
			NOTIFICATION DATE	DELIVERY MODE			
			02/14/2017	ELECTRONIC			

# 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.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

docketing@casimirjones.com pto.correspondence@casimirjones.com

	Application No.	Applicant(s)
	15/180,431	
Response to Rule 312 Communication	Examiner	Art Unit
The MAILING DATE of this communication a	ppears on the cover sheet	with the correspondence address –
<ol> <li>The amendment filed on <u>08 February 2017</u> under 37 CF</li> <li>a)</li></ol>	R 1.312 has been considere	ed, and has been:
b)  entered as directed to matters of form not affecting	g the scope of the invention.	
c)  disapproved because the amendment was filed aff	ter the payment of the issue	fee.
Any amendment filed after the date the issue for and the required fee to withdraw the application	ee is paid must be accompan	
d) disapproved. See explanation below.		
e)  entered in part. See explanation below.		
L.HILL		
PUBLISHING DIVISION		

U.S. Patent and Trademark Office PTOL-271 (Rev. 04-01)

Attorney Docket No.: AKBM-14409/US-12/CON

# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

 Serial No.:
 15/180,431
 Confirmation No.:
 2763

 Filed:
 13-Jun-2016
 Art Unit:
 1672

First Inventor: Bruheim et al. Examiner: CUTLIFF, Yate Kai Rene

Title: BIOEFFECTIVE KRILL OIL COMPOSITIONS

# RESPONSE TO THE NOTICE TO FILE CORRECTED APPLICATION PAPERS MAILED JANUARY 20, 2017

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

In response to the Notice to File Corrected Application Papers mailed January 20, 2017, Applicant submits the following:

Amendments to the Specification begin on page 2 of this paper; and Remarks begin on page 3 of this paper.

#### AMENDMENTS TO THE SPECIFICATION

Please amend the specification at page 50, lines 8-20 as follows:

The purpose of this experiment was to investigate the effect of dietary krill oil on metabolic parameters in high-fat fed mice and to compare the effect of dietary krill oil with that of fish oil containing the same amount of omega-3 fatty acids. Four groups of C57BL/6 mice (n = 10 per group) were fed 1) chow (N), 2) high fat diet comprising 21% butter fat and 0.15% cholesterol (HF), 3) high fat diet + krill oil (HFKO) or 4) high fat diet + fish oil (HFFO). Treatment 3 contained 2.25% (w/w) krill oil as prepared in example 5 (except that the astaxanthin content was 500 ppm) which were equivalent to 0.36% omega-3 fatty acids. Treatment 4 also contained 0.36% omega-3 fatty acids obtained from regular 18-12 fish oil. The diets were fed to the mice for 7 weeks with free access to drinking water. Data represented in this example means  $\pm$  SE. Columns not sharing a common letter are significantly different (P < 0.05) by ANOVA followed by Tukey's multiple comparison test. N = normal chow diet (n = 10); HF = high-fat diet (n = 10); HFFO = high-fat diet supplemented with fish oil (n = 9); HFKO = high-fat diet supplemented with krill oil (n = 8). The data are presented in Figures 12-1948-25.

## **REMARKS**

In response to the Notice to File Corrected Application Papers mailed January 20, 2017, Applicant has amended the specification to correct the figure designations in Example 12. No new matter has been added.

No fees are believed to be due in connection with this filing. Nevertheless, if the Director finds any additional fees to be due in connection with this, or any other filing, authorization is given to charge said fees to Deposit Account No. 50-4302, referencing attorney docket number AKBM-14409/US-12/CON.

Respectfully,

Date: February 8, 2017

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

Fax: (608) 662-1276

Electronic Acknowledgement Receipt					
EFS ID:	28295230				
Application Number:	15180431				
International Application Number:					
Confirmation Number:	2763				
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS				
First Named Inventor/Applicant Name:	Inge Bruheim				
Customer Number:	72960				
Filer:	John Mitchell Jones/Mallory Checkett				
Filer Authorized By:	John Mitchell Jones				
Attorney Docket Number:	AKBM-14409/US-12/CON				
Receipt Date:	08-FEB-2017				
Filing Date:	13-JUN-2016				
Time Stamp:	14:33:51				
Application Type:	Utility under 35 USC 111(a)				

# **Payment information:**

Submitted with Payment	no
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# File Listing:

Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
			118219		
1		14409US12CON_RNFCAP.pdf	870dc594ca7148858442502f6d7f80aa3ab7 f516	yes	3

	Multipart Description/PDF files in .zip description								
	Document Description	Start	End						
	Amendment after Notice of Allowance (Rule 312)	1	1						
	Specification	2	2						
	Applicant Arguments/Remarks Made in an Amendment	3	3						
Warnings:		•							

Information:

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		_	_					 					

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

#### 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.

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
15/180,431	06/13/2016	Inge Bruheim	AKBM-14409/US-12/CON	2763
72960 Casimir Jones,	7590 01/20/2017		EXAM	INER
	G WAY, SUITE 310		CUTLIFF, YA1	TE KAI RENE
MIDDLETON	, W1 33302		ART UNIT	PAPER NUMBER
			1672	
			NOTIFICATION DATE	DELIVERY MODE
			01/20/2017	ELECTRONIC

# 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.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

docketing@casimirjones.com pto.correspondence@casimirjones.com

Commissioner for Patents United States Patent and Trademark Office P.O. Box 1450 Alexandria, VA 22313-1450 www.uspto.gov

Application No.: 15180431

Applicant: Bruheim Filing Date: 06/13/2016 Date Mailed: 01/20/2017

# NOTICE TO FILE CORRECTED APPLICATION PAPERS

# Notice of Allowance Mailed

This application has been accorded an Allowance Date and is being prepared for issuance. The application, however, is incomplete for the reasons below.

Applicant is given two (2) months from the mail date of this Notice within which to respond. This time period for reply is extendable under 37 CFR 1.136(a) for only TWO additional MONTHS.

The informalities requiring correction are indicated in the attachment(s). If the informality pertains to the abstract, specification (including claims) or drawings, the informality must be corrected with an amendment in compliance with 37 CFR 1.121 (or, if the application is a reissue application, 37 CFR 1.173). Such an amendment may be filed after payment of the issue fee if limited to correction of informalities noted herein. See Waiver of 37 CFR 1.312 for Documents Required by the Office of Patent Publication, 1280 Off. Gaz. Patent Office 918 (March 23, 2004). In addition, if the informality is not corrected until after payment of the issue fee, for purposes of 35 U.S.C. 154(b)(1)(iv), "all outstanding requirements" will be considered to have been satisfied when the informality has been corrected. A failure to respond within the above-identified time period will result in the application being ABANDONED.

## See attachment(s).

A copy of this notice <u>MUST</u> be returned with the reply. Please address response to "Mail Stop Issue Fee, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450".

/Quang Nguyen/ Publication Branch Office of Data Management (571) 272-4200

# **Application No. <u>15180431</u>**

# IDENTIFICATION OF SPECIFICATION/DRAWING INCONSISTENCIES

	On Page of the specification there is a brief description of FIG., but the drawings filed do not include a drawing with that designation. Applicant must respond either by supplying the omitted drawing or by amending the specification to remove all references to that drawing.
	The drawings filed include FIG., but the specification's brief description of the drawings does not describe a drawing with that designation. Applicant must respond either by amending the specification to add a brief description of that drawing or by correcting the drawings to remove the drawing in question.
	Drawings are present in the application and are referred to in the detailed description of the invention, but the specification does not contain a brief description of the drawings as required by 37 CFR 1.74 and 37 CFR 1.77(b)(8).
X	Page 50, line 20 of the specification refers to FIG. 20-25, but no drawing with that designation is described in the brief description of the drawings and no drawing with that designation is present in the application. Applicant must respond either by amending the specification to remove all references to that drawing, or by supplying that drawing and amending the specification to add a brief description of it.
	In the reissue application, FIG., is labeled as "New" but is not described in the reissue specification's brief description of the drawings. Applicant must respond by amending the reissue specification's brief description of the drawings to add a brief description of the new drawing.
	OTHER:
	COMMENTS:

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450

www.uspto.go

# NOTICE OF ALLOWANCE AND FEE(S) DUE

72960 12/21/2016 Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON, WI 53562

**EXAMINER** CUTLIFF, YATE KAI RENE ART UNIT PAPER NUMBER

1672

DATE MAILED: 12/21/2016

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
15/180.431	06/13/2016	Inge Bruheim	AKBM-14409/US-12/CON	2763

TITLE OF INVENTION: BIOEFFECTIVE KRILL OIL COMPOSITIONS

APPLN. TYPE	ENTITY STATUS	ISSUE FEE DUE	PUBLICATION FEE DUE	PREV. PAID ISSUE FEE	TOTAL FEE(S) DUE	DATE DUE
nonprovisional	UNDISCOUNTED	\$960	\$0	\$0	\$960	03/21/2017

THE APPLICATION IDENTIFIED ABOVE HAS BEEN EXAMINED AND IS ALLOWED FOR ISSUANCE AS A PATENT. PROSECUTION ON THE MERITS IS CLOSED. THIS NOTICE OF ALLOWANCE IS NOT A GRANT OF PATENT RIGHTS. THIS APPLICATION IS SUBJECT TO WITHDRAWAL FROM ISSUE AT THE INITIATIVE OF THE OFFICE OR UPON PETITION BY THE APPLICANT. SEE 37 CFR 1.313 AND MPEP 1308.

THE ISSUE FEE AND PUBLICATION FEE (IF REQUIRED) MUST BE PAID WITHIN THREE MONTHS FROM THE MAILING DATE OF THIS NOTICE OR THIS APPLICATION SHALL BE REGARDED AS ABANDONED. STATUTORY PERIOD CANNOT BE EXTENDED. SEE 35 U.S.C. 151. THE ISSUE FEE DUE INDICATED ABOVE DOES NOT REFLECT A CREDIT FOR ANY PREVIOUSLY PAID ISSUE FEE IN THIS APPLICATION. IF AN ISSUE FEE HAS PREVIOUSLY BEEN PAID IN THIS APPLICATION (AS SHOWN ABOVE), THE RETURN OF PART B OF THIS FORM WILL BE CONSIDERED A REQUEST TO REAPPLY THE PREVIOUSLY PAID ISSUE FEE TOWARD THE ISSUE FEE NOW DUE.

#### HOW TO REPLY TO THIS NOTICE:

I. Review the ENTITY STATUS shown above. If the ENTITY STATUS is shown as SMALL or MICRO, verify whether entitlement to that entity status still applies.

If the ENTITY STATUS is the same as shown above, pay the TOTAL FEE(S) DUE shown above.

If the ENTITY STATUS is changed from that shown above, on PART B - FEE(S) TRANSMITTAL, complete section number 5 titled "Change in Entity Status (from status indicated above)".

For purposes of this notice, small entity fees are 1/2 the amount of undiscounted fees, and micro entity fees are 1/2 the amount of small entity

II. PART B - FEE(S) TRANSMITTAL, or its equivalent, must be completed and returned to the United States Patent and Trademark Office (USPTO) with your ISSUE FEE and PUBLICATION FEE (if required). If you are charging the fee(s) to your deposit account, section "4b" of Part B - Fee(s) Transmittal should be completed and an extra copy of the form should be submitted. If an equivalent of Part B is filed, a request to reapply a previously paid issue fee must be clearly made, and delays in processing may occur due to the difficulty in recognizing the paper as an equivalent of Part B.

III. All communications regarding this application must give the application number. Please direct all communications prior to issuance to Mail Stop ISSUE FEE unless advised to the contrary.

IMPORTANT REMINDER: Utility patents issuing on applications filed on or after Dec. 12, 1980 may require payment of maintenance fees. It is patentee's responsibility to ensure timely payment of maintenance fees when due.

#### PART B - FEE(S) TRANSMITTAL

# Complete and send this form, together with applicable fee(s), to: Mail Mail Stop ISSUE FEE

Commissioner for Patents P.O. Box 1450

Alexandria, Virginia 22313-1450 (571)-273-2885 or <u>Fax</u>

INSTRUCTIONS: This form should be used for transmitting the ISSUE FEE and PUBLICATION FEE (if required). Blocks 1 through 5 should be completed where appropriate. All further correspondence including the Patent, advance orders and notification of maintenance fees will be mailed to the current correspondence address as indicated unless corrected below or directed otherwise in Block 1, by (a) specifying a new correspondence address; and/or (b) indicating a separate "FEE ADDRESS" for

maintenance fee notifications. Note: A certificate of mailing can only be used for domestic mailings of the Fee(s) Transmittal. This certificate cannot be used for any other accompanying papers. Each additional paper, such as an assignment or formal drawing, must have its own certificate of mailing or transmission. CURRENT CORRESPONDENCE ADDRESS (Note: Use Block 1 for any change of address) Certificate of Mailing or Transmission I hereby certify that this Fee(s) Transmittal is being deposited with the United States Postal Service with sufficient postage for first class mail in an envelope addressed to the Mail Stop ISSUE FEE address above, or being facsimile transmitted to the USPTO (571) 273-2885, on the date indicated below. 72960 7590 12/21/2016 Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON, WI 53562 (Depositor's name (Signature (Date APPLICATION NO. FILING DATE FIRST NAMED INVENTOR ATTORNEY DOCKET NO. CONFIRMATION NO. 15/180.431 06/13/2016 Inge Bruheim AKBM-14409/US-12/CON 2763 TITLE OF INVENTION: BIOEFFECTIVE KRILL OIL COMPOSITIONS APPLN. TYPE ISSUE FEE DUE PUBLICATION FEE DUE PREV. PAID ISSUE FEE **ENTITY STATUS** TOTAL FEE(S) DUE DATE DUE UNDISCOUNTED \$0 03/21/2017 \$960 \$0 \$960 nonprovisional **EXAMINER** ART UNIT CLASS-SUBCLASS CUTLIFF, YATE KAI RENE 1672 554-210000 1. Change of correspondence address or indication of "Fee Address" (37 CFR 1.363). 2. For printing on the patent front page, list (1) The names of up to 3 registered patent attorneys ☐ Change of correspondence address (or Change of Correspondence Address form PTO/SB/122) attached. or agents OR, alternatively, (2) The name of a single firm (having as a member a registered attorney or agent) and the names of up to 2 registered patent attorneys or agents. If no name is listed, no name will be printed. ☐ "Fee Address" indication (or "Fee Address" Indication form PTO/SB/47; Rev 03-02 or more recent) attached. Use of a Customer Number is required. 3. ASSIGNEE NAME AND RESIDENCE DATA TO BE PRINTED ON THE PATENT (print or type) PLEASE NOTE: Unless an assignee is identified below, no assignee data will appear on the patent. If an assignee is identified below, the document has been filed for recordation as set forth in 37 CFR 3.11. Completion of this form is NOT a substitute for filing an assignment. (A) NAME OF ASSIGNEE (B) RESIDENCE: (CITY and STATE OR COUNTRY) Please check the appropriate assignee category or categories (will not be printed on the patent): 🔲 Individual 📮 Corporation or other private group entity 🖵 Government 4a. The following fee(s) are submitted: 4b. Payment of Fee(s): (Please first reapply any previously paid issue fee shown above) ☐ Issue Fee A check is enclosed. ☐ Publication Fee (No small entity discount permitted) Payment by credit card. Form PTO-2038 is attached. Advance Order - # of Copies \_ The director is hereby authorized to charge the required fee(s), any deficiency, or credits any overpayment, to Deposit Account Number 5. Change in Entity Status (from status indicated above) NOTE: Absent a valid certification of Micro Entity Status (see forms PTO/SB/15A and 15B), issue fee payment in the micro entity amount will not be accepted at the risk of application abandonment. Applicant certifying micro entity status. See 37 CFR 1.29 ☐ Applicant asserting small entity status. See 37 CFR 1.27  $\underline{NOTE}$ : If the application was previously under micro entity status, checking this box will be taken to be a notification of loss of entitlement to micro entity status. <u>NOTE:</u> Checking this box will be taken to be a notification of loss of entitlement to small or micro entity status, as applicable. Applicant changing to regular undiscounted fee status. NOTE: This form must be signed in accordance with 37 CFR 1.31 and 1.33. See 37 CFR 1.4 for signature requirements and certifications.

# Page 2 of 3 RIMFROST EXHIBIT 1111 Page 0052

Date \_

Registration No. \_

Authorized Signature \_ Typed or printed name \_



# UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450

P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
15/180,431	06/13/2016	Inge Bruheim	AKBM-14409/US-12/CON	2763
72960 75	590 12/21/2016		EXAM	INER
Casimir Jones, S			CUTLIFF, YA	TE KAI RENE
2275 DEMING W	AY, SUITE 310			
MIDDLETON, W			ART UNIT	PAPER NUMBER
			1672	

DATE MAILED: 12/21/2016

# Determination of Patent Term Adjustment under 35 U.S.C. 154 (b)

(Applications filed on or after May 29, 2000)

The Office has discontinued providing a Patent Term Adjustment (PTA) calculation with the Notice of Allowance.

Section 1(h)(2) of the AIA Technical Corrections Act amended 35 U.S.C. 154(b)(3)(B)(i) to eliminate the requirement that the Office provide a patent term adjustment determination with the notice of allowance. See Revisions to Patent Term Adjustment, 78 Fed. Reg. 19416, 19417 (Apr. 1, 2013). Therefore, the Office is no longer providing an initial patent term adjustment determination with the notice of allowance. The Office will continue to provide a patent term adjustment determination with the Issue Notification Letter that is mailed to applicant approximately three weeks prior to the issue date of the patent, and will include the patent term adjustment on the patent. Any request for reconsideration of the patent term adjustment determination (or reinstatement of patent term adjustment) should follow the process outlined in 37 CFR 1.705.

Any questions regarding the Patent Term Extension or Adjustment determination should be directed to the Office of Patent Legal Administration at (571)-272-7702. Questions relating to issue and publication fee payments should be directed to the Customer Service Center of the Office of Patent Publication at 1-(888)-786-0101 or (571)-272-4200.

#### OMB Clearance and PRA Burden Statement for PTOL-85 Part B

The Paperwork Reduction Act (PRA) of 1995 requires Federal agencies to obtain Office of Management and Budget approval before requesting most types of information from the public. When OMB approves an agency request to collect information from the public, OMB (i) provides a valid OMB Control Number and expiration date for the agency to display on the instrument that will be used to collect the information and (ii) requires the agency to inform the public about the OMB Control Number's legal significance in accordance with 5 CFR 1320.5(b).

The information collected by PTOL-85 Part B is required by 37 CFR 1.311. 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 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, Virginia 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, Virginia 22313-1450. Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number.

### **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.
- 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.
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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. Page 0054

# Notice of Allowability Application No. 15/180,431 BRUHEIM ET AL. Examiner YATE' K. CUTLIFF Art Unit 1672 Alla (First Inventor to File) Status No

The MAILING DATE of this communication appears on the All claims being allowable, PROSECUTION ON THE MERITS IS (OR REM herewith (or previously mailed), a Notice of Allowance (PTOL-85) or other a NOTICE OF ALLOWABILITY IS NOT A GRANT OF PATENT RIGHTS. Tof the Office or upon petition by the applicant. See 37 CFR 1.313 and MPE	AINS) CLOSED in this application. If not included appropriate communication will be mailed in due course. THIS his application is subject to withdrawal from issue at the initiative		
<ol> <li>This communication is responsive to <u>10/12/2016</u>.</li> <li>A declaration(s)/affidavit(s) under <b>37 CFR 1.130(b)</b> was/were filed</li> </ol>	d on		
2. An election was made by the applicant in response to a restriction recrequirement and election have been incorporated into this action.	uirement set forth during the interview on; the restriction		
3. The allowed claim(s) is/are <u>1 - 20</u> . As a result of the allowed claim(s), <b>Highway</b> program at a participating intellectual property office for the http://www.uspto.gov/patents/init_events/pph/index.jsp or send an inc	corresponding application. For more information, please see		
4. Acknowledgment is made of a claim for foreign priority under 35 U.S.	C. § 119(a)-(d) or (f).		
Certified copies:			
a) ☐ All b) ☐ Some *c) ☐ None of the:			
1. Certified copies of the priority documents have been rec			
2. Certified copies of the priority documents have been rec			
3. Copies of the certified copies of the priority documents h	nave been received in this national stage application from the		
International Bureau (PCT Rule 17.2(a)).  * Certified copies not received:			
ocitined copies not received			
Applicant has THREE MONTHS FROM THE "MAILING DATE" of this connoted below. Failure to timely comply will result in ABANDONMENT of the THIS THREE-MONTH PERIOD IS NOT EXTENDABLE.			
5. CORRECTED DRAWINGS ( as "replacement sheets") must be subm	itted.		
including changes required by the attached Examiner's Amenda Paper No./Mail Date	nent / Comment or in the Office action of		
Identifying indicia such as the application number (see 37 CFR 1.84(c)) sho each sheet. Replacement sheet(s) should be labeled as such in the header			
<ol> <li>DEPOSIT OF and/or INFORMATION about the deposit of BIOLOGIC attached Examiner's comment regarding REQUIREMENT FOR THE D</li> </ol>			
Attachment(s)			
1. Notice of References Cited (PTO-892)	5.  Examiner's Amendment/Comment		
2. Information Disclosure Statements (PTO/SB/08),	6. ☐ Examiner's Statement of Reasons for Allowance		
Paper No./Mail Date  3. Examiner's Comment Regarding Requirement for Deposit	7.  Other		
of Biological Material			
4. Interview Summary (PTO-413), Paper No./Mail Date			
/YATE' K. CUTLIFF/			
Primary Examiner, Art Unit 1672			

U.S. Patent and Trademark Office PTOL-37 (Rev. 08-13) 20161210

Notice of Allowability

Part of Paper No./Mail Date

Application/Control Number: 15/180,431 Page 2

Art Unit: 1672

# **DETAILED ACTION**

## Notice of Pre-AIA or AIA Status

1. The present application is being examined under the pre-AIA first to invent provisions.

# Response to Amendment

2. The amendment to claims 1, 12 - 17, 19 and 20, submitted October 12, 2016 is acknowledged and entered.

# Response to Arguments

- 3. Applicant's arguments, see page 4, filed October 12, 2016, with respect to the objection of claims 13 17, 19 and 20 have been fully considered and are persuasive. The objection of claims 13 17, 19 and 20 has been withdrawn.
- 4. Applicant's arguments, see page 4, filed October 12, 2016, with respect to the rejection claims 1 11, 13 17, 19 and 20 under 35 USC 112, second paragraph have been fully considered and are persuasive in view of the claim amendment. The rejection of claims 1 11, 13 17, 19 and 20 under 35 USC 112, second paragraph has been withdrawn.

# Double Patenting

5. Applicant's arguments, see page 4, filed October 12, 2016, with respect to the rejection claims 1 - 20 under non-statutory double patenting have been fully considered and are persuasive in view of the acceptance of the Terminal Disclaimer received October 12, 2016. The rejection of claims 1 - 20 under non-statutory double patenting has been withdrawn.

Application/Control Number: 15/180,431 Page 3

Art Unit: 1672

## Terminal Disclaimer

6. The terminal disclaimer filed on October 12, 2016 disclaiming the terminal portion of any patent granted on this application which would extend beyond the expiration date of U.S. Patents 9,119,864; 9,028,877; 9,375,453 and 9,034,388 has been reviewed and is accepted. The terminal disclaimer has been recorded.

# Allowable Subject Matter

- 7. Claims 1 20 are allowed.
- 8. The following is an examiner's statement of reasons for allowance: claims are allowed for reasons of record.

Any comments considered necessary by applicant must be submitted no later than the payment of the issue fee and, to avoid processing delays, should preferably accompany the issue fee. Such submissions should be clearly labeled "Comments on Statement of Reasons for Allowance."

# Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to YATE' K. CUTLIFF whose telephone number is (571)272-9067. The examiner can normally be reached on M-F 8:30 a.m. - 5:30 p.m..

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Fereydoun Sajjadi can be reached on (571) 272-3311. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Application/Control Number: 15/180,431 Page 4

Art Unit: 1672

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.

/YATE' K. CUTLIFF/ Primary Examiner, Art Unit 1672

# Search Notes



Application/Control No.	Applicant(s)/Patent Under Reexamination
15180431	BRUHEIM ET AL.
Examiner	Art Unit
YATE K CUTLIFF	1672

CPC- SEARCHED		
Symbol	Date	Examiner
C11B 1/02, 16: see search history	7/14/2016	
C11b 3/006: see search history	7/14/2016	ykc
C07C 51/48: see search history	7/14/2016	ykc
A61K 31/122, 685: see search history	7/14/2016	ykc

CPC COMBINATION SETS - SEARC	CHED	
Symbol	Date	Examiner

US CLASSIFICATION SEARCHED				
Class	Subclass	Date	Examiner	

SEARCH NOTES				
Search Notes	Date	Examiner		
Palm Inventor Search	12/10/2016	ykc		
East	12/10/2016	ykc		
STN: caplus, agricola USpat2, US patfull, IFIall	7/14/2016	ykc		
C11B 1/02, 16: see search history	7/14/2016			
C11b 3/006: see search history	7/14/2016	ykc		
C07C 51/48: see search history	7/14/2016	ykc		
A61K 31/122, 685: see search history	7/14/2016	ykc		

	INTERFERENCE SEARCH		
US Class/ CPC Symbol	US Subclass / CPC Group	Date	Examiner
C11B	1/02,16: SEE SEARCH HISTORY	12/10/2016	YKC
C11B	3/006: SEE SEARCH HISTORY	12/10/2016	YKC
C07C	51/48: SEE SEARCH HISTORY	12/10/2016	YKC

INTERFERENCE SEARCH								
US Class/ CPC Symbol	US Subclass / CPC Group	Date	Examiner					
A61K	31/122, 685	12/10/2016	YKC					

Doc code: IDS Doc description: Information Disclosure Statement (IDS) Filed PTO/SB/08a (03-15)
Approved for use through 07/31/2016. OMB 0651-0031
U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

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.

	Application Number		15180431		
	Filing Date		2016-06-13		
INFORMATION DISCLOSURE	First Named Inventor Ing		Inge Bruheim		
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit		1672		
(Not lot submission under or or it 1.00)	Examiner Name	CUTL	IFF, YATE KAI RENE		
	Attorney Docket Number	er	AKBM-14409/US-12/CON		

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# INFORMATION DISCLOSURE STATEMENT BY APPLICANT

( Not for submission under 37 CFR 1.99)

Application Number		15180431			
Filing Date		2016-06-13			
First Named Inventor	Inge E	Bruheim			
Art Unit		1672			
Examiner Name	CUTL	IFF, YATE KAI RENE			
Attorney Docket Number	er	AKBM-14409/US-12/CON			

1	Action Closing Prosecution, 348 Patent, mailed May 14, 2013							
2	Buchi R-220 Rotovapor® Manual, dated November 16, 2009, pages 1-50							
3	Certified translation of Ex. 1074: Japanese Patent No. 60-153779, entitled "Nutritional Supplement" ("Fukuoka"); Certificate of Translation provided as Ex. 1075, dated August 16, 2013, 1 page							
4	Certificate of translation of Ex. 1076: Japanese Patent Publication No. H08-231391, entitled "Medicine for Improvement of Dementia Symptoms" ("Yasawa"); Certificate of Translation provided as Ex. 1077, dated August 16, 2013, 1 page							
5	Certification of translation of Ex. 1070: Japanese Unexamined Patent Application Publication No. 02-215351, titled Krill Phospholipids Fractioning Method ("Maruyama,"); Certificate of Translation provided as Ex. 1071; dated July 9, 2013, 1 page							
6	Declaration of Bjorn Ole Haugsgjerd in support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Haugsgjerd"), dated September 30, 2013, 12 pages							
7	Database FSTA [Online] International Food Information Service, Frankfurt-Main; SHIBATA N. "Effect of fishing season on lipid content and composition of Antarctic krill (translated)" Database accession no. FS-1985-04-r-0091, 1983, one page, abstract only							
8	Third Party Observation against corresponding AU Patent Application No. 2014256345, filed May 23, 2016, 50 pages							
9	Third Party Observation against corresponding AU Patent Application NO. 2013227998, filed July 15, 2016, 6 pages							
10	Evidence in Support of Opposition, AU Patent Application No. 2013227998, filed September 22, 2016, 22 pages							
11	Notice of Acceptance of Application, AU Patent Application No. 2013227998, mailed October 5, 2016, 2 pages							
	1 2 3 4 5 6 7 8							

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Examiner Name	CUTL	IFF, YATE KAI RENE			
Attorney Docket Numb	er	AKBM-14409/US-12/CON			

If you wish to add additional non-patent literature document citation information please click the Add button Add								
EXAMINER SIGNATURE								
Examiner Signature	/YATE' K CUTLIFF/	K CUTLIFF/ Date Considered 12/1						
*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.								
Standard ST.3). 3 For Japan	D Patent Documents at <u>www.USPTO.GOV</u> or MPEP 901.04. <sup>2</sup> Enter offinese patent documents, the indication of the year of the reign of the Emppropriate symbols as indicated on the document under WIPO Standard is attached.	peror must precede the seri	ial number of the patent document.					

# INFORMATION DISCLOSURE STATEMENT BY APPLICANT

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Application Number		15180431			
Filing Date		2016-06-13			
First Named Inventor Inge E		Bruheim			
Art Unit		1672			
Examiner Name CUTL		IFF, YATE KAI RENE			
Attorney Docket Numb	er	AKBM-14409/US-12/CON			

#### **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.

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- 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)	2016-10-12
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.** 

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  to the Patent Cooperation Treaty.
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UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

# **BIB DATA SHEET**

# **CONFIRMATION NO. 2763**

SERIAL NUM	IBER	FILING O			CLASS	GR	OUP ART	UNIT	ATTC	RNEY DOCKET
15/180,43	31	06/13/2			554		1672 д		КВМ-	<b>NO.</b> 14409/US-12/CO
		RUL	E							
APPLICANTS AKER BIOMARINE ANTARCTIC AS, Stamsund, NORWAY;										
Inge Bruh Snorre Ti	INVENTORS Inge Bruheim, Volda, NORWAY; Snorre Tilseth, Bergen, NORWAY; Daniele Mancinelli, Orsta, NORWAY;									
This appl wh wh and and and ** <b>FOREIGN A</b>	** CONTINUING DATA ********************************  This application is a CON of 14/020,162 09/06/2013 PAT 9375453  which is a CON of 12/057,775 03/28/2008 PAT 9034388  which claims benefit of 60/920,483 03/28/2007  and claims benefit of 60/975,058 09/25/2007  and claims benefit of 60/983,446 10/29/2007  and claims benefit of 61/024,072 01/28/2008  ** FOREIGN APPLICATIONS ************************************									
Foreign Priority claims		Yes No			STATE OR	S.	HEETS	тот		INDEPENDENT
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Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON, WI 53562 UNITED STATES										
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	Application/Control No.	Applicant(s)/Patent Under Reexamination
Index of Claims	15180431	BRUHEIM ET AL.
	Examiner	Art Unit
	YATE K CUTLIFF	1672

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U.S. Patent and Trademark Office Part of Paper No.: 20161210

# **EAST Search History**

# **EAST Search History (Prior Art)**

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
S1	36	(("BRUHEIM") near3 ("Inge")).INV.	US- PGPUB; USPAT; USOCR	OR	OFF	2016/07/12 16:47
S2	27	(("TILSETH") near3 ("Snorre")).I <b>NV</b> .	US- PGPUB; USPAT; USOCR	OR	OFF	2016/07/12 16:47
S3	20	(("MANCINELLI") near3 ("Daniele")).INV.	US- PGPUB; USPAT; USOCR	OR	OFF	2016/07/12 16:47
S4	7	("20030044495"   "20040241249"   "20080166419"   "4119619"   "5266564"   "5434183"   "6537787"   "6800299"   "7666447"   "8030348").PN.	USPAT	OR	OFF	2016/07/13 13:44
S5	19	("20030044495"   "20040241249"   "20080166419"   "4119619"   "5266564"   "5434183"   "6537787"   "6800299"   "7666447"   "8030348").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:44
S6	5	("4714571"   "8278351"   "8383675").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:44
S7	28	("2652235"   "4036993"   "4251557"   "4505936"   "5006281"   "6214396"   "6346276").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:45
S8	21	("20020076468"   "20030113432"   "20060078625"   "20080166419"   "20080166420"   "20100143571"   "20100160659"   "20110130458"   "4133077"   "4749522"   "4814111"   "7488503"   "8697138").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:46
S9	1	("20060078625"   "8057825").PN.	USPAT	OR	OFF	2016/07/13 13:46
S10	3	("20060078625"   "8057825").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:46
S11	10	("20050003073"   "20110160161"   "20110256216"   "4038722"   "8586567").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:47
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S12	0	("5266564," "8030348," "7666447," "4714571," "8278351," "8383675," "2652235," " 5006281").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:11
S13	0	("5266564," "8030348," "7666447," "4714571," "8278351," "8383675," "2652235," "5006281").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:11
S14	24	("5266564," "8030348," "7666447," "4714571," "8278351," "8383675," "2652235," "5006281").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:12
S15	22	("4251557," "4505936," "6214396," "4036993," "6346276," "8697138," "7488503," "4749522," "4133077").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:14
S16	12	("20060193962," "20080166419," "20110130458," "20080166420," "20060078625," "20020076468," "20030113432," "20100143571," "20100160659," "20080166419").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:16
S17	9	("9119864," "9072752," "9034388," "9028877," "9078905," "8372812," "8697138").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:18
S18	0	("20140274968," "20150030718," "20140088043," "20140088047," "20140080791," "20150164841," "20140363517," "20100226977," "20150050403," "20140005421," "20140107072," "20090061067," "20140010888").pn.	USPAT	OR	OFF	2016/07/13 14:21
S19	22	("20140274968," "20150030718," "20140088043," "20140088047," "20140080791," "20150164841," "20140363517," "20100226977," "20150050403," "20140005421," "20140107072," "20090061067," "20140010888").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:21
S20	10	(S5 or S6 or S7 or S8 or S10) and krill	USPAT	OR	OFF	2016/07/13 14:22
S21	15	(S11 or S14 or S15 or S16 or S17 or S19) and krill	USPAT	OR	OFF	2016/07/13 14:23
S22	16	(S20 or S21) and (extraction or extract or extracting) and oil	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:24
S23	20	(S5 or S6 or S7 or S8 or S10) and krill	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:24
S24	40	(S11 or S14 or S15 or S16 or S17 or S19) and krill	US- PGPUB;	OR OST EX	OFF	2016/07/13 14:25 <b>1111 P</b> a

			USPAT; USOCR; FPRS			
S25	42	(S23 or S24) and (extraction or extract or extracting) and oil	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:25
S26	10	S25 and (polar near2 solvent)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:27
S27	3	("20100226977"   "20140274968"   "20140370115").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:47
S28	0	S27 and krill	USPAT	OR	OFF	2016/07/14 09:47
S29	0	S27 and (extraction or extracting or extract) and pholpholipid?	USPAT	OR	OFF	2016/07/14 09:48
S30	0	S27 and (extraction or extracting or extract)	USPAT	OR	OFF	2016/07/14 09:48
S31	3	S27 and (extraction or extracting or extract)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:49
S32	3	S27 and (extraction or extracting or extract) and ((polar near2 solvent) or ethanol)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:50
<b>S</b> 33	3	S27 and krill	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:53
S34	696	krill and oil and astaxanthin	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 10:55
<b>S</b> 35	201	(krill and oil and astaxanthin) and phosphatidylcholine	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 10:56
S36	13	denature near3 lipase	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 11:17
S37	21	denature near3 lipases	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 11:17

31	\f	3	30	7	31	¥==
S38		S34 and S37	USPAT	OR	OFF	2016/07/14 11:17
S39	52	(krill adj oil) and (extraction or extracting or extract) and (ship or boat)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 13:36
S40	0	S39 and @ay<="2007"	USPAT	OR	OFF	2016/07/14 13:36
S41	1	S39 and @ay<="2008"	USPAT	OR	OFF	2016/07/14 13:36
S42	50	((krill adj oil) same (extraction or extracting or extract)) and (ship or boat)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 13:37
S43	179	c11b3/006.cpc.	USPAT	OR	OFF	2016/07/14 15:55
S44	0	c11b1/02,16,cpc.	USPAT	OR	OFF	2016/07/14 15:56
S45	133	c11b1/02,16.cpc.	USPAT	OR	OFF	2016/07/14 15:57
S46	664	c07c51/48.cpc.	USPAT	OR	OFF	2016/07/14 15:57
S47	1226	a61k31/122,685.cpc.	USPAT	OR	OFF	2016/07/14 15:57
S48	236	a61k9/48.cpc.	USPAT	OR	OFF	2016/07/14 15:58
S49	201	(krill and oil and astaxanthin) and phosphatidylcholine	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 16:01
S50	9	S43 and S49	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 16:01
S51	6	S45 and S49	USPAT	OR	OFF	2016/07/14 16:03
S52	7	S45 and (krill near2 oil)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:04
S53	1	S46 and (krill near2 oil)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:08
S54	55	S47 and (krill near2 oil)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:09
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S55	7	S47 and (krill near2 oil) and (denature near3 lipases)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:09
S56	7	S48 and (krill near2 oil) and (denature near3 lipases)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:10

## Issue Classification



15180431 BRUHEIM ET AL.

Examiner Art Unit

YATE K CUTLIFF 1672

CPC						
Symbol			Туре	Version		
C11B	3	006	F	2013-01-01		
A61K	9	/ 4858	1	2013-01-01		
A61K	31	122	I	2013-01-01		
A61K	31	23	1	2013-01-01		
A61K	31	/ 683	1	2013-01-01		
A61K	31	685		2013-01-01		
A61K	35	612	I	2013-01-01		
A61K	45	<i>f</i> 06	1	2013-01-01		
A61K	31	202	1	2013-01-01		
A61K	9	48	1	2013-01-01		
A61K	31	20		2013-01-01		
A61K	31	235	1	2013-01-01		
A61K	9	0053	1	2013-01-01		
A61K	9	4825		2013-01-01		

CPC Combination Sets							
Symbol			Туре	Set	Ranking	Version	
A61K	31	23	1	1	1	2013-01-01	
A61K	2300	00	А	1	2	2013-01-01	
A61K	31	683	1	2	1	2013-01-01	
A61K	2300	<i>I</i> 00	A	2	2	2013-01-01	
A61K	31	685	1	3	1	2013-01-01	
A61K	2300	00	A	3	2	2013-01-01	
A61K	31	122	1	4	1	2013-01-01	
A61K	2300	00	A	4	2	2013-01-01	

NONE	Total Claims Allowed:			
(Assistant Examiner)	(Date)	2	0	
/YATE K CUTLIFF/ Primary Examiner.Art Unit 1672	12/10/2016	O.G. Print Claim(s)	O.G. Print Figure	
(Primary Examiner)	(Date)	1	1	

U.S. Patent and Trademark Office Part of Paper No. 20161210

## Issue Classification

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Application/Control No.	Applicant(s)/Patent Under Reexamination
15180431	BRUHEIM ET AL.
Examiner	Art Unit
YATE K CUTLIFF	1672

US ORIGINAL CLASSIFICATION					INTERNATIONAL CLASSIFICATION										
	CLASS SUBCLASS				CLAIMED						NON-CLAIMED				
210				С	1	1	В	3 / 00 (2006.01.01)							
	CR	OSS REFI	ERENCE(	S)											
CLASS	SUB	CLASS (ONE	(ONE SUBCLASS PER BLOCK)												

NONE		Total Clain	ıs Allowed:
(Assistant Examiner)	(Date)	2	0
/YATE K CUTLIFF/ Primary Examiner.Art Unit 1672	12/10/2016	O.G. Print Claim(s)	O.G. Print Figure
(Primary Examiner)	(Date)	1	1

## Issue Classification



Application/Control No.	Applicant(s)/Patent Under Reexamination
15180431	BRUHEIM ET AL.
Examiner	Art Unit
VATE K CUTUEE	1679

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	16														

NONE		Total Clain	ıs Allowed:
(Assistant Examiner)	(Date)	2	0
/YATE K CUTLIFF/ Primary Examiner.Art Unit 1672	12/10/2016	O.G. Print Claim(s)	O.G. Print Figure
(Primary Examiner)	(Date)	1	1

Doc Code: DIST.E.FILE Document Description: Electroni	c Terminal Disclaimer - Filed	PTO/SB/26 U.S. Patent and Trademark Office Department of Commerce						
Electronic Petition Request	TERMINAL DISCLAIMER TO OB "PRIOR" PATENT	VIATE A DOUBLE PATENTING REJECTION OVER A						
Application Number	15180431	15180431						
Filing Date	13-Jun-2016							
First Named Inventor	Inge Bruheim	Inge Bruheim						
Attorney Docket Number	AKBM-14409/US-12/CON	AKBM-14409/US-12/CON						
Title of Invention	BIOEFFECTIVE KRILL OIL COMPO	OSITIONS						
Office Action	oes not obviate requirement for respainter is not being used for a Joint Re	ponse under 37 CFR 1.111 to outstanding						
Owner	·	Percent Interest						
AKER BIOMARINE ANTARCTIC AS		100%						
	of any patent granted on the instant	on hereby disclaims, except as provided below, the application which would extend beyond the expiration						
9119864								
9028877								
9375453								
9034388								

as the term of said prior patent is presently shortened by any terminal disclaimer. The owner hereby agrees that any patent so granted on the instant application shall be enforceable only for and during such period that it and the prior patent are commonly owned. This agreement runs with any patent granted on the instant application and is binding upon the grantee, its successors or assigns. In making the above disclaimer, the owner does not disclaim the terminal part of the term of any patent granted on the instant application that would extend to the expiration date of the full statutory term of the prior patent, "as the term of said prior patent is presently shortened by any terminal disclaimer," in the event that said prior patent later: expires for failure to pay a maintenance fee; is held unenforceable; - is found invalid by a court of competent jurisdiction; - is statutorily disclaimed in whole or terminally disclaimed under 37 CFR 1.321; has all claims canceled by a reexamination certificate; - is in any manner terminated prior to the expiration of its full statutory term as presently shortened by any terminal disclaimer. Terminal disclaimer fee under 37 CFR 1.20(d) is included with Electronic Terminal Disclaimer request. **(•**) I certify, in accordance with 37 CFR 1.4(d)(4), that the terminal disclaimer fee under 37 CFR 1.20(d) ()required for this terminal disclaimer has already been paid in the above-identified application. Applicant claims the following fee status: Small Entity Micro Entity Regular Undiscounted  $\odot$ I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon. THIS PORTION MUST BE COMPLETED BY THE SIGNATORY OR SIGNATORIES I certify, in accordance with 37 CFR 1.4(d)(4) that I am: An attorney or agent registered to practice before the Patent and Trademark Office who is of record in ◉ this application Registration Number 44174 A sole inventor A joint inventor; I certify that I am authorized to sign this submission on behalf of all of the inventors as evidenced by the power of attorney in the application A joint inventor; all of whom are signing this request Signature /J. Mitchell Jones/ Name J. Mitchell Jones

\*Statement under 37 CFR 3.73(b) is required if terminal disclaimer is signed by the assignee (owner). Form PTO/SB/96 may be used for making this certification. See MPEP § 324.

Electronic Patent Application Fee Transmittal					
Application Number:	15180431				
Filing Date:	13-Jun-2016				
Title of Invention:	BIC	DEFFECTIVE KRILL OI	L COMPOSITIO	NS	
First Named Inventor/Applicant Name:	Inge Bruheim				
Filer:	John Mitchell Jones/Mallory Checkett				
Attorney Docket Number:	AKBM-14409/US-12/CON				
Filed as Large Entity					
Filing Fees for Utility under 35 USC 111(a)					
Description		Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Basic Filing:					
STATUTORY OR TERMINAL DISCLAIMER		1814	1	160	160
Pages:					
Claims:					
Miscellaneous-Filing:					
Petition:					
Patent-Appeals-and-Interference:					
Post-Allowance-and-Post-Issuance:					

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Extension-of-Time:				
Miscellaneous:				
	Total in USD (\$)		160	

Doc Code: DISQ.E.FILE Document Description: Electronic Terminal Disclaimer – Approved
Application No.: 15180431
Filing Date: 13-Jun-2016
Applicant/Patent under Reexamination: Bruheim et al.
Electronic Terminal Disclaimer filed on October 12, 2016
This patent is subject to a terminal disclaimer
☐ DISAPPROVED
Approved/Disapproved by: Electronic Terminal Disclaimer automatically approved by EFS-Web
U.S. Patent and Trademark Office

Electronic Acknowledgement Receipt			
EFS ID:	27186672		
Application Number:	15180431		
International Application Number:			
Confirmation Number:	2763		
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS		
First Named Inventor/Applicant Name:	Inge Bruheim		
Customer Number:	72960		
Filer:	John Mitchell Jones/Mallory Checkett		
Filer Authorized By:	John Mitchell Jones		
Attorney Docket Number:	AKBM-14409/US-12/CON		
Receipt Date:	12-OCT-2016		
Filing Date:	13-JUN-2016		
Time Stamp:	14:28:43		
Application Type:	Utility under 35 USC 111(a)		

## **Payment information:**

Submitted with Payment	yes
Payment Type	Deposit Account
Payment was successfully received in RAM	\$160
RAM confirmation Number	832
Deposit Account	504302
Authorized User	Jones, J. Mitchell

The Director of the USPTO is hereby authorized to charge indicated fees and credit any overpayment as follows:

File Listing	:				
Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
			34891		
1 Electronic Terminal Disclaimer-Filed	e Terminal-Disclaimer.pdf	6ae6e574a5579bef602d709a816d7ebb553 251a9	no	3	
Warnings:	+				
Information:					
			30522		
2	2 Fee Worksheet (SB06) fee-info.pdf	fee-info.pdf	6e29fc3c11b92658213538b1e53f49710aef 466c	no	2
Warnings:			1		
Information:					

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

Total Files Size (in bytes):

#### 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.

Attorney Docket No.: AKBM-14409/US-12/CON

#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: Bruheim et al. Art Unit: 1672
Serial No.: 15/180,431 Examiner: Cutliff
Filed: 06/13/2016 Confirmation: 2763

Entitled: BIOEFFECTIVE KRILL OIL COMPOSITIONS

# RESPONSE TO OFFICE ACTION MAILED JULY 21, 2016

#### **EFS WEB FILED**

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

#### Examiner Cutliff:

This communication is responsive to the Office Action mailed July 21, 2016. 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. **AKBM-14409/US-12/CON**. This paragraph is intended to be a CONSTRUCTIVE PETITION FOR EXTENSION OF TIME in accordance with 37 C.F.R. § 1.136(a)(3).

#### **CLAIM AMENDMENTS:**

- 1. (Currently amended) A method of production of krill oil comprising:
  - a) providing krill;
- b) treating said krill to denature lipases and phospholipases in said krill to provide a denatured krill product;
  - c) storing said denatured krill product for a storage period of from 1 to 24 months;
- d) <u>after said storage period</u>, extracting oil from said denatured krill product with a polar solvent to provide a krill oil with from about 3% to about 15% ether phospholipids w/w of said krill oil astaxanthin esters in amount of greater than about 100 mg/kg of said krill oil.
- 2. (Original) The method of claim 1, wherein said steps a and b are performed on a ship.
- 3. (Original) The method of claim 1, wherein said treating comprises heating.
- 4. (Original) The method of claim 1, wherein said denatured krill product is a krill meal.
- 5. (Original) The method of claim 1, wherein said krill is freshly harvested.
- 6. (Original) The method of claim 1, further comprising encapsulating said krill oil.
- 7. (Original) The method of claim 1, wherein said krill is Antarctic krill.
- 8. (Original) The method of claim 1, wherein said Antarctic krill is Euphausia superba.
- 9. (Original) The method of claim 1, wherein said krill oil contains astaxanthin esters in an amount of greater than about 200 mg/kg of said krill oil.
- 10. (Original) The method of claim 1, wherein said krill oil comprises at least 30% total phospholipids w/w of said krill oil.

- 11. (Original) The method of claim 1, wherein said krill oil comprises at least 30% phosphatidylcholine w/w of said krill oil.
- 12. (Currently amended) A method of production of krill oil comprising:
- a) obtaining a denatured krill product produced by treating freshly harvested krill krill to denature lipases and phospholipases in said krill and that has been stored from 1 to 24 months; and
- b) extracting oil from said denatured krill product that has been stored from 1 to 24 months with a polar solvent to provide a krill oil with from about 3% to about 15% ether phospholipids w/w of said krill oil astaxanthin esters in amount of greater than about 100 mg/kg of said krill oil.
- 13. (Currently amended) The method of claim 12, wherein said treating comprises heating.
- 14. (Currently amended) The method of claim 12, wherein said denatured krill product is a krill meal.
- 15. (Currently amended) The method of claim 12, wherein said krill is freshly harvested.
- 16. (Currently amended) The method of claim 12, further comprising encapsulating said krill oil.
- 17. (Currently amended) The method of claim 12, wherein said krill is Antarctic krill.
- 18. (Original) The method of claim 10, wherein said Antarctic krill is Euphausia superba.
- 19. (Currently amended) The method of claim 12, wherein said krill oil contains astaxanthin esters in an amount of greater than about 200 mg/kg of said krill oil.
- 20. (Currently amended) The method of claim 12, wherein said krill oil comprises at least 30% total phospholipids w/w of said krill oil.

**REMARKS** 

Claims 1-20 are pending and under examination following entry of this amendment. 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.

arguments or rejections, and solely for the purpose of expediting the patent application process

and without waiving the right to prosecute the cancelled claims (or similar claims) in the future.

The pending rejections are addressed in order below.

**Indefiniteness.** The claims have been amended to clarify that the oil is extracted after the storage period in claim 1 and to correct dependencies. Applicant believes that the claims as

amended traverse the indefiniteness rejection and are in condition for allowance.

**Double patenting.** The claims are rejected are rejected under provisional double

patenting over co-owned patents 9,034,388; 9,375,453; 9,028,877 and 9,119,864. Applicant has

filed an electronic terminal disclaimer over the listed patents.

**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:	October 12, 2016	/J. Mitchell Jones/
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John Mitchell Jones Registration No. 44,174

Casimir Jones, S.C. 2275 Deming Way, Suite 310 Middleton, WI, 53562 (608) 662-1277

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

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#### (10) International Publication Number WO 2011/050474 A1

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- (51) International Patent Classification: A61K 31/683 (2006.01) A61P 9/00 (2006.01) A61P 3/00 (2006.01)
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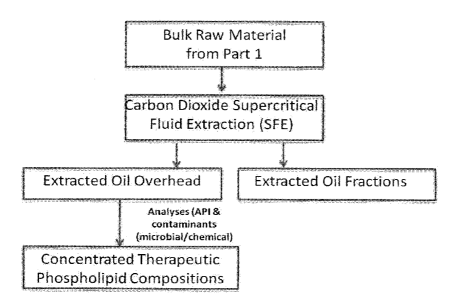
29 October 2009 (29.10.2009) 61/256,106

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[Continued on next page]

(54) Title: CONCENTRATED THERAPEUTIC PHOSPHOLIPID COMPOSITIONS



(57) Abstract: The invention relates to concentrated therapeutic phospholipid compositions; methods for treating or preventing diseases associated with cardiovascular disease, metabolic syndrome, inflammation and dieases associated therewith, neurodevelopmental diseases, and neurodegenerative diseases, comprising administering an effective amount of a concentrated therapeutic phospholipid composition.

FIGURE 1B

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#### CONCENTRATED THERAPEUTIC PHOSPHOLIPID COMPOSITIONS

#### **CROSS-REFERENCE TO RELATED APPLICATIONS**

[0001] This application claims the benefit of Provisional Application Serial No. 61/256,106, filed October 29, 2009, the contents of which are incorporated by reference in their entirety. All patents, patent applications, and publications cited herein are incorporated herein by reference in their entireties.

#### FIELD OF THE INVENTION

[0002] The invention relates to concentrated therapeutic compositions. More particularly, the invention relates to concentrated therapeutic phospholipid compositions useful for treating or preventing diseases.

#### BACKGROUND OF THE INVENTION

[0003] Genetic traits, coupled with a Western diet and lifestyle, have made cardiometabolic disorders / metabolic syndrome (MetS) a growing global epidemic. Cardiometabolic syndrome refers to a cluster of cardiovascular risk factors that include central obesity, high blood pressure, impaired glucose tolerance, hyperglycemia and dyslipidemia. Dyslipidemia is a major modifiable risk factor leading to atherosclerotic and related cardiovascular diseases (CVD), the nation's number one killer.

#### Cardiovascular Disease

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[0004] Cardiovascular disease affects one in three people in the United States during their lifetime, and accounts for nearly a third of the deaths that occur each year (Rosamond W, et al., Circulation, 115, e69–e171, (2007)). Cardiovascular diseases are defined as diseases which affect the heart or blood vessels

[0005] Statins are considered as first-line therapy for subjects at risk for CVD focusing predominantly on the reduction in low-density lipoprotein cholesterol (LDL-C or "bad cholesterol"), to recommended target levels. However statins have minimal effect in raising high-density lipoprotein cholesterol (HDL C or "good cholesterol"), now recognized as a major risk factor for developing cardiovascular disease. Treatment options to raise HDL-C are very limited and include Niaspan® (branded niacin) which is known to cause flushing

and is reported to cause hepatic enzyme abnormalities, and Tricor® (branded fenofibrates) which causes a 40% increase in LDL C and significant increase in liver enzymes, hematological changes, gall stones, pancreatitis, as well as myopathy. Some treatment options lower plasma triglycerides but have a negligible effect on HDL-C (Lovaza®). Other treatment options increase HDL-C, but are less effective on triglycerides.

10 [0006] Others have tried to increase HDL C (good cholesterol) without deleteriously affecting LDL, TG, or causing hypertension, but have not been successful. For example, torcetrapib appeared to raise HDL levels, but had no effect on TGs and LDL. However, torcetrapib caused severe hypertension and high mortality in phase III trials. Despite advancements in lowering total cholesterol, lipid abnormalities as well as other severe negative side effects still prevail. Treatment gaps in the management of dyslipidemia, considered one of the top five major modifiable risk factors of CVD, represent critical unmet medical needs. While most treatment methods only target the intrinsic LDL-C synthesis in the liver, other treatments are needed to further reduce triglycerides while increasing HDL-C and not increasing LDL-C.

20 Neurodevelopmental and neurodegenerative disease

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[0007] Neurodevelopmental and neurodegenerative diseases/disorders and neurological imbalance (in neurotransmitters) affect many people, and are defined as chronic progressive neuropathy characterized by selective and generally symmetrical loss of neurons in motor, sensory, or cognitive systems. One progressive neurodegenerative disorder, Alzheimer's disease (AD), is irreversible, and is characterized by gradual cognitive deterioration, changes in behavior and personality. These symptoms are related to neurochemical changes, neural death, and the breakdown of the inter-neural connections. Loss of short-term memory is often the first sign, followed by cognitive deficits involving multiple functions. Early stages of AD and mild cognitive impairment are characterized as milder forms of memory loss or cognitive impairment that could precede the onset of dementia and AD. Prevention of further cognitive decline in subjects with these possible precursor conditions is of paramount importance given that reversibility of AD is not possible.

[0008] It is estimated there are currently about 5.1 million people with Alzheimer's disease (AD) in the United States (Alzheimer's Association, 2007) and this number is expected to reach 13.2 million by 2050 (Hebert et al., 2003). Alzheimer's is ranked as the 7th leading cause of death in the US for people of all ages and the 5th for people aged 65 or older

5 (National Center for Health Statistics, 2004). In Canada it is 280,000 people over 65 that are estimated to have AD, and over 750,000 are expected to have the disease by 2031 (Alzheimer Society of Canada, 2006). It is estimated to 10% of all North Americans over the age of 70 years have early stage AD or mild cognitive impairment.

[0009] Alzheimer's disease is characterized by two main pathological features of the brain: intracellular neurofibrillary tangles formed by abnormal protein  $\tau$  (tau); and 10 extracellular neuritic plaques formed by  $\beta$ -amyloid peptides (A $\beta$ ) (Kuo et al., 1996). The overproduction of Aβ42 is genetically induced but environmental risk factors are required to get fully symptomatic AD (Grant et al., 2002). Among these risk factors, low docosahexaenoic acid (DHA) is one of the most important dietary risk factor for AD (Morris et al., 2005). The reasons for the impact of DHA on learning and memory and the association 15 with AD are unclear but could result from its loss in synapses (Montine et al., 2004), which are normally rich in DHA (Salem et al., 2001), where it is particularly important for postsynaptic transmission and neuroprotection (Bazan, 2003). Studies in animal models have consistently showed that brain n-3 fatty acid content is highly dependent on dietary intake and aging (Favrere et al., 2000; Youdim et al., 2000; Calon & Cole, 2007). However, some 20 reports claim higher concentrations of DHA have a deleterious effect in neurological patients.

#### Omega-3 Fatty Acids and Inflammation

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[0010] Several animal studies, has shown that increased DHA intake has been found to increase hippocampal acetycholine levels and its derivatives, neuroprotectin DI, which decreased cell death (Aid et al, 2005; Lukiw et al., 2005). A study conducted on aged mice showed that DHA intake improved memory performance (Lim et al. 2001). In another Alzheimer's disease mouse model, reduction in dietary DHA showed loss of postsynaptic proteins associated with increased oxidation, which was localized in the dendrites. However, when a group of DHA-restricted mice where given DHA, they showed signs that the DHA intake protected them against dendritic pathology, implying that DHA could be useful in preventing cognitive impairment in Alzheimer's Disease (Calon et al., 2004).

[0011] Several epidemiological studies have shown a protective effect associated with increased fish intake (a direct source of omega 3 fatty acids) against dementia and cognitive impairment decline (Kalmijin et al. 1997, Barberger-Gateau et al. 2002; Morris et al 2003). Recently, one large randomized double-blind placebo-controlled study found 1.6 g DHA and 0.7 EPA may be beneficial in reducing risk for AD (Freund-Levi et al, 2006). In addition,

there is mounting evidence that dietary supplementation with Omega 3 fatty acids may be beneficial in different psychiatric conditions such as mood behaviour, depression and dementia (Bourre et al., 2005; Peet and Stokes, 2005; Stoll et al., 1999).

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[0012] The anti-inflammatory effects of omega-3 fatty acids have been widely studied with positive results for several chronic inflammatory diseases. C-reactive protein (CRP) is a protein that increases dramatically during inflammatory processes and is commonly measured as a marker of inflammation. Greater intake of omega-3 polyunsaturated fatty acid is related to a lower prevalence of elevated CRP levels. Animal models of colitis indicate that fish oil, a natural source of omega 3 fatty acids, decreases colonic damage and inflammation. Fish oil supplements in subjects with IBD have shown to modulate levels of inflammatory mediators and may be beneficial for the induction and maintenance of remission in ulcerative colitis. In the management of RA and other inflammatory conditions, side effects limit the use of NSAIDs, such as salicylates, ibuprofen and naproxen. A clinical trial showed that 39 percent of subjects with RA supplemented with cod liver oil were able to reduce their daily NSAID requirement by greater than 30 percent. Omega-3 fatty acids have been used to reduce the risk for sudden death caused by cardiac arrhythmias.

[0013] Furthermore, omega-3 fatty acids have been shown to improve insulin sensitivity and glucose tolerance in normoglycemic men and in obese individuals. Omega-3 fatty acids have also been shown to improve insulin resistance in obese and non-obese subjects with an inflammatory phenotype. Lipid, glucose and insulin metabolism have been show to be improved in overweight hypertensive subjects through treatment with omega-3 fatty acids.

[0014] Omega-3 fatty acids can be obtained from marine organisms such as squid, fish, krill, etc. and are sold as dietary supplements. However, the uptake of omega-3 fatty acids by the body is not efficient and these raw oils contain other substances such a triglycerides and cholesterol which are known to cause deleterious side effects such as an increase in LDL-C. Certain fish oils have been developed as pharmaceutical-grade OM3-acid ethyl esters. One such OM3-acid ethyl ester is presently sold under the brand name Lovaza®. Studies have shown that Lovaza® can decrease plasma triglycerides levels in patients, however, Lovaza® has a negligible effect on raising good cholesterol (HDL-C). AMR101 is another ethyl ester

form of OM3 fatty acids based on EPA with little or no DHA that is presently in clinical trials. AMR101 also appears to decrease triclycerides but also has a negligible effect on raising HDL-C.

[0015] A phospholipid composition of OM3 fatty acids has been disclosed in US 2004/0234587. This phospholipid composition has OM3 fatty acids esterified to the phospholipid. This phospholipid composition is reported to be at a concentration of about 40% phospholipids (w/w composition) and contains high concentrations of triglycerides (about 45%) and free fatty acids (about 15%). When tested in subjects, this composition demonstrated very little effect on lowering triglyceride plasma levels (less than 11% reduction).

15 [0016] Marine oil compositions comprising free fatty acids and lipids, including OM3 fatty acids and phospholipids, have been disclosed in WO 2000/23546, however the compositions do not disclose OM3 fatty acids esterified to diglycerol phosphate and have very high concentrations of triglycerides and free fatty acids, and for these reasons would not be expected to reduce triglycerides even to the level of the composition disclosed in US 2004/0234587, described above.

[0017] Therefore, new forms of omega-3 fatty acids are needed that are usefulefor treating or preventing disease. Described herein are novel concentrated therapeutic phospholipid compositions, as well as pharmaceutical compositions comprising same, and methods of their use.

#### SUMMARY OF THE INVENTION

[0018] Accordingly, in one aspect concentrated therapeutic phospholipid compositions are described, the compositions comprising compounds of the Formula I:

$$H_2C \longrightarrow O \longrightarrow R_1$$
 $R_2 \longrightarrow O \longrightarrow CH$ 
 $H_2C \longrightarrow O \longrightarrow P \longrightarrow O \longrightarrow X$ 
 $O$ 

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wherein for each compound of Formula I in the compositions

each  $R_1$  is independently selected from hydrogen or any fatty acid;

each  $R_2$  is independently selected from hydrogen or any fatty acid;

wherein at least one of R<sub>1</sub> and R<sub>2</sub> in each compound of Formula I is a fatty acid; and

each X is independently selected from -CH<sub>2</sub>CH<sub>2</sub>NH<sub>3</sub>, -CH<sub>2</sub>CH<sub>2</sub>N(CH<sub>3</sub>)<sub>3</sub> or

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wherein the total amount of the compounds of Formula I in the composition being at a concentration of between 45% (w/w) to about 99% (w/w).

[0019] In some embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition are in a concentration of between about 45% (w/w (phospholipids/total composition)) up to 70% (w/w (phospholipids/total composition)). In still further embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition are in a concentration of between about 50% (w/w
(phospholipids/total composition)) up to 70% (w/w (phospholipids/total composition)). In other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition are in a concentration of between about 60% (w/w (phospholipid/total composition)) up to 70% (w/w (phospholipids/total composition)). In still other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition are in a concentration of about 66% (w/w (phospholipids/total composition)).

[0020] In other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition are in a concentration of above 70% (w/w (phospholipids/total composition)) to about 99% (w/w (phospholipids/total composition)). In still other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipids/total composition) to about 98% (w/w (phospholipids/total composition)). In still other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition are in a concentration of between about 85% (w/w (phospholipids/total composition)) to about 95% (w/w (phospholipids/total composition)). In further embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition are in a concentration of about 90% (w/w (phospholipids/total composition)).

[0021] In some embodiments, R1 is a monounsaturated fatty acid. In other embodiments, R1 is a polyunsaturated fatty acid. In some embodiments, R2 is a monounsaturated fatty acid. In other embodiments, R2 is a polyunsaturated fatty acid. In other embodiments, the polyunsaturated fatty acid is an omega 3 fatty acid. In still other embodiments, both R1 and R2 are each independently selected from an omega 3 fatty acid. When at least one of R1 and R2 is an omega 3 fatty acid, the concentrated therapeutic phospholipid composition comprising compounds of Formula I is known as an OM3:PL.

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In other embodiments, R1 is docosahexaenoic acid (DHA). In other embodiments, R2 is a monounsaturated fatty acid and R1 is DHA. In other embodiments, R2 is a polyunsaturated fatty acid and R1 is DHA. In other embodiments, R2 is an omega 3 fatty acid and R1 is DHA. In still further embodiments, R2 is EPA and R1 is DHA. In still further embodiments, R2 is DHA and R1 is DHA.

- 10 [0023] In other embodiments, R1 is eicosapentaenoic acid (EPA). In other embodiments, R2 is a monounsaturated fatty acid and R1 is EPA. In other embodiments, R2 is a polyunsaturated fatty acid and R1 is EPA. In other embodiments, R2 is an omega 3 fatty acid and R1 is EPA. In still further embodiments, R2 is DHA and R1 is EPA. In still further embodiments, R2 is EPA and R1 is EPA.
- 15 [0024] In another embodiment, R2 is DHA. In other embodiments, R1 is a monounsaturated fatty acid and R2 is DHA. In other embodiments, R1 is a polyunsaturated fatty acid and R2 is DHA. In other embodiments, R1 is an omega 3 fatty acid and R2 is DHA.
  - [0025] In other embodiments, R2 is EPA. In other embodiments, R1 is a monounsaturated fatty acid and R2 is EPA. In other embodiments, R1 is a polyunsaturated fatty acid and R2 is EPA. In other embodiments, R1 is an omega 3 fatty acid and R2 is EPA.

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[0026] In some embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition have predominantly DHA at the R2 position of Formula I. In other embodiments, there is more DHA in the compounds of Formula I in the concentrated therapeutic phospholipid composition than EPA. In some embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition have greater than 60% DHA. In other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition have greater than 70% DHA. In other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition have greater than 80% DHA. In other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition have greater than 90% DHA. In other embodiments, the compounds of Formula I in the concentrated therapeutic phospholipid composition have greater than 95% DHA.

In some embodiments, there are free fatty acids in the concentrated therapeutic phospholipid composition, in addition to the fatty acids esterified to the phosphate. In other embodiments, there are essentially no free fatty acids (also expressed as 0% free fatty acids (or FFA)) in the concentrated therapeutic phospholipid composition.

- [0028] In other embodiments, the ratio of the total amount of DHA to EPA in the concentrated therapeutic phospholipid composition is between about 1:1 and 1:0.1. In some embodiments, the ratio is between about 1:0.7 and about 1:0.3. In other embodiments, the ratio is about 1:0.5.
  - [0029] In some embodiments, the ratio of the total amount of EPA to DHA in the compounds of Formula I in the concentrated therapeutic phospholipid composition is between about 1:1 and 1:0.1. In some embodiments, the ratio is between about 1:0.7 and about 1:0.3. In other embodiments, the ratio is about 1:0.5.

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- [0030] In some embodiments, the total amount of OM3 fatty acids in the concentrated therapeutic phospholipid composition is between about 20% and about 50%. In other embodiments, the total amount of OM3 fatty acids in the concentrated therapeutic phospholipid composition is between about 30% and about 45%. In other embodiments, the total amount of OM3 fatty acids in the concentrated therapeutic phospholipid composition is about 40%.
- [0031] In some embodiments, the total amount of DHA in the concentrated therapeutic phospholipid composition is between about 5% and 20%. In some embodiments, the total amount of DHA in the concentrated therapeutic phospholipid composition is between about 10% and 15%. In some embodiments, the total amount of DHA in the concentrated therapeutic phospholipid composition is about 14%.
- [0032] In some embodiments, the total amount of EPA in the concentrated therapeutic phospholipid composition is between about 10% and 30%. In some embodiments, the total amount of DHA in the concentrated therapeutic phospholipid composition is between about 15% and 25%. In some embodiments, the total amount of EPA in the concentrated therapeutic phospholipid composition is about 22%.

5 [0033] In some embodiments, X is -CH<sub>2</sub>CH<sub>2</sub>NH<sub>3</sub>. In other embodiments, X is -CH<sub>2</sub>CH<sub>2</sub>N(CH<sub>3</sub>)<sub>3</sub>. In some embodiments, X is

[0034] In some embodiments, the concentrated therapeutic phospholipid composition comprises predominantly phospholipids containing -CH<sub>2</sub>CH<sub>2</sub>N(CH<sub>3</sub>)<sub>3</sub> (also known as a phosphoyidyl-N-trimethylethanolamine).

[0035] In other embodiments, the concentrated therapeutic phospholipid composition further comprises an antioxidant. In some embodiments, the antioxidant is a carotenoid. In other embodiments, the carotenoid is pro-vitamin A. In other embodiments, the antioxidant is a flavonoid. In other embodiments, the flavonoid is selected from naringin, naringenin, hesperetin/kaempferol, rutin, luteolin, neohesperidin, quecertin. In other embodiments, the flavonoid is

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20 **[0036]** In some embodiments, the concentration of the flavonoid is between about 1 mg/kg (w/w of composition) and about 20 mg/kg (w/w of composition). In other embodiments, the concentration of the flavonoid is greater than about 10 mg/kg (w/w of composition).

5 [0037] In further embodiments, the concentrated therapeutic phospholipid composition has a concentration of astaxanthin greater than 2000 mg/kg (w/w of composition). In still other embodiments, the concentration of astaxanthin is between about 2,000 mg/kg (w/w of composition) and about 5,500 mg/kg (w/w of composition).

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[0038] In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration below about 22% (w/w of composition). In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration below about 15% (w/w of composition). In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentrated therapeutic phospholipid composition has a free fatty acid concentrated therapeutic phospholipid composition has a free fatty acid concentrated therapeutic phospholipid composition has a free fatty acid concentrated therapeutic phospholipid composition has a free fatty acid concentration of about 1% (w/w of composition). In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration below 1% (w/w of composition). In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration below 1% (w/w of composition). In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration of 0% (w/w of composition).

[0039] In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration of between about 1% (w/w) and about 20% (w/w). In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration of between about 5% (w/w) and about 17% (w/w). In some embodiments, the concentrated therapeutic phospholipid composition has a free fatty acid concentration of between about 10% (w/w) and about 15% (w/w).

[0040] In some embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentration between about 0% (w/w) and about 30% (w/w). In other embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentration between about 5% and about 20%. In still further embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentration between about 10% and about 15%.

In some embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentration below about 15%. In some embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentration below about 10%. In some embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentrated therapeutic phospholipid composition has a triglyceride concentrated therapeutic phospholipid composition has a triglyceride concentration below 1%. In some embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentration below 1%. In some embodiments, the concentrated therapeutic phospholipid composition has a triglyceride concentration of about 0%.

[0042] In other embodiments, the concentrated therapeutic phospholipid composition comprises at least 50% compounds of Formula I (w/w), wherein at least 15% of the fatty acid content is EPA, at least 9 % of the fatty acid content is DHA, and at least 0.1% astaxanthin (w/w). In other embodiments, the concentrated therapeutic phospholipid composition comprises at least 66% compounds of Formula I (w/w), wherein at least 20% of the fatty acid content is EPA, at least 12 % of the fatty acid content is DHA, and at least 0.4% astaxanthin (w/w). In other embodiments, the concentrated therapeutic phospholipid composition comprises at least 90% compounds of Formula I (w/w), at least 22% of the fatty acid content is EPA, at least 12% of the fatty acid content is DHA, and 0.4% astaxanthin (w/w).

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[0043] In other embodiments, the concentrated therapeutic phospholipid composition comprises at least 50% compounds of Formula I (w/w composition), wherein at least 15% of the fatty acid content is EPA, at least 9 % of the fatty acid content is DHA. In other embodiments, the concentrated therapeutic phospholipid composition comprises at least 66% compounds of Formula I (w/w), wherein at least 20% of the fatty acid content is EPA, at least 12 % of the fatty acid content is DHA. In other embodiments, the concentrated therapeutic phospholipid composition comprises above 70% compounds of Formula I(w/w), wherein at least 22% of the fatty acid content is EPA, at least 12% of the fatty acid content is DHA. In other embodiments, the concentrated therapeutic phospholipid composition comprises above 90% compounds of Formula I(w/w), wherein at least 22% of the fatty acid content is EPA, at least 12% of the fatty aci

In one aspect, a concentrated therapeutic phospholipid composition is described comprising compounds of Formula at a concentration of about 66% (w/w (phospholipids/total composition) a free fatty acid (FFA) concentration of less than 6% (w/w FFA/total composition) and a triglyceride concentration of about 0%, the composition being useful for treating and preventing cardiometabolic disorders / metabolic syndrome. In some embodiments, 1 g of the concentrated therapeutic phospholipid composition comprises about 387 mg of total OM3 fatty acids wherein EPA is at about 215 mg and DHA is at about 136 mg) and astaxanthine at about 5 mg.

[0045] In one aspect, a concentrated therapeutic phospholipid composition is described comprising compounds of Formula at a concentration of above 70% (w/w (phospholipids/total composition), a free fatty acid (FFA) concentration of about 0% and a triglyceride concentration of about 0%, the composition being useful for treating and preventing neurodegenerative and neurodevelopmental disorders and diseases.

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[0046] In one aspect, the invention is based in part on the unexpected and surprising discovery that concentrated therapeutic phospholipid compositions are useful in modulating plasma triglyceride levels as well as plasma HDL C levels, while not elevating LDL C levels. This unexpected and surprising discovery is useful in the treatment or prevention of disorders associated with increased triglyceride levels, increased LDL-C levels and decreased HDL-C levels. Such diseases and disorders inlude but are not limited to cardiometabolic disorders / metabolic syndrome (MetS), neurodevelopmental and neurodegenerative diseases/disorders, and inflammation disorders.

[0047] In another aspect, a method of treating or preventing a cardiometabolic disorder / metabolic syndrome is described, the method comprising administering to a subject in need thereof a concentrated therapeutic phospholipid composition. In some embodiments, the cardiometabolic disorder is selected from atherosclerosis, arteriosclerosis, coronary heart (carotid artery) disease (CHD or CAD), acute coronary syndrome (or ACS), valvular heart disease, aortic and mitral valve disorders, arrhythmia / atrial fibrillation, cardiomyopathy and heart failure, angina pectoris, acute myocardial infarction (or AMI), hypertension, orthostatic hypotension, shock, embolism (pulmonary and venous), endocarditis, diseases of arteries, the aorta and its branches, disorders of the peripheral vascular system (peripherial arterial disease or PAD), Kawasaki disease, congenital heart disease (cardiovascular defects) and stroke (cerebrovascular disease), dyslipidemia, hypertriglyceridemia, hypertension, heart failure,

5 cardiac arrhythmias, low HDL levels, high LDL levels, stable angina, coronary heart disease, acute myocardial infarction, secondary prevention of myocardial infarction, cardiomyopathy, endocarditis, type 2 diabetes, insulin resistance, impaired glucose tolerance, hypercholesterolemia, stroke, hyperlipidemia, hyperlipoprotenemia, chronic kidney disease, intermittent claudication, hyperphosphatemia, omega-3 deficiency, phospholipid 10 deficiency, carotid atherosclerosis, peripheral arterial disease, diabetic nephropathy, hypercholesterolemia in HIV infection, acute coronary syndrome (ACS), non-alcoholic fatty liver disease/non-alcoholic steatohepatitis (NAFLD/NASH), arterial occlusive diseases, cerebral atherosclerosis, arteriosclerosis, cerebrovascular disorders, myocardial ischemia, coagulopathies leading to thrombus formation in a vessel and diabetic autonomic neuropathy. 15 In some instances, the methods described above for treating or preventing a cardiometabolic disorder / metabolic syndrome may utilize concentrated therapeutic phospholipid compositions having a concentration of 66% (w/w (phospholipids/composition)).

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[0048] In another aspect, methods of treating, preventing, or improving cognition and /or a cognitive disease, disorder or impairment (memory, concentration, learning (deficit)), or of treating or preventing neurodegenerative disorders are described, the method comprising administering to a subject in need thereof a concentrated therapeutic phospholipid composition. In some embodiments, the cognitive disease, disorder or impairment is selected from Attention Deficit Disorder (ADD), Attention Deficit Hyperactivity Disorder (ADHD), autism/autism spectrum disorder (ASD), (dyslexia, age-associated memory impairment and learning disorders, amnesia, mild cognitive impairment, cognitively impaired non-demented, pre-Alzheimer's disease, Alzheimer's disease, epilepsy, Pick's disease, Huntington's disease, Parkinson disease, Lou Gehrig's disease, pre-dementia syndrome, Lewy body dementia dementia, dentatorubropallidoluysian atrophy, Freidreich's ataxia, multiple system atrophy, types 1, 2, 3, 6, 7 spinocerebellar ataxia, amyotrophic lateral sclerosis, familial spastic paraparesis, spinal muscular atrophy, spinal and bulbar muscular atrophy, age-related cognitive decline, cognitive deterioration, moderate mental impairment, mental deterioration as a result of ageing, conditions that influence the intensity of brain waves and/or brain glucose utilization, stress, anxiety, concentration and attention impairment, mood deterioration, general cognitive and mental well being, neurodevelopmental, neurodegenerative disorders, hormonal disorders, neurological imbalance or any combinations thereof. In a specific embodiment, the cognitive disorder is memory impairment. In some instances, the methods described above for treating, preventing, or

improving cognition and /or a cognitive disease, disorder or impairment (memory, concentration, learning (deficit)), or of treating or preventing neurodegenerative disorders may utilize concentrated therapeutic phospholipid compositions having a concentration of greater than 70% (w/w (phospholipids/composition)).

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[0049] In another aspect, a method for inhibiting, preventing, or treating inflammation or an inflammatory disease is described, the method comprising administering to a subject in need thereof, a concentrated therapeutic phospholipid composition. In some embodiments, the inflammation or inflammatory disease is selected from organ transplant rejection; reoxygenation injury resulting from organ transplantation (see Grupp et al., J. Mol. Cell Cardiol. 31: 297-303 (1999)) including, but not limited to, transplantation of the following organs: heart, lung, liver and kidney; chronic inflammatory diseases of the joints, including arthritis, rheumatoid arthritis, osteoarthritis and bone diseases associated with increased bone resorption; inflammatory bowel diseases (IBD) such as ileitis, ulcerative colitis (UC), Barrett's syndrome, and Crohn's disease (CD); inflammatory lung diseases such as asthma, acute respiratory distress syndrome (ARDS), and chronic obstructive pulmonary disease (COPD); inflammatory diseases of the eye including corneal dystrophy, trachoma, onchocerciasis, uveitis, sympathetic ophthalmitis and endophthalmitis; chronic inflammatory diseases of the gum, including gingivitis and periodontitis; inflammatory diseases of the kidney including uremic complications, glomerulonephritis and nephrosis; inflammatory diseases of the skin including sclerodermatitis, psoriasis and eczema; inflammatory diseases of the central nervous system, including chronic demyelinating diseases of the nervous system, multiple sclerosis, AIDS-related neurodegeneration and Alzheimer's disease, infectious meningitis, encephalomyelitis, Parkinson's disease, Huntington's disease, Epilepsy, amyotrophic lateral sclerosis and viral or autoimmune encephalitis, preeclampsia; chronic liver failure, brain and spinal cord trauma, and cancer. The inflammatory disease can also be a systemic inflammation of the body, exemplified by gram-positive or gram negative shock, hemorrhagic or anaphylactic shock, or shock induced by cancer chemotherapy in response to proinflammatory cytokines, e.g., shock associated with proinflammatory cytokines. Such shock can be induced, e.g., by a chemotherapeutic agent that is administered as a treatment for cancer. Other disorders include depression, obesity, allergic diseases, acute cardiovascular events, muscle wasting diseases, and cancer cachexia. Also inflammation that results from surgery and trauma can be treated with the concentrated therapeutic phospholipid compositions.

5 [0050] The details of the invention are set forth in the accompanying description below. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, illustrative methods and materials are now described. Other features, objects, and advantages of the invention will be apparent from the description and from the claims. In the specification and the appended claims, the singular forms also include the plural unless the context clearly dictates otherwise. Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. All patents and publications cited in this specification are incorporated herein by reference in their entireties.

#### **DESCRIPTION OF THE FIGURES**

- 15 **[0051]** FIG. 1A depicts a flow chart for the process for making the concentrated therapeutic phospholipid compositions.
  - [0052] FIG. 1B depicts a flow chart for the process for making the concentrated therapeutic phospholipid compositions.
  - [0053] FIG. 1C depicts shows the schematic of the supercritical CO2 extraction apparatus.

- [0054] FIG. 2 depicts circulating plasma triglyceride concentration of C57BL/6 mice treated with Composition 3.
- [0055] FIG. 3 depicts circulating plasma HDL-Cholesterol concentration of C57BL/6 mice treated with Composition 3.
- 25 [0056] FIG. 4 depicts circulating plasma percentage of HDL-Cholesterol in C57BL/6 mice treated with Composition 3.
  - [0057] FIG. 5 depicts circulating plasma LDL-Cholesterol concentration of C57BL/6 mice treated with Composition 3.
- [0058] FIG. 6 depicts circulating plasma percentage of LDL-Cholesterol in C57BL/630 mice treated with Composition 3.
  - [0059] FIG. 7 depicts circulating plasma NEFA concentration of C57BL/6 mice treated with Composition 3.
  - [0060] FIG. 8 depicts circulating plasma Glucose concentration of C57BL/6 mice treated with Composition 3.
- 35 [0061] FIG. 9 depicts circulating plasma Phospholipid concentration of C57BL/6 mice treated with Composition 3.

5 [0062] FIG. 10 depicts circulating plasma ALT concentration of C57BL/6 mice treated with Composition 3.

- [0063] FIG. 11 depicts liver Total Cholesterol concentration of C57BL/6 mice treated with Composition 3.
- [0064] FIG. 12 depicts liver Triglyceride concentration of C57BL/6 mice treated with Composition 3.
  - [0065] FIG. 13 depicts circulating plasma triglyceride concentration of LDLr KO mice treated with Composition 3.
  - [0066] FIG. 14 depicts circulating plasma HDL-Cholesterol concentration of LDLr KO mice treated with Composition 3.
- 15 **[0067]** FIG. 15 depicts circulating plasma percentage of HDL-Cholesterol in LDLr KO mice treated with Composition 3.
  - [0068] FIG. 16 depicts liver Total Cholesterol concentration of LDLr KO mice treated with Composition 3.
  - [0069] FIG. 17 depicts liver Triglyceride concentration of LDLr KO mice treated with Composition 3.

- [0070] FIG. 18 depicts circulating plasma Triglyceride concentration of ApoA-1 CET Tg mice treated with Composition 3.
- [0071] FIG. 19 depicts circulating plasma total cholesterol concentration of adult male SD, ZDF, SHR and JCR:LA rats.
- 25 **[0072]** FIG. 20 depicts circulating plasma total cholesterol concentration of adult male SD, ZDF, SHR and JCR:LA rats.
  - [0073] FIG. 21 depicts circulating plasma HDL/LDL concentration of adult male SD, ZDF, SHR and JCR:LA rats.
- [0074] FIG. 22 depicts circulating plasma total cholesterol/HDL concentration of adult male SD, ZDF, SHR and JCR:LA rats.
  - [0075] FIG. 23 depicts prothrombin time of adult male SD, ZDF, SHR and JCR:LA rats.
  - [0076] FIG. 24 depicts OGTT area under the curve data in ZDF male rats treated with Composition 3 for 28 days.
- [0077] FIG. 25 depicts OGTT area under the curve data in ZDF male rats treated with Composition 3 for 28 days.
  - [0078] FIG. 26 depicts OGTT area under the curve data in SD male rats treated with Composition 3 for 28 days.

5 [0079] FIG. 27 depicts OGTT area under the curve data in ZDF male rats treated with Composition 3 for 28 days.

- [0080] FIG. 28 depicts the effects of Composition 3 on plasma total cholesterol in male ZDF rats compared to age-matched controls.
- [0081] FIG. 29 depicts the effects of Composition 3 on plasma HDL-cholesterol in male ZDF rats compared to age-matched controls.
  - [0082] FIG. 30 depicts the effects of Composition 3 on plasma triglycerides in male ZDF rats compared to age-matched controls.
  - [0083] FIG. 31 depicts the effects of Composition 3 on glucose intolerance in male ZDF rats.
- 15 [0084] FIG. 32 depicts the effects of Composition 3 on glucose intolerance in male ZDF rats.
  - [0085] FIG. 33 depicts the effects of Composition 3 on glucose intolerance in male SD rats.
  - [0086] FIG. 34 depicts the effects of Composition 3 on glucose intolerance in male SD rats.
  - [0087] FIG. 35 depicts the comparative effects of Composition 3 and Lovaza (R) on the Omega-3 Index.

### **DETAILED DESCRIPTION OF THE INVENTION**

[0088] It has been unexpectedly discovered that concentrated therapeutic phospholipid compositions demonstrate surprising effects in the treatment of metabolic disorders, cardiovascular disease, neurodevelopmental disorders and neurodegenerative diseases, and inflammation disorders.

#### **DEFINITIONS**

- [0089] The following definitions are used in connection with the concentrated therapeutic phospholipid compositions:
  - [0090] The term "concentrated therapeutic phospholipid composition" and "concentrated therapeutic phospholipid compositions" as used herein refer to the concentrated therapeutic phospholipid compositions comprising compounds of Formula I.

5 [0091] The articles "a" and "an" are used in this disclosure to refer to one or more than one (i.e., to at least one) of the grammatical object of the article. By way of example, "an element" means one element or more than one element.

[0092] The term "and/or" is used in this disclosure to mean either "and" or "or" unless indicated otherwise.

10 **[0093]** The term "about" when used in this disclosure along with a recited value means the value recited and includes the range of + or -5% of the value. For example, the phrase about 80% means 80% and + or -5% of 80, i.e. 76% to 84%. The recited value "about 0%" as used herein means that the detectable amount is less than one part per thousand.

[0094] The term "fatty acid" or "fatty acid residue" as used herein means a carboxylic acid with a long unbranched aliphatic chain, which is either saturated or unsaturated. Saturated fatty acids have the general formula C<sub>n</sub>H<sub>2n</sub>+1 COOH. Examples of saturated fatty acids include but are not limited to: Propanoic acid, Butanoic acid, Pentanoic acid, Hexanoic acid, Heptanoic acid, Octanoic acid, Nonanoic acid, Decanoic acid, Undecanoic acid, Dodecanoic acid, Tridecanoic acid, Tetradecanoic acid, Pentadecanoic acid, Hexadecanoic acid, Heptadecanoic acid, Octadecanoic acid, Nonadecanoic acid, Eicosanoic acid,

Heneicosanoic acid, Docosanoic acid, Tricosanoic acid, Tetracosanoic acid, Pentacosanoic acid, Hexacosanoic acid, Heptacosanoic acid, Octacosanoic acid, Nonacosanoic acid, Triacontanoic acid, Henatriacontanoic acid, Dotriacontanoic acid, Tritriacontanoic acid, Tetratriacontanoic acid, Pentatriacontanoic acid, Hexatriacontanoic acid. An unsaturated fat

is a fat or fatty acid in which there are one or more double bonds in the fatty acid chain. A fat molecule is monounsaturated if it contains one double bond, and polyunsaturated if it contains more than one double bond. Examples of unsaturated fatty acids include but are not limited to: Myristoleic acid, Palmitoleic acid, Sapienic acid, Oleic acid, Linoleic acid, α-Linolenic acid, Arachidonic acid, Eicosapentaenoic acid (EPA), Erucic acid,

30 Docosahexaenoic acid (DHA), and Docosapentaenoic acid.

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[0095] A "subject" is a mammal, e.g., a human, mouse, rat, guinea pig, dog, cat, horse, cow, pig, or non-human primate, such as a monkey, chimpanzee, baboon or rhesus.

[0096] Representative "pharmaceutically acceptable salts" include, e.g., water-soluble and water-insoluble salts, such as the acetate, amsonate (4,4-diaminostilbene-2, 2 -

disulfonate), benzenesulfonate, benzonate, bicarbonate, bisulfate, bitartrate, borate, bromide, butyrate, calcium, calcium edetate, camsylate, carbonate, chloride, citrate, clavulariate, dihydrochloride, edetate, edisylate, estolate, esylate, fiunarate, gluceptate, gluconate, glutamate, glycollylarsanilate, hexafluorophosphate, hexylresorcinate, hydrabamine,

hydrobromide, hydrochloride, hydroxynaphthoate, iodide, isothionate, lactate, lactobionate, laurate, magnesium, malate, maleate, mandelate, mesylate, methylbromide, methylnitrate, methylsulfate, mucate, napsylate, nitrate, N-methylglucamine ammonium salt, 3-hydroxy-2-naphthoate, oleate, oxalate, palmitate, pamoate (1,1-methene-bis-2-hydroxy-3-naphthoate, einbonate), pantothenate, phosphate/diphosphate, picrate, polygalacturonate, propionate,
 p-toluenesulfonate, salicylate, stearate, subacetate, succinate, sulfate, sulfosalicylate, suramate, tannate, tartrate, teoclate, tosylate, triethiodide, and valerate salts.

[0097] The term "metabolic disorder" as used herein refers to disoders, diseases and syndromes involving dyslipidemia, and the terms metabolic disorder, metabolic disease, and metabolic syndrome are used interchangeably herein.

[0098] An "effective amount" when used to describe an amount of a concentrated therapeutic phospholipid composition useful for treating or preventing a disease or disorder, is an amount that is efficacious with respect to the disease or disorder connected with that particular effective amount.

[0099] The term "carrier", as used in this disclosure, encompasses carriers, excipients, and diluents and means a material, composition or vehicle, such as a liquid or solid filler, diluent, excipient, solvent or encapsulating material, involved in carrying or transporting a pharmaceutical agent from one organ, or portion of the body, to another organ, or portion of the body.

[0100] The term "treating", with regard to a subject, refers to improving at least one symptom of the subject's disorder. Treating can be curing, improving, or at least partially ameliorating the disorder.

[0101] The term "disorder" is used in this disclosure to mean, and is used interchangeably with, the terms disease, condition, or illness, unless otherwise indicated.

[0102] The term "administer", "administering", or "administration" as used in this disclosure refers to either directly administering a compound or pharmaceutically acceptable salt of the compound or a composition to a subject, or administering a prodrug derivative or analog of the compound or pharmaceutically acceptable salt of the compound or composition to the subject, which can form an equivalent amount of active compound within the subject's body.

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5 Methods of Making the Concentrated Therapeutic Phospholipid Compositions The concentrated therapeutic phospholipid compositions can be made or produced [0103] by any method known to one of skill in the art. For example, phospholipid containing oils can be isolated from natural sources (see US 2004/0234587, US 2009/0074857, and US 2008/0274203, the disclosures of which are incorporated by reference in their entireties), 10 which can then be further processed. Alternatively, following the process outlined in Figure la results in bulk raw material krill oil ready for further processing. These phospholipid containing oils can be further processed using countercurrent supercritical CO<sub>2</sub> extraction (Lucien, F. P., et al., Australas Biotechnol. 1993, 3, 143-147) to concentrate the compositions to produce the concentrated therapeutic phospholipid compositions described herein (see Figure 1b). For example, countercurrent supercritical CO<sub>2</sub> extraction at 70 C and 30 MPa 15 and with a CO<sub>2</sub>/oil ratio of 72 can be used to remove certain biomolecules such as all triglycerides from the bulk raw material krill oil as well as some of the free fatty acids (Figure 1b). As more of the TGs and FFAs are removed from the bulk raw material krill oil, the concentration of the phospholipids increases. When the TGs have been removed through 20 this process the phospholipid composition is at about 66% concentration (w/w (phospholipids/composition)) and contains less than 5% free fatty acids (w/w). As more of the FFAs are removed using this process, a concentrated therapeutic phospholipid composition results having a phospholipid concentration above 70% up to about 90% (w/w (phospholipids/composition)) having about 1% or less TG and about 0% FFA. Other aquatic 25 and/or marine biomasses may be used as starting materials, such as, for example, squid or blue mussels. Additional components can be added before, during, or after processing. Alternatively, phospholipids can be synthesized; a typical way to synthesize would be, among others, according to the procedure described in US 7,034,168, the disclosure of which is incorporated herein its entirety.

30 Methods for using the Concentrated Therapeutic Phospholipid Compositions

[0104] Described herein are methods of reducing circulating plasma concentrations of triglycerides, LDL-cholesterol, total cholesterol and NEFA, the method comprising administering to a subject in need thereof an effective amount of a Compsition of the Invention.

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5 [0105] Also provided are methods of increasing plasma concentrations of HDL-cholesterol and hepatic concentrations of triglycerides and total cholesterol, the method comprising administering to a subject in need thereof an effective amount of a concentrated therapeutic phospholipid composition.

- 10 **[0106]** In another aspect, a method of reducing TG without the risk of increasing LDL is described, the method comprising administering to a subject in need thereof, a concentrated therapeutic phospholipid composition.
- [0107] Also provided are methods for inhibiting, preventing, or treating a metabolic disorder, or symptoms of a metabolic disease, in a subject, the method comprising 15 administering to a subject in need thereof an effective amount of a concentrated therapeutic phospholipid composition. Examples of such disorders include, but are not limited to atherosclerosis, dyslipidemia, hypertriglyceridemia, hypertension, heart failure, cardiac arrhythmias, low HDL levels, high LDL levels, stable angina, coronary heart disease, acute 20 myocardial infarction, secondary prevention of myocardial infarction, cardiomyopathy, endocarditis, type 2 diabetes, insulin resistance, impaired glucose tolerance, hypercholesterolemia, stroke, hyperlipidemia, hyperlipoprotenemia, chronic kidney disease, intermittent claudication, hyperphosphatemia, carotid atherosclerosis, peripheral arterial disease, diabetic nephropathy, hypercholesterolemia in HIV infection, acute coronary 25 syndrome (ACS), non-alcoholic fatty liver disease, arterial occlusive diseases, cerebral atherosclerosis, arteriosclerosis, cerebrovascular disorders, myocardial ischemia, and diabetic autonomic neuropathy.
- [0108] Also provided are methods for inhibiting, preventing, or treating inflammation or an inflammatory disease in a subject. The inflammation can be associated with an inflammatory disease. Inflammatory diseases can arise where there is an inflammation of the body tissue. These include local inflammatory responses and systemic inflammation. Examples of such diseases include, but are not limited to: organ transplant rejection; reoxygenation injury resulting from organ transplantation (see Grupp et al., J. Mol. Cell
   Cardiol. 31: 297-303 (1999)) including, but not limited to, transplantation of the following organs: heart, lung, liver and kidney; chronic inflammatory diseases of the joints, including arthritis, rheumatoid arthritis, osteoarthritis and bone diseases associated with increased bone resorption; inflammatory bowel diseases such as ileitis, ulcerative colitis, Barrett's syndrome,

5 and Crohn's disease; inflammatory lung diseases such as asthma, adult respiratory distress syndrome, and chronic obstructive airway disease; inflammatory diseases of the eye including corneal dystrophy, trachoma, onchocerciasis, uveitis, sympathetic ophthalmitis and endophthalmitis; chronic inflammatory diseases of the gum, including gingivitis and periodontitis; inflammatory diseases of the kidney including uremic complications, 10 glomerulonephritis and nephrosis; inflammatory diseases of the skin including sclerodermatitis, psoriasis and eczema; inflammatory diseases of the central nervous system, including chronic demyelinating diseases of the nervous system, multiple sclerosis, AIDSrelated neurodegeneration and Alzheimer's disease, infectious meningitis, encephalomyelitis, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis and viral or autoimmune encephalitis. Metabolic disease such as type II diabetes mellitus; the prevention 15 of type I diabetes; dyslipedemia; diabetic complications, including, but not limited to glaucoma, retinopathy, nephropathy, such as microaluminuria and progressive diabetic nephropathy, polyneuropathy, atherosclerotic coronary arterial disease, peripheral arterial disease, nonketotic hyperglycemichyperosmolar coma, mononeuropathies, autonomic 20 neuropathy, joint problems, and a skin or mucous membrane complication, such as an infection, a shin spot, a candidal infection or necrobiosis lipoidica diabeticorum; immunecomplex vasculitis, systemic lupus erythematosus; inflammatory diseases of the heart such as cardiomyopathy, ischemic heart disease hypercholesterolemia, and atherosclerosis; as well as various other diseases that can have significant inflammatory components, including preeclampsia; chronic liver failure, brain and spinal cord trauma, and cancer. The 25 inflammatory disease can also be a systemic inflammation of the body, exemplified by grampositive or gram negative shock, hemorrhagic or anaphylactic shock, or shock induced by cancer chemotherapy in response to proinflammatory cytokines, e.g., shock associated with proinflammatory cytokines. Such shock can be induced, e.g., by a chemotherapeutic agent 30 that is administered as a treatment for cancer. Other disorders include depression, obesity, allergic diseases, acute cardiovascular events, muscle wasting diseases, and cancer cachexia. Also inflammation that results from surgery and trauma can be treated with the concentrated therapeutic phospholipid compositions.

5 **[0109]** Also provided are methods for inhibiting, preventing, or treating hypertriglyceridemia in subject. In some embodiments, the hypertriglyceridemia is moderate hypertriglyceridemia. In some embodiments, the subject is diagnosed with moderate hypertriglyceridemia. Moderate hypertriglyceridemia is defined as a subject having a TG level of > 3.9 mmol/L (>350 mg/dL).

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**[0110]** Also provided are methods for reducing fasting plasma levels of Low-density Lipoprotein Cholesterol (LDL-C) in a subject. In some embodiments of reducing fasting plasma levels of Low-density Lipoprotein Cholesterol (LDL-C), the subject is diagnosed with moderate hypertriglyceridemia.

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[0111] Also provided are methods for increasing fasting plasma levels of High-density Lipoprotein Cholesterol (HDL-C) in a subject. In some embodiments of increasing fasting plasma levels of High-density Lipoprotein Cholesterol (HDL-C), the subject is diagnosed with moderate hypertriglyceridemia.

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[0112] Also provided are methods for increasing the Omega-3 index (OM3I) in a subject. The Omega-3 Index is defined as the percentage of EPA+DHA in red blood cells (RBC) which can be represented by the formula: OM3I = (EPA + DHA) / Total fatty acids in RBC. Low levels of EPA+DHA in erythrocytes are associated with increased risk for sudden cardiac death and can be viewed as a marker of increased risk (an actual risk factor) for death from coronary heart disease (Harris, 2010). In other embodiments, the method provided elevates the omega-3 index (OM3I) and reduces oral glucose intolerance (OGTT). In some embodiments of increasing omega-3 index, the subject is diagnosed with moderate hypertriglyceridemia.

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[0113] Also provided are methods for reducing high sensitivity C-reactive protein (hs-CRP) in a subject. In some embodiments of reducing high sensitivity C-reactive protein (hs-CRP), the subject is diagnosed with moderate hypertriglyceridemia.

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5 [0114] Also provided are methods for inhibiting, preventing, or treating cardiovascular disease in a subject. Cardiovascular diseases include atherosclerosis, arteriosclerosis, coronary artery disease, heart valve disease, arrhythmia, heart failure, hypertension, orthostatic hypotension, shock, endocarditis, diseases of the aorta and its branches, disorders of the peripheral vascular system, and congenital heart disease.

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[0115] Also provided are methods for inhibiting, preventing, or treating metabolic syndrome in a subject. Metabolic syndrome is a combination of medical disorders that increase the risk of developing cardiovascular disease and diabetes. It affects one in five people, and prevalence increases with age. Some studies estimate the prevalence in the USA to be up to 25% of the population. Metabolic syndrome is also known as metabolic syndrome X, syndrome X, insulin resistance syndrome, Reaven's syndrome, and CHAOS (Australia).

Also provided are methods for inhibiting, preventing, or treating a cognitive [0116] 20 disorder, in a subject. The term "cognitive disease or disorder" as used herein should be understood to encompass any cognitive disease or disorder. Non-limiting examples of such a cognitive disease or disorder are Attention Deficit Disorder (ADD), Attention Deficit Hyperactivity Disorder (ADHD), dyslexia, age-associated memory impairment and learning disorders, amnesia, mild cognitive impairment, cognitively impaired non-demented, pre-25 Alzheimer's disease, autism, dystonias and Tourette syndrome, dementia, age related cognitive decline, cognitive deterioration, moderate mental impairment, mental deterioration as a result of ageing, conditions that influence the intensity of brain waves and/or brain glucose utilization, stress, anxiety, concentration and attention impairment, mood deterioration, general cognitive and mental well being, neurodegenerative disorders, 30 hormonal disorders or any combinations thereof. In a specific embodiment, the cognitive disorder is memory impairment.

[0117] The term "improving a condition in a subject suffering from a cognitive disease or a cognitive disorder" as used herein should be understood to encompass: ameliorating undesired symptoms associated with a disease, disorder, or pathological condition; preventing manifestation of symptoms before they occur; slowing down progression of a disease or disorder; slowing down deterioration of a disease or disorder; slowing down irreversible damage caused in a progressive (or chronic) stage of a disease or disorder; delaying onset of a

5 (progressive) disease or disorder; reducing severity of a disease or disorder; curing a disease or disorder; preventing a disease or disorder from occurring altogether (for example in an individual generally prone to the disease) or a combination of any of the above. For example, in a subject suffering from memory impairment, for example as a result of Alzheimer's Disease, symptoms including deterioration of spatial short-term memory, memory recall and/or memory recognition are improved by use of a lipid concentrated therapeutic phospholipid composition.

[0118] Also provided are methods for inhibiting, preventing, or treating neurodegenerative disorder in a subject. Neurodegenerative disorder is defined as a chronic progressive neuropathy characterized by selective and generally symmetrical loss of neurons in motor, sensory, or cognitive systems. Non limiting examples of neurodegenerative disorders include but are not limited to Alzheimer's disease, Pick's disease, Lewy body dementia Basal ganglia—Huntington's disease, Parkinson's disease, dentatorubropallidoluysian atrophy, Freidreich's ataxia, multiple system atrophy, types 1, 2, 3, 6, 7 spinocerebellar ataxia Motor—amyotrophic lateral sclerosis, familial spastic paraparesis, spinal muscular atrophy, spinal and bulbar muscular atrophy, Lou Gehrig's disease, pre-dementia syndrome, Lewy body dementia, age-related cognitive decline, cognitive deterioration, moderate mental impairment, mental deterioration as a result of ageing, dentatorubropallidoluysian atrophy, Freidreich's ataxia, multiple system atrophy, types 1, 2, 3, 6, 7 spinocerebellar ataxia, amyotrophic lateral sclerosis, and familial spastic paraparesis.

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[0119] Also provided are methods for reducing the decline of global cognitive function in a subject. In some embodiments, the reduction in decline of global cognitive function can be measured by the Neuropsychological Test Battery (NTB). In some embodiments, the subject is diagnosed with early stage Alzheimer's disease.

[0120] Also provided are methods for reducing worsening of neuropsychiatric symptoms in a subject. In some embodiments, the reduction is measured by the Neuropsychiatric Inventory questionnaire (NPI). In some embodiments, the subject is diagnosed with early stage Alzheimer's disease.

[0121] Also provided are methods for maintaining self-care and activities of daily living function in a subject suffering from Alzheimer's disease. In some embodiments, the subject is diagnosed with early stage Alzheimer's disease. In some embodiments, the maintaining self-care and activities of daily living function is measured by the Disability Assessment in Dementia caregiver-based interview (DAD).

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[0122] Additional health disorders or conditions which may be treated or improved by the concentrated therapeutic phospholipid compositions include, but are not limited to, high blood cholesterol levels, high triglycerides levels, high blood fibrinogen levels, low HDL/LDL ratio, menopausal or post-menopausal conditions, hormone related disorders, vision disorders, immune disorders, liver diseases, chronic hepatitis, steatosis, lipid peroxidation, dysrhythmia of cell regeneration, destabilization of cell membranes, high blood pressure, cancer, hypertension, aging, kidney disease, skin diseases, edema, gastrointestinal diseases, peripheral vascular system diseases, allergies, airways diseases, and psychiatric diseases.

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#### Combination Therapies

[0123] In some embodiments, the subject is administered an effective amount of a concentrated therapeutic phospholipid composition. In other embodiments, the treatment comprises a combination of a concentrated therapeutic phospholipid composition and treatment agents such as anti-dyslipidemic agents. Anti-dyslipidemic agents include but are not limited to atorvastatin, fluvastatin, lovastatin, pitavastatin, pravastatin, rosuvastatin and simvastatin.

[0124] In other embodiments, the treatment comprises a combination of a concentrated therapeutic phospholipid composition and a cholinesterase inhibitor. Cholinesterase inhibitors include but are not limited to metrifonate (irreversible), carbamates, physostigmine, neostigmine, pyridostigmine, ambenonium, demarcarium, rivastigmine, phenanthrene derivatives, galantamine, piperidines, donepezil, tacrine, edrophonium, huperzine A, ladostigill and ungeremine.

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5 [0125] In some embodiments, the subject is administered a combination of a concentrated therapeutic phospholipid composition and at least one of vitamins, minerals, cox-inhibitors, sterols, fibrates, antihypertensives, insulin, cholesterol digestion inhibitors, for example, ezetimibe, fatty acids, omega-3 fatty acids, antioxidants, and the methylphenydate class of compounds, such as for example ritalin. In other embodiments, a combination of a 10 concentrated therapeutic phospholipid composition and elements depleted during traditional chronic treatments, such as for example during chronic treatment with statins. For example, in some embodiments, a concentrated therapeutic phospholipid composition is described which contains at least one of cox-2, folic acid, vitamin B6, vitamin B12, magnesium or zinc. In other embodiments, combination therapies comprising a concentrated therapeutic 15 phospholipid composition and potassium are described. Potassium is usually depleted during treatment with diuretics. Combination therapies reduce risk of side effects, increase benefits, increase solubility, and/or increase bioavailability.

## Modes of Administration

20 **[0126]** Administration of the concentrated therapeutic phospholipid compositions can be accomplished via any mode of administration for therapeutic agents. These modes include systemic or local administration such as oral, parenteral, transdermal, subcutaneous, or topical administration modes.

#### 25 Pharmaceutical Formulations

[0127] Depending on the intended mode of administration, the compositions can be in solid, semi-solid or liquid dosage form, such as, for example, injectables, tablets, pills, time-release capsules, elixirs, tinctures, emulsions, syrups, liquids, suspensions, or the like, sometimes in unit dosages and consistent with conventional pharmaceutical practices.

30 Likewise, they can also be administered in intravenous (both bolus and infusion), intraperitoneal, subcutaneous or intramuscular form, all using forms well known to those skilled in the pharmaceutical arts.

[0128] Illustrative pharmaceutical compositions are tablets and gelatin capsules

comprising a concentrated therapeutic phospholipid composition neat, or if required, contains a pharmaceutically acceptable carrier, such as a) a diluent, *e.g.*, purified water, triglyceride oils, such as hydrogenated or partially hydrogenated vegetable oil, or mixtures thereof, corn oil, olive oil, sunflower oil, safflower oil, lactose, dextrose, sucrose, mannitol, sorbitol,

5 cellulose, sodium, saccharin, glucose and/or glycine; b) a lubricant, e.g., silica, talcum, stearic acid, its magnesium or calcium salt, sodium oleate, sodium stearate, magnesium stearate, sodium benzoate, sodium acetate, sodium chloride and/or polyethylene glycol; for tablets also; c) a binder, e.g., magnesium aluminum silicate, starch paste, gelatin, tragacanth, methylcellulose, sodium carboxymethylcellulose, magnesium carbonate, natural sugars such 10 as glucose or beta-lactose, corn sweeteners, natural and synthetic gums such as acacia, tragacanth or sodium alginate, waxes and/or polyvinylpyrrolidone, if desired; d) a disintegrant, e.g., starches, agar, methyl cellulose, bentonite, xanthan gum, algiic acid or its sodium salt, or effervescent mixtures; e) absorbent, colorant, flavorant and sweetener; f) an emulsifier or dispersing agent, such as Tween 80, Labrasol, HPMC, DOSS, caprovl 909, 15 labrafac, labrafil, peceol, transcutol, capmul MCM, capmul PG-12, captex 355, gelucire, vitamin E TGPS or other acceptable emulsifier; and/or g) an agent that enhances absorption of the compound such as cyclodextrin, hydroxypropyl—cyclodextrin, PEG400, PEG200.

[0129] Liquid, particularly injectable, compositions can, for example, be prepared by dissolution, dispersion, *etc*. For example, the concentrated therapeutic phospholipid composition is dissolved in or mixed with a pharmaceutically acceptable solvent such as, for example, water, saline, aqueous dextrose, glycerol, ethanol, and the like, to thereby form an injectable isotonic solution or suspension. Proteins such as albumin, chylomicron particles, or serum proteins can be used to solubilize the concentrated therapeutic phospholipid composition.

[0130] Other illustrative topical preparations include creams, ointments, lotions, aerosol sprays and gels, wherein the concentration of the concentrated therapeutic phospholipid composition ranges from about 0.1 % to about 15 %, w/w or w/v.

30 Dosing

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[0131] The dosage regimen utilizing the concentrated therapeutic phospholipid compositions is selected in accordance with a variety of factors including type, species, age, weight, sex and medical condition of the subject; the severity of the condition to be treated; the route of administration; the renal or hepatic function of the subject; and the particular concentrated therapeutic phospholipid composition employed. A physician or veterinarian of ordinary skill in the art can readily determine and prescribe the effective amount of the drug required to prevent, counter or arrest the progress of the condition.

Effective dosage amounts of the present invention, when used for the indicated effects, range from about 20 mg to about 10000 mg of the concentrated therapeutic phospholipid composition per day. Dosages for *in vivo* or *in vitro* use can contain about 20, 50, 75, 100, 150, 250, 500, 750, 1000, 1250, 2500, 3500, 5000, 7500 or 10000 mg of the concentrated therapeutic phospholipid composition. Effective blood plasma levels after administration of the concentrated therapeutic phospholipid composition to a subject can range from about 0.002 mg to about 100 mg per kg of body weight per day. Appropriate dosages of the concentrated therapeutic phospholipid composition can be determined as set forth in L.S.

concentrated therapeutic phospholipid composition can be determined as set forth in L.S. Goodman, et al., The Pharmacological Basis of Therapeutics, 201-26 (5th ed.1975).

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[0133] The concentrated therapeutic phospholipid compositions can be administered in a single daily dose, or the total daily dosage can be administered in divided doses of two, three or four times daily. To be administered in the form of a transdermal delivery system, the dosage administration can be continuous rather than intermittent throughout the dosage regimen. In some embodiments, of combination therapy, the concentrated therapeutic phospholipid composition and the therapeutic agent can be administered simultaneously. In other embodiments, the concentrated therapeutic phospholipid composition and the therapeutic agent can be administered sequentially. In still other embodiments of combination therapy, the concentrated therapeutic phospholipid composition can be administered daily and the therapeutic agent can be administered less than daily. In still other embodiments of combination therapy, the concentrated therapeutic phospholipid composition can be administered daily and the therapeutic agent can be administered more than once daily.

#### **EXAMPLES**

The disclosure is further illustrated by the following examples, which are not to be construed as limiting this disclosure in scope or spirit to the specific procedures herein described. It is to be understood that the examples are provided to illustrate certain embodiments and that no limitation to the scope of the disclosure is intended thereby. It is to be further understood that resort may be had to various other embodiments, modifications, and equivalents thereof which may suggest themselves to those skilled in the art without departing from the spirit of the present disclosure and/or scope of the appended claims.

### Concentrated Therapeutic Phospholipid Compositions

[0135] The following non-limiting examples of therapeutic compositions serve to illustrate further embodiments of the concentrated therapeutic phospholipid composition. It is to be understood that any embodiments listed in the Examples section are embodiments of the concentrated therapeutic phospholipid composition and, as such, are suitable for use in the methods and compositions described above.

[0136] The following methods can be used to make the concentrated therapeutic phospholipid compositions (Figure 1A and 1B)

Step 1:

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[0137] Frozen krill is mechanically crushed and incubated with a solvent in a ratio of 9:1 acetone water for 60-90 minutes at 8°C to extract different proportions of the lipids (PL, TG and FFA) from the krill biomass. Lipids are subsequently separated from proteins and krill material by filtration under pressure (50-60 kpa). The solid phase is discarded. The soluble extract is evaporated by a continuous distillation column under vacuum to remove the solvent (acetone). The major part of the aqueous (water) fraction is separated from the lipid fraction by decantation and the remaining water removed by evaporation under vacuum and gentle heating. Those fractions are dosed, analyzed, and blended to constitute an intermediary krill oil product which is re-analyzed to achieve desired specifications  $\pm$  5%: EPA (15g/100g), DHA (9g/100g), total phospholipids (42g/100g) and astaxanthin's forms (125mg/100g).

Step 2:

[0138] 100.5g of received krill oil from step 1 was charged to a 300ml extraction vessel (ID = 0.68"). The extractor was sealed, pre-heated CO2 at  $55^{\circ}$ C was introduced from the bottom, and the pressure in the extractor was maintained at 5,000 psi using a diaphragm CO2 pump. The flow of CO2 was continued in the upflow direction through the extractor and was expanded to atmospheric pressure through a pressure-reduction-valve (PRV) so that the dissolved material in the CO2 precipitated and collected in the flask. The flow rate and volume of CO2 exiting the flask was measured with a flowmeter and dry test meter (DTM). A total of 7200g of CO2 was passed through the extractor (solvent to feed ratio, S/F = 72) and 34.1% of the charge was removed by the CO2. The flow of about 25 Standard Liters/min of CO2 was maintained during the course of the test and the total time of extraction was

about 160min. The extractor was isolated and the CO2 was vented to atmosphere. The extractor was opened and the un-extracted material (raffinate-product) was removed from the vessel.

Step 3:

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[0139] SC CO2 Extraction to produce 90+% OM3:PLs. 9.44g of oil was mixed with inert packing and charged to the extractor. The procedure carried out similar to that desribed in step 2 except more aggressive extraction conditions were used with the pressure and temperature in the extractor maintained at 10,000 psi and 70°C. A total S/F ratio of 200 was used f; therefore, about 1900g of CO2 was flowed through the extractor. A flow rate of CO2 of about 10 Standard Liters/min was maintained; therefore, the total run time for this test was 105min. A total of 56.3% of the charge was extracted from this oil during the course of the run. The extractor was isolated, the CO2 vented to atmosphere, the vessel opened, and the resulting raffinate-product scraped off the inert packing. This un-extracted material analyzed to be 91% OM3:PLs.

	47 % OM3:PL			
*Total lipids as FA <sub>TG</sub>	g/100g oil	61.3	(100)	
*Omega-3	g/100g oil	14.1	(22,5)	
*EPA	g/100g oil	7.4	(11,6)	
*DHA	g/100g oil	3.8	(6,1)	
*DPA	g/100g oil	0.2	(0,3)	
*Omega-6	g/100g oil	10.8	(18,3)	
*Linoleic acid	g/100g oil	10.6	(18,0)	
*Omega-9	g/100g oil	6.6	(11,6)	
*Oleic acid	g/100g oil	6.1	(10,8)	
*Sat. FA <sub>TG</sub>	g/100g oil	21.4	(36,1)	
*Monounsat. FA <sub>TG</sub>	g/100g oil	13.9	(23,1)	
*Polyunsat. <sub>TG</sub>	g/100g oil	26.0	(40,7)	
*EPA as FA <sub>TG</sub>	g/100g oil	7.7		
*DHA as FA <sub>TG</sub>	g/100g oil	3.9		
Water	%	0.8		
Color	-	Red ora	nge	
Odor	-	Slightly r		
Total carotenoids	mg/100g oil	36.0		
Astaxanthine	mg/100g oil	65.3		
Astaxanthine	% diester	83.1		
	% monoester	16.9	1	
	% free	0.0		
Peroxide Index	mEq peroxide/kg	1.0		
p-Anisidine Index	-	2.0		
Iodine Index	gI <sub>2</sub> /100g oil	101.1		
Saponification Index	mg KOH/g oil	214.1		
Indice acide	mg KOH/g oil	17.2		
Total fat	%			
Free fatty acid	% as oleic acid	5.2		
Triglycerides	%	36.5		
Viscosity	cP	1323.	.0	
ash	%	5.0		
Vitamin A	UI/g Oil	40.4		
Vitamin E	UI/g Oil	0.1		
Total phospholipids	g/100g oil	47.2		
Phospholipid profile	TLC	-		
	% LPC	3.7		
	% PC	53.6		
	% PS	24.7		
	% PE	16.4		
	% PA	1.7		
Molecular mass PL	g/mol	773.8	3	

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53%	OM3:PL	
Total lipids as fatty acids (FA) TG	g / 100 g oil	69.80
Total Omega-3	g / 100 g oil	31.30
C 20:5 (n=3) EPA	g / 100 g oil	13.90
C 22:6 (n=3) DHA	g / 100 g oil	10.10
C 22:5 (n=3) DPA	g / 100 g oil	0.40
Total Omega-6	g / 100 g oil	1,60
linoleic acid – LA	g / 100 g oil	1.30
Total Omega-9	g / 100 g oil	6.10
oleic acid – OA	g / 100 g oil	5.70
Saturated FA	g / 100 g oil	21.10
Monounsaturated FA	g / 100 g oil	14.50
Polyunsaturated FA	g / 100 g oil	34.20
EPA as FA	g / 100 g oil	14.40
DHA as FA	g / 100 g oil	10.50
PHOSPHOLIPID PROFILE		
total	g / 100 g oil	52.30
lysophosphatidyl choline - LPC	%	10.80
sphingomyeline - SM	%	0.10
phsophatidyl choline - PC	%	79.70
phsophatidyl serine - PS	%	
phsophatidyl inositol - PI	%	
phosphatidyl ethanolamine - PE	%	9.40
PA	%	0.00
CAROTENOIDS		
total	mg / 100 g oil	92.60
total astaxanthin - AST	mg / 100 g oil	161.60
AST diester		62.00
AST monoester		35.00
AST free		3.00

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66% OM3:PL		
Total lipids as fatty acids (FA) TG	g / 100 g oil	74.2
Total Omega-3	g / 100 g oil	39.8
C 20:5 (n=3) EPA	g / 100 g oil	21.7
C 22:6 (n=3) DHA	g / 100 g oil	14.1
C 22:5 (n=3) DPA	g / 100 g oil	0.5
Total Omega-6	g / 100 g oil	1.7
linoleic acid - LA	g / 100 g oil	1.3
Total Omega-9	g / 100 g oil	5.8
oleic acid - OA	g / 100 g oil	5.1
Saturated FA	g / 100 g oil	18.0
Monounsaturated FA	g / 100 g oil	13.2
Polyunsaturated FA	g / 100 g oil	43.1
EPA as FA	g / 100 g oil	22.6
DHA as FA	g / 100 g oil	14.6
PHOSPHOLIPID PROFILE		
total	g / 100 g oil	66.2
lysophosphatidyl choline - LPC	%	10.7
phsophatidyl choline - PC	%	75.3
phosphatidyl ethanolamine - PE	%	11.8
other	%	2.2
CAROTENOIDS		
total	mg / 100 g oil	273.4
total astaxanthin - AST	mg / 100 g oil	466.8
AST diester	%	57.4
AST monoester	%	40.7
AST free	%	1.9

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80% OM3:PI	<u> </u>	
Total lipids as fatty acids (FA) TG	g / 100 g oil	68.35
Total Omega-3	g / 100 g oil	37.90
C 20:5 (n=3) EPA	g / 100 g oil	20.40
C 22:6 (n=3) DHA	g / 100 g oil	12.95
C 22:5 (n=3) DPA	g / 100 g oil	0.48
Total Omega-6	g / 100 g oil	1.45
linoleic acid - LA	g / 100 g oil	1.26
Total Omega-9	g / 100 g oil	4.93
oleic acid - OA	g / 100 g oil	4.35
Saturated FA	g / 100 g oil	16.15
Monounsaturated FA	g / 100 g oil	11.21
Polyunsaturated FA	g / 100 g oil	40.99
EPA as FA	g / 100 g oil	21.30
DHA as FA	g / 100 g oil	13.50
PHOSPHOLIPID PROFILE		
total	g / 100 g oil	80.00
lysophosphatidyl choline - LPC	%	9.20
sphingomyeline - SM	%	0.20
phsophatidyl choline - PC	%	80.60
phsophatidyl serine - PS	%	1.10
phsophatidyl inositol - PI	%	0.10
phosphatidyl ethanolamine - PE	%	7.50
PA	%	1.30
CAROTENOIDS		
total	mg / 100 g oil	180.4
total astaxanthin - AST	mg / 100 g oil	325.5
AST diester	%	68.45
AST monoester	%	29.27
AST free	%	2.28

9	0% OM3:PL	
*Total lipids as FA <sub>TG</sub>	g/100g oil	63.9
*Omega-3	g/100g oil	35.1
*EPA	g/100g oil	18.9
*DHA	g/100g oil	12.2
*DPA	g/100g oil	0.5
*Omega-6	g/100g oil	1.3
*Linoleic acid	g/100g oil	1.2
*Omega-9	g/100g oil	4.6
*Oleic acid	g/100g oil	3.9
*Sat. FA <sub>TG</sub>	g/100g oil	15.8
*Monounsat. FA <sub>TG</sub>	g/100g oil	10.2
*Polyunsat. <sub>TG</sub>	g/100g oil	37.9
*EPA as FA <sub>TG</sub>	g/100g oil	19.7
*DHA as FA <sub>TG</sub>	g/100g oil	12.7
Acetone	Ppm	1.6
Humidity and volatiles	%	1.7
Water	%	1.9
Color	-	Red chili
Odor	-	shellfish
Total carotenoids	mg/100g oil	168.9
Astaxanthine	mg/100g oil	309.3
Astaxanthine	% Diester	73.1
	% Monoester	25.3
	% Libre	1.6
Index p-Anisidine	-	3.1
Index acid	mg KOH/g oil	33.6
Index iodine	gI <sub>2</sub> /100g oil	
Index saponification	mg KOH/g oil	
Index Peroxide	mEq peroxyde/kg	0.1
Vitamin A	UI/g Oil	15.2
Vitamin E	UI/g Oil	0.3
Fatty acid total	%	97.6
Viscosity	сР	
Total phospholipids	g/100g oil	90.6
Phospholipid profile	TLC	-
	% LPC	13.5
	% SM	0.4
	% PC	76.3
	% Autres PL	1.2
	% PE	7.9
	% PA	0.8
Triglycerides	%	0.0

70% OM3:PL	derived from Squid	
*Total lipids as FA <sub>TG</sub>	g/100g oil	54,5
*Omega-3	g/100g oil	29,1
*EPA	g/100g oil	8,9
*DHA	g/100g oil_	18,3
*DPA	g/100g oil	0,2
*Omega-6	g/100g oil_	0,7
*Linoleic acid	g/100g oil_	0,3
*Omega-9	g/100g oil	4,2
*Oleic acid	g/100g oil	2,0
*Sat. FA <sub>TG</sub>	g/100g oil	16,9
*Monounsat, FA <sub>TG</sub>	g/100g oil	6,8
*Polyunsat. <sub>TG</sub>	g/100g oil	30,9
*EPA as FA <sub>TG</sub>	g/100g oil	9.3
	g/100g oil	19.1
*DHA as FA <sub>TG</sub> Humidity (calmar)	9/100g on 9/0	17.1
Indice acide	mg KOH/g oil	55.7
Vitamin A	UI/g Oil	33.1
Vitamin A  Vitamin E	UI/g Oil	
Fatty acid total	%	2.7
Total carotenoids	mg/100g oil	8.3
Astaxanthine	mg/100g oil	13.2
Astaxanthine	% diester	42.5
Astavantime	% monoester	35.6
	% libre	21.9
Total phospholipids	g/100g oil	70,8*
Phospholipid profile	TLC	- 70,8
i nosphonpia prome	% LPC	12,4
	% SM	7.8
	% PC	55.8
	% other	2.0
	% PE	22.0
	% PA	6.4
Triglycerides	%	25.0
Free fatty acid	% as oleic acid	3.2
Index p-Anisidine	, us office dela	4.3
Index Peroxide	mEq peroxyde/kg	0.6
	mining peroxydering	
**PM Phospholipids (g/mol)		847.14
Profile of Fatty Acids of the PL		
*Total lipids as FA	g/100g PL	53,9
*Oméga-3	g/100g PL	28,6
*EPA	g/100g PL	9,0
*DHA	g/100g PL	18,3
*DPA	g/100g PL	0,2
*Omega-6	g/100g PL	0,5
*Linoleic acid	g/100g PL	0,2
*Omega-9	g/100g PL	3,6

70% OM3:PL derived from Squid				
*Oleic acid	g/100g PL	1,5		
*Sat. FA	g/100g PL	18,3		
*Monounsat. FA	g/100g PL	5,4		
*Polyunsat.FA	g/100g PL	30,2		

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#### **Biological Examples**

## Example 1

#### Managing Dyslipidemia in Three Murine Phenotypes

[0140] The aim of this study was to examine the effects of Composition 3 in three age/sex-matched murine phenotypes representative of (1) normal healthy non-obese
normoglycemic control (C57BL6) versus (2) hyperdislipidemic LDL-receptor gene knockout
(LDLr -/-) or (3) human apoA-I transgenic mice (Jackson Labs) at 12-w of age: 27.5±0.7 vs
25.6±0.7 vs 29.2±0.8 gr., respectively; n=7-10/gr. kept according to local and national ethic
regulations, fed a normal vs a Western-diet regime and water *ad libitum*. Data are presented
as mean±sem and statistics assessed by t- test (unpaired, two-tail) (v5-GraphPad Prism).

The profile of plasma lipids concentrations (mg/dL) in the above-three untreated [0141] adult male murine models were as reported in the literature: Total cholesterol (TC): 71.1±3.3 vs 215.3±10.4 vs 50.3±1.3; triglycerides (TGs): 59.5±4.5 vs 65.1±3.8 vs 53.0±12.9; lowdensity lipoprotein (LDL): 13.3±1.2 vs 101.6±6.7 vs 12.2±1.6; high-density lipoprotein (HDL): 53.4±3.2 vs 88.8±3.6 vs 24.8±2.6. Six (6) weeks of QD treatment with Composition 3 (104 vs 208 vs 417 mg/kg (Human equivalent dosing of 500, 1,000 and 2,000 mg/day) in C57BL6 led to significant dose-dependent decrease in plasma TGs (up to 60%), reduced LDL (up to 28%), elevated HDL (by 17%) but did not affect TC (see Figures 2-12 and Table 1). In severely dyslipidemic LDLr-KO mice, Composition 3 led to significant dose-dependent decrease in plasma TGs, elevated further HDL, caused a slight elevation in TC (only at middose) and did not affect LDL (see Figures 13-17 and Table 1). In hApoA-I transgenic mice, Composition 3 led to significant decrease in plasma TGs, elevated HDL and did not affect TC (see Figure 18 and Table 1). The liver concentrations of TC were the same in all three phenotypes but TGs were reduced (19%; p<0.05) in LDLr-KO and elevated (153%; p<0.01) in hApoA-I, compared to control C57BL6. Treatment with Composition 3 elevated liver TC

and TGs by up to 12% and 27%, respectively, in C57BL6, by up to 10% and 36% in LDLr-KO and by up to 10% for TC but mixed effects for TGs (-13 to + 12%) in hApoA-I mice, respectively (see Table 2).

Table 1: Effects mediated by 6-weeks treatment with Composition 3 on plasma lipids.

Murine				High-dose	
phenotype	Lipids	104 mg/kg	208 mg/kg	417 mg/kg	
		(HED: 500mg/day)	(HED: 1000mg/day)	(HED: 2000mg/day)	
C57BL6	TC	75.9±2.0	80.6±3.3	78.1±2.7	
		up 6.8% (NS)	up 13.4% (NS)	up 9.8% (NS)	
	TGs	32.2±1.6	26.0±2.8	$23.8 \pm 1.2$	
		down 46% (p<0.001)	down 56% (p<0.001)	down 60% (p<0.001)	
	LDL	11.1±1.4	11.2±1.0	9.6±0.7	
		down 15% (NS)	down 16% (p≤0.05)	down 28% (p<0.05)	
	HDL	59.4±1.9	63.4±3.5	62.3±2.4	
		up 11% (NS)	up 19% (NS)	up 17% (p<0.05)	
LDLr-KO	TC	219.3±7.4	244.4±7.9	238.5 ±6.9	
		up 2% (NS)	up 14% (p<0.01)	up 11% (NS)	
	TGs	45.7±2.8	41.7±4.9	36.7±1.6	
		down 30% (p<0.001)	down 36% (p<0.01)	down 44% (p<0.001)	
	LDL	99.2±5.7	116.2 ±4.4	97.4±6.5	
		down 2% (NS)	up 14% (NS)	down 4% (NS)	
	HDL	90.9±2.5	88.4 ±4.6	111.2±2.5	
		up 2% (NS)	= (NS) up 25% (p		
hApoA-I	TC	51.1±1.5	57.2±2.6	50.1±1.4	
transgenic		up 2% (NS)	up 14% (p<0.05)	= (NS)	
	TGs	19.3±2.8	43.2±11.7	43.0±8.2	
		down 64% (p<0.05)	down 18% (NS)	down 19% (NS)	
	LDL	10.8±1.0	12.8±2.3	13.9±1.0	
		down 11% (NS)	= (NS)	up 14% (NS)	
	HDL	27.9±2.4	28.6±2.6	21.6±12.8	
		up 13% (NS)	up 15% (NS)	down 13% (NS)	

NS; not significant

## 10 Table 2: Plasma lipids at baseline between murine phenotype

Plasma concentrations (mg / dL)	C57BL6 control	LDLr-KO	Variation vs control	hApoA-I transgenic	Variation vs control
total cholesterol (TC)	71.1±3.3	215.3±10.4	up 3-fold	50.3±1.3	down 29.3%
		·	p<0.001		NS
triglycerides (TGs)	59.5±4.5	65.1±3.8	up 9.4%	53.0±12.9	down 10.9%
			NS		NS
low density lipoprotein	13.3±1.2	101.6±6.7	up 7.6-fold	12.2±1.6	down 8.3%
(LDL-C)			p<0.001		NS
high-density lipoprotein	53.4±3.2	88.8±3.6	up 166%	24.8±2.6	down 54%
(HDL-C)			p<0.001		p<0.001
Liver concentrations					
(µg / mg)					
total cholesterol (TC)	23.1±0.8	23.6±0.6	= (NS)	25.1±0.5	up 9%
			, ,		(NS)
triglycerides (TGs)	53.0±3.1	42.9±2.1	down 19%	81.3±7.4	up 153%
			p<0.05		p<0.001

NS; not significant

5 [0142] These data indicate that the Composition 3 is an effective modulator of lipid metabolism, mainly at reducing plasma TGs and LDL and elevating HDL. These data indicate that in some embodiments, concentrated therapeutic phospholipid compositions can be effective as a therapy against moderate to severe hypertriglyceridemia. In some embodiments, concentrated therapeutic phospholipid compositions in combination with other anti-dyslipidemic agents can be effective at lowering refractory hypertriglyceridemia.

## Example 2

Increase of the circulating plasma concentration of high-density Lipoprotein-Cholesterol (HDL-C) and Reduced Total Cholesterol (TC) / HDL ratio in 12-week old Male Zucker Diabetic Fatty Rats.

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[0143] The purpose of this study was to examine the effects of the concentrated therapeutic phospholipid compositions in the Zucker Diabetic Fatty rat rodent model for type 2 diabetes with obesity, hyperlipidaemia and insulin resistance (ZDF; Gmi-fa/fa) vs age-/sex-matched normal healthy non-obese normoglycemic lean control SD rat (from Charles River Labs; 12w, 359±17 vs 439±13 gr.; n=9-12/gr.). Lipid profile (total cholesterol (TC), tryglycerides (TGs), High-Density Lipoprotein-Cholesterol (HDL-C) and TC / HDL ratio were assessed before, 1 and 2 month after QD treatment with Composition 3 (52 versus 260 mg/kg (HED of 500 and 2,500 mg), and kept according to local and national ethic regulations (Formulab high fat 5008 (ZDF) vs normal 5001 (SD) diet regime and water ad libitum). Data are presented as Mean±SD (n=2-10) and statistical differences calculated by unpaired two-tailed t test (v5-GraphPad Prism). At 12-w of age, the circulating plasma concentrations of TC, TGs, HDL and TC/HDL ratio were: 4.6±0.9, 11.6±5.9, 2.3±1.1 mmol / L and 2.16±0.62, respectively. Lipids' profile in SD rats were significantly lower at 1.9±0.4, 1.2±0.4, 1.3±0.2 mmol/L and 1.45±0.11, respectively. Daily low and high dose treatment for 60 days did not affect TC and TGs concentrations but increased by 1.7- to 1.8-fold (p<0.01) "good" HDL-cholesterol and decreased the TC / HDL ratio by 26-32% (p<0.01-0.05), respectively.

5 Example 3

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# **Improved Glucose Intolerance in Zucker Diabetic Fatty Rats after Administration of Composition 3**

[0144] The purpose of this study was to investigate the effects of Composition 3 in an overtly dislipidemic, obese, type 2 diabetic rat model. Zucker Diabetic Fatty (ZDF; Gmi-fa/fa) rats were used *versus* age-/sex-matched normal healthy non-obese normoglycemic lean control SD rat (from Charles River Labs; 12-w, 359±17 vs. 439±13 gr.; n=9-12/gr.). Glucose intolerance was assessed conducting an oral glucose tolerance test (OGTT; overnight fasting then single gavage of glucose 2 g/kg rat b.w.) over 180 minutes using glucometer strips (Accu-Chek Aviva, Roche Diagnostics), before and 90 days after treatment with Composition 3 given by QD gavage at 52 versus 260 mg/kg (HED of 500 and 2,500 mg), and kept according to local and national ethic regulations (Formulab high fat 5008 (ZDF) vs normal 5001 (SD) diet regime and water *ad libitum*). Data are presented as Mean±SD and statistical differences calculated by unpaired two-tailed t test (v5-GraphPad Prism).

At 12-w of age  $(T_0)$  fasting circulating plasma concentrations of glucose were  $7.8\pm2.1 \text{ vs } 5.0\pm0.6 \text{ mmol/L } (p<0.001) \text{ in untreated ZDF vs SD rats. Non-fasting ZDF and$ SD rats glucose levels were 22.0±4.2 vs 8.6±0.6 mmol / L, respectively. One month later, baseline values increased by 1.9-fold (p<0.0001) in fasted ZDF while remaining unchanged in fasted SD. Aging did not affect glucose levels in non-fasted rats. Glucose challenge led to a maximum 2.5-fold (p<0.0001) and 1.6-fold (p<0.0001) increase in plasma glucose concentrations in untreated fasted ZDF and SD, at 30 and 60 minutes, respectively, returning mostly to initial values after 180 mins. At 16 weeks of age, thirty days (T<sub>30</sub>) of treatment did not affect either the profile (AUC) or maximum elevation in glucose in SD rats but treatment of ZDF shifted to the right the maximum elevation in plasma glucose (from 30 to 60 minutes), reduced by 61-72% (p<0.02) the peak elevation at 30 minutes and reduced by 50-60% (p<0.0001) the AUC at either doses of Composition 3, thus back to the AUC observed in untreated glucose challenged SD rats. At 20 weeks of age, 60 days of treatment, either dosing did not further attenuate glucose intolerance. None of the treatment profile affected the plasma and urinary concentrations of glucose (hyperglycemia and glucosuria) in nonfasted ZDF or SD. These data indicate that a short term and low dose chronic administration of Composition 3 significantly improves glycemic control in a model of severe hyperglycemia.

5 Example 4

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A randomized, placebo-controlled, double-blind, dose-ranging and multi-centered trial to evaluate the safety and efficacy of concentrated therapeutic phospholipid compositions in the treatment of moderate hypertriglyceridemia.

10 Subjects with moderate hypertriglyceridemia treated by physician according to the Canadian Lipid Treatment Guidelines who are treated over 12 weeks with concentrated phospholipid given at doses of 1.0, 2.0 or 4.0g. The primary measure of efficacy will be the percent change in fasting blood circulating serum triglycerides (TGs) between baseline (Week 1) and 12 weeks of treatment. Secondary Outcomes: between baseline and after six weeks and 12 15 weeks of treatment: 1) absolute change in fasting plasma TGs; 2) percentage (%) of subjects achieving target TG fasting plasma levels; 3) absolute change in fasting plasma LDL-C. VLDL-C, HDL-C, HDL2-C, HDL3-C, Total Cholesterol, hs-CRP and non-HDL; 4) percentage (%) change in fasting plasma concentrations of LDL-C, VLDL-C, HDL-C, HDL2-C, HDL3-C, TC, hs-CRP and non-HDL; 5) calculated Ratios: a) total cholesterol: 20 HDL-C; b) LDL-C: HDL-C; c) TGs: HDL-C; 6) LDL-C-related parameters: a) particle number; b) particle size; c) oxidation; 7) absolute and percent (%) change in fasting plasma concentrations of biomarkers; a) glycated Hemoglobin (HbA1c), b) apolipoprotein A-I (ApoA-I), c) apolipoprotein B-100 (ApoB-100), d) apolipoprotein E (ApoE), e) lipoprotein(a) (Lp(a)), f) adiponectin, g) glucose, h) insulin; 8) calculated ApoB: ApoA-I ratio; 9) fasting 25 plasma lipoprotein-associated phospholipase A2 activity (Lp-PLA2); 10) HOMA-IR (homeostasis model assessment of insulin resistance: [glucose (mmol / L) x IRI (microIU / L) / 22.5]; 11) plasma concentrations of total EPA and DHA (PK/PD – 25 subjects/group); 12) OM3I (Omega-3 index); 13) Subjects Genetic Polymorphism: a) Lecithin: Cholesterol Acyltransferase (LCAT), b) cholesteryl ester transfer protein (CETP), c) scavenger receptor

type B-1 (SR-B1), d) ATP Binding Cassette transporter 1 (ABCA1).

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5 Example 5

Preclinical non-GLP assessment of efficacy of concentrated therapeutic phospholipid compositions alone or in combination with a statin on modulating blood lipids and the development of atherosclerotic lesions in ApoE null mice fed a Western-type diet.

Male adult mice (n=135 (15 mice / group ) 5-6 weeks of age) weighing about 18-20g each homozygous for the Apoetm1Unc mutation are administered HOW either Vehicle (Water or 0.2%-0.5% Carboxymethylcellulose); composition 3 (1,000 mg/daily HED); composition 3 (2,000 mg/daily HED); or Lipitor (20 mg/daily HED); or composition 3 (1,000 mg/daily HED) + Lipitor (20 mg/daily HED). At 0, 3 months, or 6 months the following assessment of values is made relative to: blood Lipids: TC, TGs, LDL, HDL, non-HDL, VLDL (0, 3 and 6 months) (2)Aortic Atherosclerosis (0, 3 and 6 months): a. Thoracic and abdominal aorta will be isolated, trimmed of fat, laid out and pinned on black matrix for photography, and stained with Sudan IV or Oil Red-O. b.Vessel will be imaged for surface involvement using a
computerized image analysis system (Image ProPlus or NIH Package Software). The data will be computed by group and statistically analyzed. c.Lipid extraction: Following staining

and morphometric analysis, aortas will be extracted (Bligh/Dyer). (3) Red blood cells Omega-3 Index (0, 3 and 6 months); (4) Circulating plasma concentration of CRP (0, 3 and 6 months).

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#### Example 6

#### Comparison of Composition 3 with Lovaza® on the Omega 3 Index

Male sdult (14 weeks) Sprague-Dawley (SD) rats with an avereage body weight of >375-425 were fed normal rat chow(diet 5075- normal, standard rat chow). Number of test subjects / group: n=56; n=8 rats / gr. Dosing was QD (single daily dosing/morning) for 12 weeks with either (i) Vehicle (ii) Composition 3 52 mpk 500 mg / day HED; (iii) Composition 3 104 mpk 1000 mg / day HED; (iv) Composition 3 416 mpk 4000 mg / day HED; (v) Lovaza® 416 mpk = 4000 mg / day HED. Results are shown in Figure 35.

5 Example 7

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Montherapy study of Concentrated Phospholipids in early stage Alzheimer's Disease Subjects will be randomly assigned to receive either concentrated therapeutic phospholipid composition 1 g, fish oil (135 mg EPA: 108 mg DHA) 1 g, or placebo (soy oil) 1 g once daily. The primary outcome measure will be the change in NTB between baseline and 24 weeks of treatment. The Neuropsychological Test Battery (NTB) will be used to monitor and evaluate important cognitive changes. The following 9 components of the NTBare used to determine the outcome for the subject: (1) Wechsler Memory Scale, visual immediate (score range, 0-18), (2) Wechsler Memory Scale verbal immediate (score range, 0-24), (3) Rey Auditory Verbal Learning, Test (RAVLT) immediate (score range, 0-105), (4) Wechsler, Memory Digit Span (score range, 0-24), (5) Controlled Word Association, Test (COWAT), (6) Category Fluency Test (CFT), (7) Wechsler, Memory Scale visual delayed (score range, 0-6), (8) Wechsler, Memory Scale verbal delayed (score range, 0-8), and (9) RAVLT, delayed (score range, 0-30) (Harrison et al. 2007). The RAVLT delayed measure is composed of delayed recall and recognition performance components that are summed to yield a score ranging from 0 to 30, yielding 9 measures of subject performance. Secondary outcome measures will include the change in the NPI and DAD at 24 weeks of treatment. The NPI evaluates 12 neuropsychiatric disturbances common in dementia: delusions, hallucinations, agitation, dysphoria, anxiety, apathy, irritability, euphoria, disinhibition, aberrant motor behaviour, night-time behaviour disturbances, and appetite and eating abnormalities. The DAD is a caregiver-based interview instrument used to evaluate instrumental and basic activities of daily living in dementia (hygiene, dressing, undressing, continence, eating, meal preparation, telephoning, going on an outing, finance, correspondence, medication, leisure and housework). The NPI also assesses the amount of caregiver distress engendered by each of the neuropsychiatric disorders. Blood is drawn and levels of EPA, DHA and phopsholipids is measured.

#### **EQUIVALENTS**

[0146] Those skilled in the art will recognize, or be able to ascertain, using no more than routine experimentation, numerous equivalents to the specific embodiments described specifically herein. Such equivalents are intended to be encompassed in the scope of the following claims.

#### **CLAIMS**

1. A concentrated therapeutic phospholipid composition comprising compounds of the Formula I:

$$H_2C - O - R_1$$
 $R_2 - O - CH O$ 
 $H_2C - O - P - O - X$ 
 $O - CH O - P - O - X$ 
 $O - CH O - P - O - X$ 
 $O - CH O - P - O - X$ 

wherein for each compound of Formula I in the composition:

each R<sub>1</sub> is independently selected from hydrogen or any fatty acid;

each R<sub>2</sub> is independently selected from hydrogen or any fatty acid;

 $\mbox{wherein at least one of } R_1 \mbox{ and } R_2 \mbox{ in each compound of Formula I is a fatty} \\ \mbox{acid; and}$ 

each X is independently selected from -CH<sub>2</sub>CH<sub>2</sub>NH<sub>3</sub>, -CH<sub>2</sub>CH<sub>2</sub>N(CH<sub>3</sub>)<sub>3</sub> or

wherein the total amount of the compounds of Formula I in the composition are at a concentration of between 45% (w/w) to about 99% (w/w).

2. The composition of claim 1, wherein the compounds of Formula I are in a concentration of between about 50% (w/w (phospholipids/total composition)) up to 70% (w/w (phospholipids/total composition)).

- 3. The composition of claim 2, wherein the compounds of Formula I are in a concentration of between about 60% (w/w (phospholipids/total composition)) to up to 70% (w/w (phospholipids/total composition)).
- 4. The composition of claim 3, wherein the compounds of Formula I are in a concentration of about 66% (w/w (phospholipids/total composition)).
- 5. The composition of claim 1, wherein the compounds of Formula I are in a concentration of above 70% (w/w (phospholipids/total composition)) to about 99% (w/w (phospholipids/total composition)).
- 6. The composition of claim 5, wherein the compounds of Formula I are in a concentration of between about 80% (w/w (phospholipids/total composition)) to about 95% (w/w (phospholipids/total composition)).
- 7. The composition of claim 6, wherein the compounds of Formula I are in a concentration of about 90% (w/w (phospholipids/total composition)).
- 8. The composition of claim 1, wherein both R1 and R2 are each independently an omega 3 fatty acid.
- 9. The composition of claim 1, wherein R1 is DHA.
- 10. The composition of claim 1, wherein R2 is DHA.
- 11. The composition of claim 1, wherein R1 is EPA.
- 12. The composition of claim 1, wherein R2 is EPA.
- 13. The composition of claim 1, wherein R2 is DHA and R1 is EPA.
- 14. The composition of claim 1, wherein R2 is EPA and R1 is EPA.

15. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition has about the same amount of EPA and DHA.

- 16. The concentrated therapeutic phospholipid composition of claim 1, wherein the the composition has predominantly DHA at the R2 position of Formula I.
- 17. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition has more DHA than EPA.
- 18. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition has greater than 60% DHA of the total fatty acids.
- 19. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition has greater than 70% DHA of the total fatty acids.
- 20. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition has greater than 80% DHA of the total fatty acids.
- 21. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition has greater than 90% DHA of the total fatty acids.
- 22. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition has greater than 95% DHA of the total fatty acids.
- 23. The concentrated therapeutic phospholipid composition of claim 1, wherein the free fatty acid concentration is between about 0% (w/w) and about 20% (w/w).
- 24. The concentrated therapeutic phospholipid composition of claim 23, wherein the free fatty acid concentration of between about 5% (w/w) and about 17% (w/w).
- 25. The concentrated therapeutic phospholipid composition of claim 24, wherein the free fatty acid concentration of between about 10% (w/w) and about 15% (w/w).
- 26. The concentrated therapeutic phospholipid composition of claim 1, wherein the triglyceride concentration is between below about 5%.

27. The concentrated therapeutic phospholipid composition of claim 1, wherein the triglyceride concentration is about 0%.

- 28. The composition of claim 1, further comprising an antioxidant.
- 29. The composition of claim 28, wherein the antioxidant is selected from astaxanthin, a carotenoid, and a flavonoid.
- 30. The composition of claim 29, wherein the antioxidant is a carotenoid.
- 31. The composition of claim 30, wherein the caretinoid is pro-vitamin A.
- 32. The composition of claim 29, wherein the antioxidant is a flavonoid.
- 33. The composition of claim 32, wherein the flavonoid is selected from naringin, naringenin, hesperetin/kaempferol, rutin, luteolin, neohesperidin, quecertin.
- 34. The composition of claim 32, wherein the flavonoid is

- 35. The composition of claim 1, wherein the antioxidant is astaxanthin.
- 36. The composition of claim 35, wherein the concentration of astaxanthin is greater than 2000 mg/kg (w/w of composition).

37. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition comprises:

compounds of Formula I are at a concentration of about 66% (w/w (phospholipids/total composition);

a free fatty acid (FFA) concentration of less than \_\_\_\_\_% (w/w FFA/total composition); and

a triglyceride concentration of 0%.

38. The concentrated therapeutic phospholipid composition of claim 1, wherein the composition comprises:

compounds of Formula I are at a concentration of above 70% (w/w (phospholipids/total composition);

- a free fatty acid (FFA) concentration of 0%; and
- a triglyceride concentration of 0%.
- 39. The concentrated therapeutic phospholipid composition of claim 38, wherein the composunds of Formula I are at a concentration of about 90%.
- 40. A method of treating or preventing a cardiometabolic disorder / metabolic syndrome, the method comprising administering to a subject in need thereof a composition of claim 1.
- 41. The method of claim 40, wherein the cardiometabolic disorder / metabolic syndrome is selected from atherosclerosis, dyslipidemia, hypertriglyceridemia, hypertension, heart failure, cardiac arrhythmias, low HDL levels, high LDL levels, stable angina, coronary heart disease, acute myocardial infarction, secondary prevention of myocardial infarction, cardiomyopathy, endocarditis, type 2 diabetes, insulin resistance, impaired glucose tolerance, hypercholesterolemia, stroke, hyperlipidemia, hyperlipoprotenemia, chronic kidney disease, intermittent claudication, hyperphosphatemia, omega-3 deficiency, phospholipid deficiency, carotid atherosclerosis, peripheral arterial disease, diabetic nephropathy,

hypercholesterolemia in HIV infection, acute coronary syndrome (ACS), non-alcoholic fatty liver disease/non-alcoholic steatohepatitis (NAFLD/NASH), arterial occlusive diseases, cerebral atherosclerosis, arteriosclerosis, cerebrovascular disorders, myocardial ischemia, coagulopathies leading to thrombus formation in a vessel and diabetic autonomic neuropathy.

- 42. A method of treating, preventing, or improving cognition and /or a cognitive disease, disorder or impairment (memory, concentration, learning (deficit)), the method comprising administering to a subject in need thereof a composition of claim 1.
- 43. The method of claim 42, wherein the cognitive disease, disorder or impairment is selected from Attention Deficit Disorder (ADD), Attention Deficit Hyperactivity Disorder (ADHD), dyslexia, age-associated memory impairment and learning disorders, amnesia, mild cognitive impairment, cognitively impaired non-demented, pre-Alzheimer's disease, autism, dystonias and Tourette syndrome, dementia, age related cognitive decline, cognitive deterioration, moderate mental impairment, mental deterioration as a result of ageing, conditions that influence the intensity of brain waves and/or brain glucose utilization, stress, anxiety, concentration and attention impairment, mood deterioration, general cognitive and mental well being, neurodegenerative disorders, hormonal disorders or any combinations thereof. In a specific embodiment, the cognitive disorder is memory impairment.
- 44. A method of inhibiting, preventing, or treating a neurodegenerative disorder, the method comprising administering to a subject in need thereof a composition of claim 1.
- Alzheimer's disease, Pick's disease, Lewy body dementia Basal ganglia—Huntington's disease, Parkinson's disease, dentatorubropallidoluysian atrophy, Freidreich's ataxia, multiple system atrophy, types 1, 2, 3, 6, 7 spinocerebellar ataxia Motor—amyotrophic lateral sclerosis, familial spastic paraparesis, spinal muscular atrophy, spinal and bulbar muscular atrophy, Lou Gehrig's disease, pre-dementia syndrome, Lewy body dementia, age-related cognitive decline, cognitive deterioration, moderate mental impairment, mental deterioration as a result of ageing, dentatorubropallidoluysian atrophy, Freidreich's ataxia, multiple system atrophy, types 1, 2, 3, 6, 7 spinocerebellar ataxia, amyotrophic lateral sclerosis, and familial spastic paraparesis.

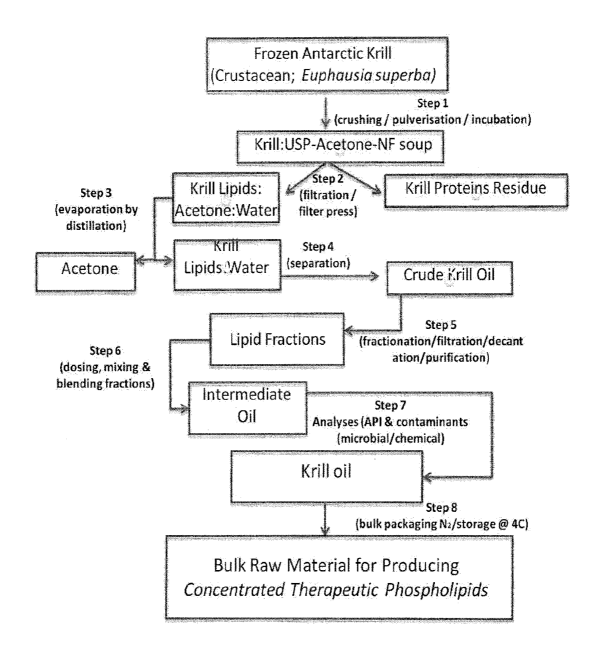


FIGURE 1A

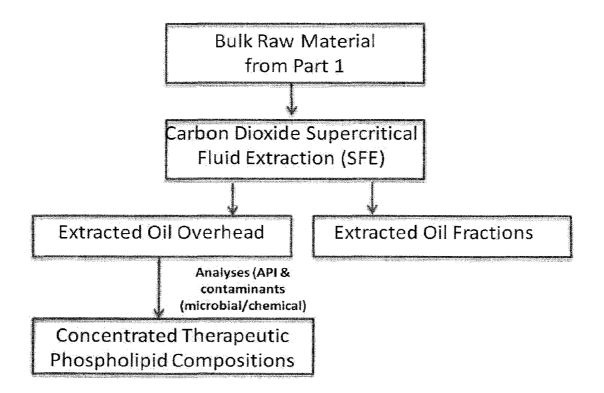


FIGURE 1B

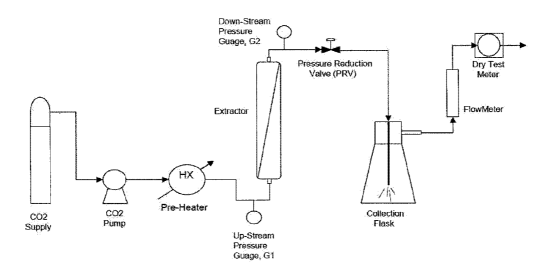


FIGURE 1C

# Circulating plasma Triglyceride concentration of C57BL/6 mice

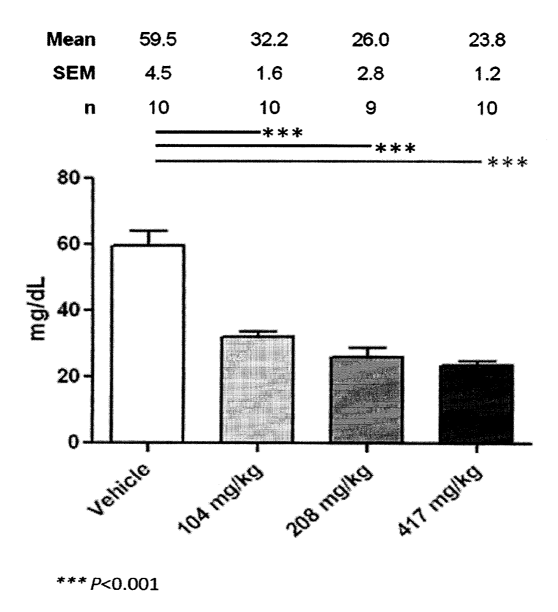


FIGURE 2

Circulating plasma HDL-Cholesterol concentration of C57BL/6 mice

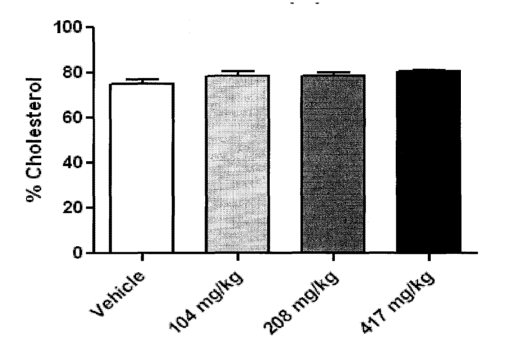
N	lean	53.4	59.4	63.4	62.3
\$	SEM	3.2	1.9	3.5	2.4
	n	10	10	10	10
₩Q/QM	<sup>80</sup> 7	<del> </del>			*
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FIGURE 3

Circulating plasma percentage of HDL-Cholesterol in C57BL/6 mice

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Mean	74.9	78.4	78.4	80.6
SEM	2.1	2.2	1.7	0.6
n	10	10	10	9



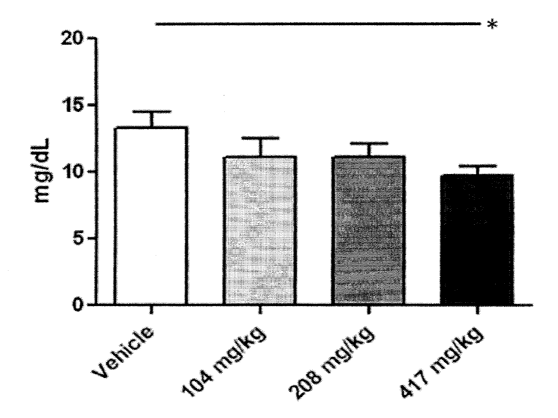
\* P<0.05

FIGURE 4

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Circulating plasma LDL-Cholesterol concentration of C57BL/6 mice

Mean	13.3	11.1	11.2	9.6
SEM	1.2	1.4	1.0	0.7
n	10	10	10	9



\* P<0.05

FIGURE 5

Circulating plasma percentage of LDL-Cholesterol in C57BL/6 mice

N	lean	18.9	14.5	14.1	12.2
5	SEM	1.8	1.7	1.4	0.5
	n	10	10	10	9
	0E			*	**
	<sup>25</sup> 7				
ठ	20-				
ester	15-				
% Cholesterol	10-				
8	5-				
	07				
	*	Jehicle 10	A MUSIKES	THEIKE AN	Ingka

\* P<0.05; \*\*P<0.01

FIGURE 6

Circulating plasma NEFA concentration of C57BL/6 mice

Mean	2.92	2.65	2.93	2.63
SEM	0.10	0.06	0.12	0.09
n	10	9	9	9
		*		<del></del> *
4 3 1/2 1 1 0 3	Ehicle . O		TOTAL STATE OF THE	malkes

\*P<0.05

FIGURE 7

10/37
Circulating plasma Glucose concentration of C57BL/6 mice

Mean	177.0	107.0	167.6	157.3
SEM	8.4	6.3	5.9	12.0
n	10	10	9	10
		***		
<sup>200</sup> 7				
150-				
100-				
50-			Control of the second	
01			THE PARTY NAMED IN COLUMN TWO IS NOT THE PARTY N	
7	Jehicle N	JA MUJIKU 208	THEIRES AN	7 mg/kg
	SEM n 200 - 150 -	SEM 8.4 n 10  200 150- 100-	SEM 8.4 6.3  n 10 10	SEM 8.4 6.3 5.9  n 10 10 9

FIGURE 8

\*\*\* P<0.001

11/37

## Circulating plasma Phospholipid concentration of C57BL/6 mice

M	ean	141.9	140.5	147.4	138.6
S	SEM	7.2	4.1	5.5	5.4
	n	10	10	9	10
	200				
•	150 -				_
mg/dl	100-				
	50 -				
	$L_{o}$	1 , 1			
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		^	ري د الم	D	

FIGURE 9

12/37

### Circulating plasma ALT concentration of C57BL/6 mice

Mean	36.1	31.0	26.9	27.8
SEM	3.9	3.5	1.8	1.2
n	10	10	10	10

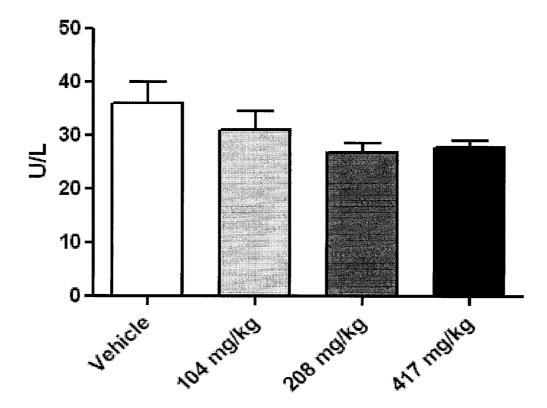


FIGURE 10

Liver Total Cholesterol concentration of C57BL/6 mice

	lean SEM	23.1 0.8	23.3 0.6	25.8 0.3	24.1 0.8
	n	10	10	9	10
	307			**	
ธิพ/ธิ <del>ป</del>	10-				
	⊥ <sub>0</sub>	enicle 40	JA MIGHKS 208	1467 - 12 B	Trigiko

\*\* P<0.01

FIGURE 11

14/37

Liver Triglyceride concentration of C57BL/6 mice

ean	53.0	58.0	67.4	66.3
EM	3.1	1.7	2.6	5.4
n	10	10	9	10
	**************************************		**	*
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	80 7 60 -	80 3.1 60 - 40 - 20 - 20 - 20 - 20 - 20 - 20 - 2	80 - 10 10 10 10 10 10 10 10 10 10 10 10 10	EM 3.1 1.7 2.6  n 10 10 9

\* P<0.05;\*\* P<0.01

FIGURE 12

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## Circulating plasma Triglyceride concentration of LDLr KO mice

M	ean	65.1	45.7	41.7	36.7
S	EM	3.8	2.8	4.9	1.6
	n	9	10	10	10
			***	**	***
	<sup>80</sup> 7				
_	60-				
	40-				
	20-				
	$_{0}\bot$			80 7 7 7 7 7 7	
	16	Enicle	OA MISIKO	malked A	7 mg/kg

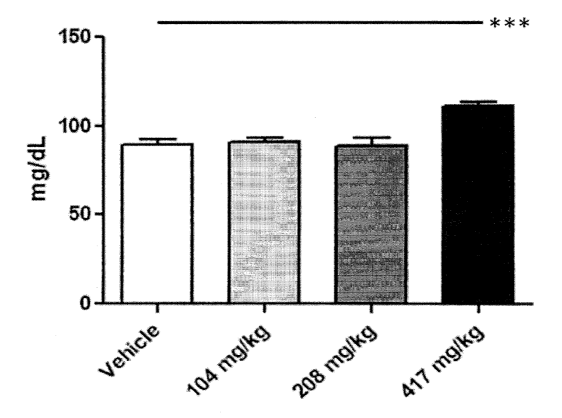
\*\* *P*<0.01; \*\*\**P*<0.001

FIGURE 13

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Circulating plasma HDL-Cholesterol concentration of LDLr KO mice

Mean	88.8	90.9	88.4	111.2
SEM	3.6	2.5	4.6	2.5
n	9	9	9	10

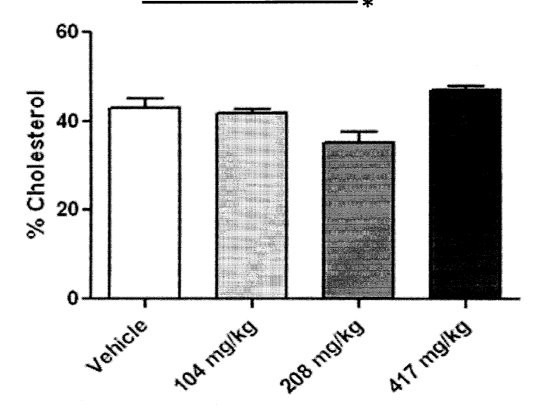


\*\*\*P<0.001

FIGURE 14

Circulating plasma percentage of HDL-Cholesterol in LDLr KO mice

Mean	43.0	41.9	35.1	46.8
SEM	2.1	0.8	2.3	1.1
n	10	9	10	10



\*P<0.05

FIGURE 15

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### Liver Total Cholesterol concentration of LDLr KO mice

Mean	23.6	24.5	25.8	24.9
SEM	0.6	0.3	0.9	0.6
n	10	9	10	10

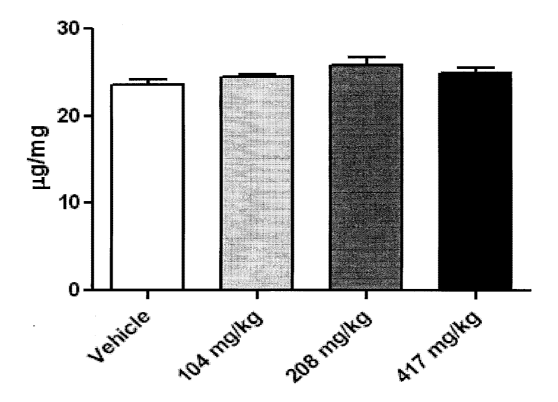


FIGURE 16

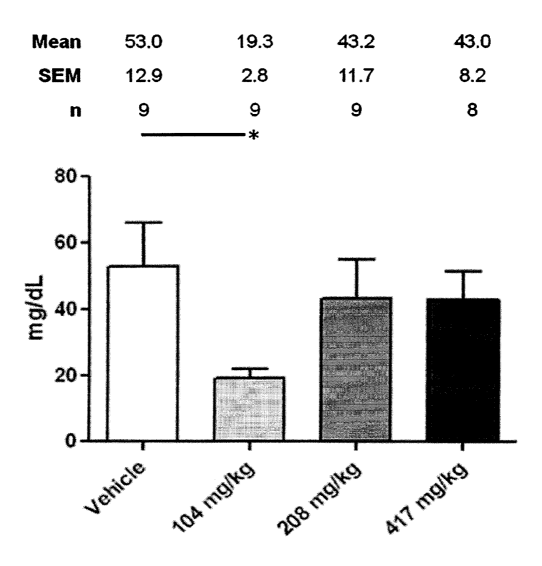
Liver Triglyceride concentration of LDLr KO mice

Me	ean	42.9	54.3	52.4	58.4
SI	EM	2.1	2.1	3.0	4.5
	n	10	9	9	10
		***************************************	**	*	
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		<b>*</b>	a do	<b>N</b>	•

\* P<0.05; \*\*P<0.01

FIGURE 17

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Circulating plasma Triglyceride concentration of ApoA-1 CETP
Tg mice



\* P<0.05

FIGURE 18

# Circulating plasma total cholesterol concentration of adult male SD, ZDF, SHR & JCR:LA rats

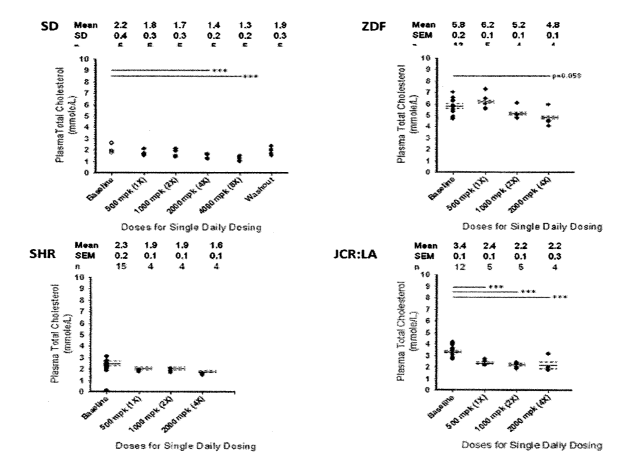
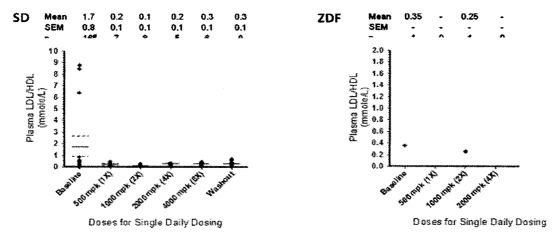


FIGURE 19

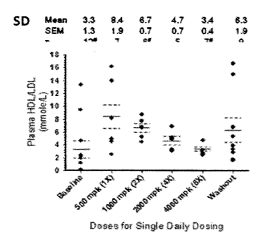
# Circulating plasma LDL/HDL concentration of adult male SD, ZDF, SHR & JCR:LA rats

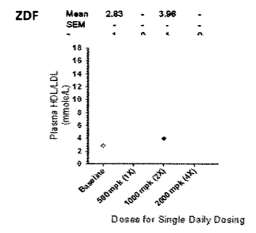


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FIGURE 20

# Circulating plasma HDL/LDL concentration of adult male SD, ZDF, SHR & JCR:LA rats

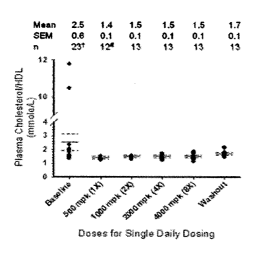


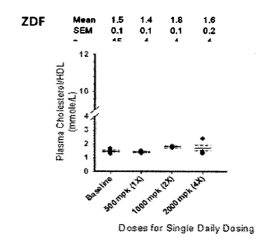


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FIGURE 21

# Circulating plasma total cholesterol/HDL concentration of adult male SD, ZDF, SHR & JCR:LA rats





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- †, 2 values removed

FIGURE 22

### Prothrombin Time of adult male SD, ZDF, SHR & JCR:LA rats

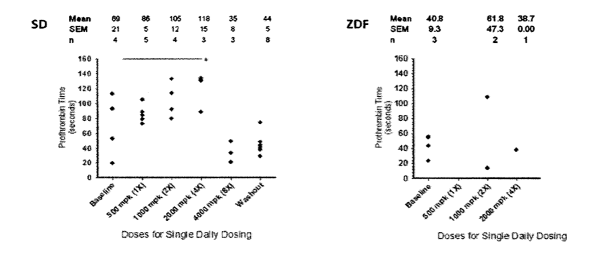
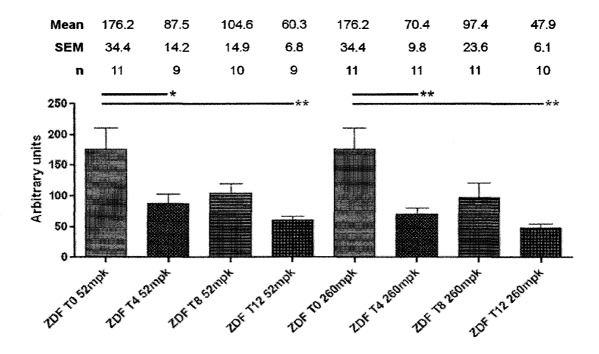


FIGURE 23

OGTT area under the curve data in ZDF male rats treated with Composition 3 for 90 days



\* P<0.05; \*\*P<0.01

FIGURE 24

OGTT area under the curve data in ZDF male rats treated with Composition 3 for 90 days

Mean	176.2	87.5	104.6	60.3	176.2	70.4	97.4	47.9
SEM	34.4	14.2	14.9	6.8	34.4	9.8	23.6	6.1
n	11	9	10	9	11	11	11	10
500 <sub>7</sub>	***************************************	*		**	***************************************			**
400	*				•			
Arbitrary units	٥				•		*	
를 200-	<b></b>		<b>A</b>					
₹ 100- 0	•	*****	***	****	***		· <u>\$</u>	-
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\* *P*<0.05; \*\**P*<0.01

FIGURE 25

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OGTT area under the curve data in SD male rats treated with Composition 3 for 90 days

Mean	63.1	52.5	51.1	50.7	63.1	86.5	65.9	66.6
SEM	8.0	7.8	6.0	5.9	8.0	21.8	7.5	7.7
n	12	6	6	6	12	6	6	6

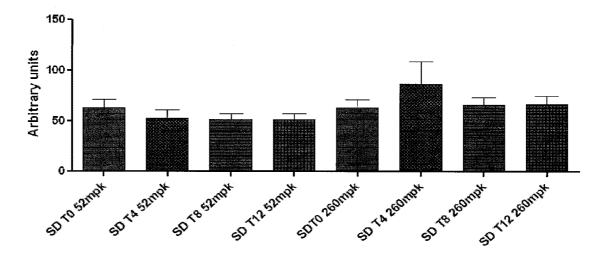


FIGURE 26

OGTT area under the curve data in SD male rats treated with Composition 3 for 90 days

Mean	63.1	52.5	51.1	50.7	63.1	86.5	65.9	66.6
SEM	8.0	7.8	6.0	5.9	8.0	21.8	7.5	7.7
n	12	6	6	6	12	6	6	6

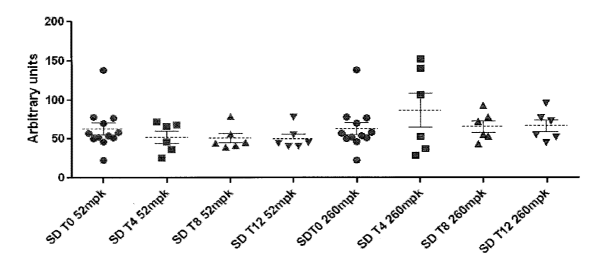


FIGURE 27

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## Effects of Composition 3 on lipid biomarkers in male ZDF rats compared to agematched controls

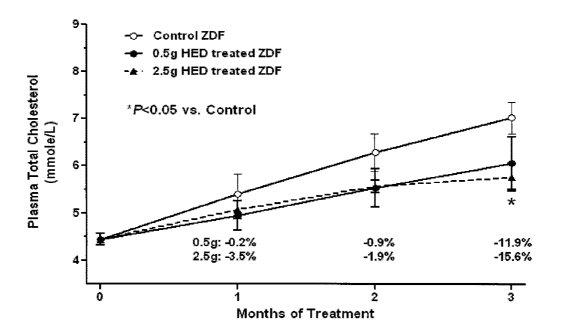


FIGURE 28

## Effects of Composition 3 on lipid biomarkers in male ZDF rats compared to agematched controls

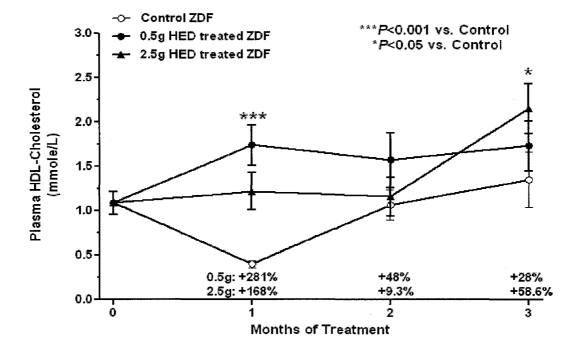


FIGURE 29

Effects of Composition 3 on lipid biomarkers in male ZDF rats compared to agematched controls

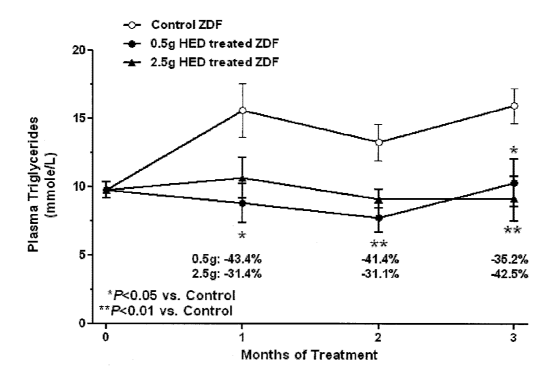


FIGURE 30

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#### Effects of Composition 3 on Glucose Intolerance in male ZDF rats

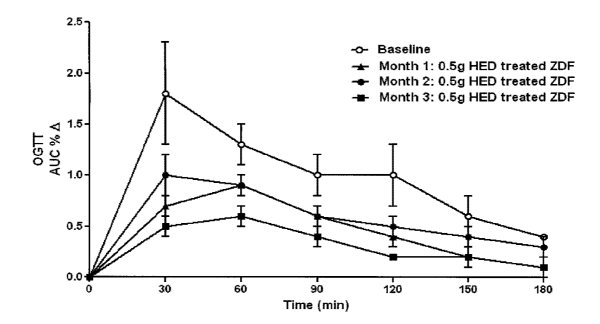


FIGURE 31

#### Effects of Composition 3 on Glucose Intolerance in male ZDF rats

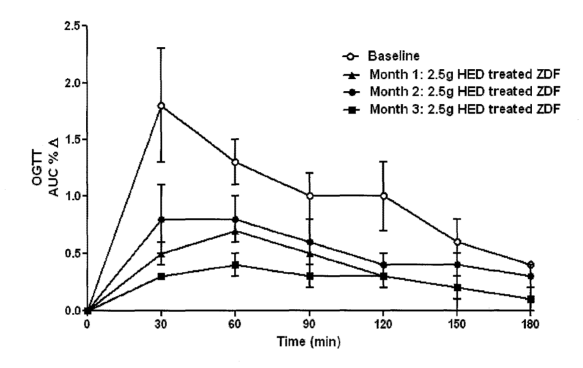


FIGURE 32

#### Effects of Composition 3 on Glucose Intolerance in male SD rats

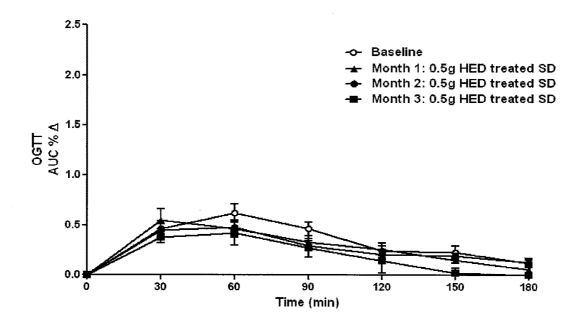


FIGURE 33

#### Effects of Composition 3 on Glucose Intolerance in male SD rats

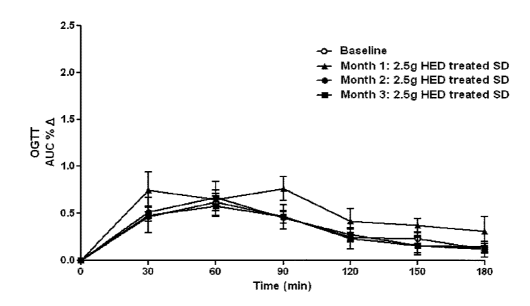
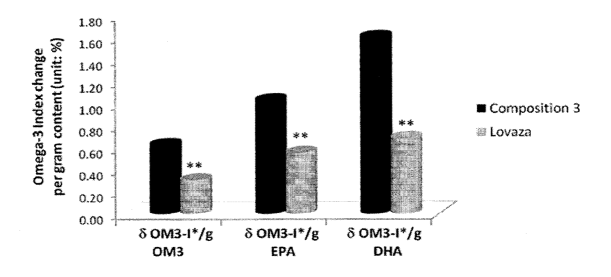


FIGURE 34

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# Comparative Effects of Composition 3 and Lovaza® on Omega-3 Index



	∆0M3I/G 0 <b>M</b> 3	∆OM3I/G EPA	∆OM3I/G DHA
Comp. 3 1g	62.85%	103.26%	160.62%
Lovaza 4g	30.65%	55.37%	68.65%
Δ Comp. 3 vs Lovaza	105.70%	86.49%	133.95%

FIGURE 35

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/CA2010/001720

#### A. CLASSIFICATION OF SUBJECT MATTER

IPC: A61K 31/683 (2006.01), A61P 3/00 (2006.01), A61P 9/00 (2006.01)

According to International Patent Classification (IPC) or to both national classification and IPC

#### **B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

A61K 31/683 (2006.01), A61K 31/685 (2006.01), A61P 3/00 (2006.01), A61P 9/00 (2006.01)

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic database(s) consulted during the international search (name of database(s) and, where practicable, search terms used) EPOQUE, Canadian Patent Database, PubMed

phospholipid, DHA, EPA, krill, squid, marine, fish supercritical fluid extraction, dyslipidemia, hyperlipidemia, hypertriglycemia, diabetes

#### C. DOCUMENTS CONSIDERED TO BE RELEVANT

Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
WO2008117062A1 (AKER BIOMARINE ASA) 02 -10-2008 ( 02 October 2008) tables 18B, 19B, 20B, 21, 22; tables 10, 13-15; tables 16 and 19C. examples 9, 12	1-5, 8-16, 23-27, 28-35 40-41
WO03011873 A2 (NEPTUNE TECHNOLOGIES & BIORES et al) 13-02-2003 (13 February 2003) p.26, 1. 3-6; table 5; example 1, Sample #804; p.30, 1.1-7; table 5. example 3	1-3, 8-16, 23-24, 26, 28-35 42-45
WO02102394A2 (SAMPALIS) 27-12-2002 (27 December 2002) whole document	1-45
	WO2008117062A1 (AKER BIOMARINE ASA) 02 -10-2008 ( 02 October 2008) tables 18B, 19B, 20B, 21, 22; tables 10, 13-15; tables 16 and 19C. examples 9, 12  WO03011873 A2 (NEPTUNE TECHNOLOGIES & BIORES et al) 13-02-2003 (13 February 2003) p.26, 1. 3-6; table 5; example 1, Sample #804; p.30, 1.1-7; table 5. example 3  WO02102394A2 (SAMPALIS) 27-12-2002 (27 December 2002)

[X] F	urther documents are listed in the continuation of Box C.	[X]	See patent family annex.
* "A"	Special categories of cited documents:  document defining the general state of the art which is not considered to be of particular relevance	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"E"	earlier application or patent but published on or after the international filing date	Λ	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
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"O"	document referring to an oral disclosure, use, exhibition or other means	"&"	•
"P"	document published prior to the international filing date but later than the priority date claimed	··&··	document member of the same patent family
Date o	f the actual completion of the international search	Date of	of mailing of the international search report
1 Febr	uary 2010 (01-02-2001)	1 Mar	ch 2011 (01-03-2011)
	and mailing address of the ISA/CA ian Intellectual Property Office	Autho	rized officer
50 Vi	du Portage I, C114 - 1st Floor, Box PCT toria Street	Yong	g-Huang Chen (819) 956-4113
	au, Quebec K1A 0C9 nile No.: 001-819-953-2476		

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Page 3 of 6

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/CA2010/001720

Box I	No.	II	Observations where certain claims were found unsearchable (Continuation of item 2 of the first sheet)
This i		rnational	search report has not been established in respect of certain claims under Article 17(2)(a) for the following
1. [	X]	Claim N	os.: 40-45
•	_		they relate to subject matter not required to be searched by this Authority, namely:
		searched	10-45 are directed to a method for treatment of the human or animal body by surgery or therapy, are not required to be nor is a written opinion required by this Authority. Regardless, this Authority has established a written opinion based on ed therapeutic effect or purpose/use of the product defined in claim 1.
2. [	]	Claim N	08. :
			they relate to parts of the international application that do not comply with the prescribed requirements to such an extent neaningful international search can be carried out, specifically:
3. [	]	Claim N	os.: they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box N			Observations where unity of invention is lacking (Continuation of item 3 of first sheet)  Searching Authority found multiple inventions in this international application, as follows:
1. [	]		quired additional search fees were timely paid by the applicant, this international search report covers all le claims.
2. [	]		archable claims could be searched without effort justifying additional fees, this Authority did not invite of additional fees.
3. [	]		some of the required additional search fees were timely paid by the applicant, this international search report nly those claims for which fees were paid, specifically claim Nos.:
4. [	]	No requi	red 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 claim Nos.:
		Rema	rk on Protest [ ] The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.
			[ ] The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.
			[ ] No protest accompanied the payment of additional search fees.

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Page 2 of 6

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/CA2010/001720

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO0023546A1 (UNIV SHERBROOKE et al) 27-04-2000 (27 April 2000) table 14; tables 15, 16; tables 17, 18.	1-2, 8-16, 28-35
X	WO2008060163A1 (PRONOVA BIOPHARMA NORGE AS) 22-05-2008 (22 May 2008) tables 3-5; p.20, i.5-9.	1-7, 8-16, 23-25, 28-35
X X	JP2215351A (MARUYAMA et al) 28-08-1990 (28 August 190) p.327, right column, para.4; also JPO machine translation. example 2	1-16, 23-27, 37-39 42-45
x	JP2000060432A (NIPPON KAGAKU SHIRYO KK) 29-02-2000 (29 February 2000) para.[009]; example 1.	1-6, 8-20

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Page 4 of 6

#### INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No. PCT/CA2010/001720

	-		<u></u>
Patent Document	Publication	Patent Family	Publication
Cited in Search Report	Date	Member(s)	Date
WO2008117062A1	02 October 2008 (02-10-2008)	AU2008231570A1 CA2682068A1 EP2144618A1 US2008274203A1	02 October 2008 (02-10-2008) 02 October 2008 (02-10-2008) 20 January 2010 (20-01-2010) 06 November 2008 (06-11-2008)
			<del>.</del>
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		WO03011873A8	07 August 2003 (07-08-2003)
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		JP2004534800T JP2010090140A JP2010090141A	18 November 2004 (18-11-2004) 22 April 2010 (22-04-2010) 22 April 2010 (22-04-2010)
		NO20035618D0 PT1406641E US2004241249A1 US2007098808A1	16 December 2003 (16-12-2003) 15 April 2009 (15-04-2009) 02 December 2004 (02-12-2004) 03 May 2007 (03-05-2007)
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		EP1123368A1 EP1123368B1 ES2306527T3	16 August 2001 (16-08-2001) 09 April 2008 (09-04-2008) 01 November 2008 (01-11-2008)
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WO2008060163A1	22 May 2008 (22-05-2008)	AR064250A1 AU2007320183A1 CA2669847A1 CL32832007A1 CN101652462A EP2094823A1 EP2094823A4 JP2010510208T KR20090085682A MX2009005227A NO20092310A PE10602008A1 US2010143571A1 WO2008060163A9 ZA200904176A	25 March 2009 (25-03-2009) 22 May 2008 (22-05-2008) 22 May 2008 (22-05-2008) 23 May 2008 (23-05-2008) 17 February 2010 (17-02-2010) 02 September 2009 (02-09-2009) 02 February 2011 (02-02-2011) 02 April 2010 (02-04-2010) 07 August 2009 (07-08-2009) 28 May 2009 (28-05-2009) 16 June 2009 (16-06-2009) 08 September 2008 (08-09-2008) 10 June 2010 (10-06-2010) 02 April 2009 (02-04-2009) 25 August 2010 (25-08-2010)
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JP2000060432A	29 February 2000 (29-02-2000)	None	

Doc code: IDS Doc description: Information Disclosure Statement (IDS) Filed PTO/SB/08a (03-15)
Approved for use through 07/31/2016. OMB 0651-0031
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INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Application Number		15180431		
	Filing Date		2016-06-13		
	First Named Inventor Inge Br		Bruheim		
	Art Unit		1672		
	Examiner Name	CUTL	LIFF, YATE KAI RENE		
	Attorney Docket Number	er	AKBM-14409/US-12/CON		

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Filing Date		2016-06-13
First Named Inventor	Inge E	Bruheim
Art Unit		1672
Examiner Name	CUTL	IFF, YATE KAI RENE
Attorney Docket Numb	er	AKBM-14409/US-12/CON

	Action Closing Prosecution, 348 Patent, mailed May 14, 2013
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;	Certified translation of Ex. 1074: Japanese Patent No. 60-153779, entitled "Nutritional Supplement" ("Fukuoka"); Certificate of Translation provided as Ex. 1075, dated August 16, 2013, 1 page
4	Certificate of translation of Ex. 1076: Japanese Patent Publication No. H08-231391, entitled "Medicine for mprovement of Dementia Symptoms" ("Yasawa"); Certificate of Translation provided as Ex. 1077, dated August 16, 2013, 1 page
	Certification of translation of Ex. 1070: Japanese Unexamined Patent Application Publication No. 02-215351, titled Krill Phospholipids Fractioning Method ("Maruyama,"); Certificate of Translation provided as Ex. 1071; dated July 9, 2013, 1 page
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8	Third Party Observation against corresponding AU Patent Application No. 2014256345, filed May 23, 2016, 50 pages
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	Evidence in Support of Opposition, AU Patent Application No. 2013227998, filed September 22, 2016, 22 pages
	Notice of Acceptance of Application, AU Patent Application No. 2013227998, mailed October 5, 2016, 2 pages
	1

Application Number		15180431
Filing Date		2016-06-13
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Application Number		15180431
Filing Date		2016-06-13
First Named Inventor	Inge E	Bruheim
Art Unit		1672
Examiner Name CUTL		IFF, YATE KAI RENE
Attorney Docket Number	er	AKBM-14409/US-12/CON

#### **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).

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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.

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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)	2016-10-12
Name/Print	J. Mitchell Jones	Registration Number	44174

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Application Number:	15	180431					
Filing Date:	13-Jun-2016						
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS						
First Named Inventor/Applicant Name:	Ing	e Bruheim					
Filer:	Joł	nn Mitchell Jones					
Attorney Docket Number:	AK	BM-14409/US-12/C	ON				
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Filing Fees for Utility under 35 USC 111(a)							
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:							
Extension-of-Time:							

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Miscellaneous:				
Submission- Information Disclosure Stmt	1806	1	180	180
	Tot	al in USD	(\$)	180

Electronic Ack	Electronic Acknowledgement Receipt					
EFS ID:	27186798					
Application Number:	15180431					
International Application Number:						
Confirmation Number:	2763					
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS					
First Named Inventor/Applicant Name:	Inge Bruheim					
Customer Number:	72960					
Filer:	John Mitchell Jones/Mallory Checkett					
Filer Authorized By:	John Mitchell Jones					
Attorney Docket Number:	AKBM-14409/US-12/CON					
Receipt Date:	12-OCT-2016					
Filing Date:	13-JUN-2016					
Time Stamp:	14:29:31					
Application Type:	Utility under 35 USC 111(a)					

### **Payment information:**

Submitted with Payment	yes
Payment Type	Deposit Account
Payment was successfully received in RAM	\$180
RAM confirmation Number	859
Deposit Account	504302
Authorized User	Jones, J. Mitchell

The Director of the USPTO is hereby authorized to charge indicated fees and credit any overpayment as follows:

File Listing	y:				
Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl
			91390		
1		161007_ROA1.pdf	01ecddcd8bcd6cc294598046fda2de40f90 9b0ef	yes	4
	Multip	! part Description/PDF files in .	zip description		
	Document De	scription	Start	Eı	nd
	Amendment/Req. Reconsiderati	1		1	
	Claims	2		3	
	Applicant Arguments/Remarks	Made in an Amendment	4		4
Warnings:					
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			4665035		
2	Foreign Reference	WO2011050474.pdf	5a1c37295df398f7a0dcaa8c0bc0f0fd14971 15b	no	95
Warnings:					
Information:					
	Oth ou Defended Detect / App / Secure	Astion Clasin - Dunas suttion 240	24322856		
3	Other Reference-Patent/App/Search documents	ActionClosingProsecution_348 patent_PART1.pdf	8699cc6595609b800e3fa2d8a452a99f4917 e73b	no	43
Warnings:					
Information:					
			10631681		
4	Other Reference-Patent/App/Search documents	ActionClosingProsecution_348 patent_PART2.pdf	65d1a0435f40c39e36d2077d4f191c34fafe bf51	no	21
Warnings:					
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			1450631		
5	Other Reference-Patent/App/Search documents	ActionClosingProsecution_348 patent_PART3_uspto.pdf	d256d0f5c5600d76327393d80f1a4e8ad2d 96fa8	no	21
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6	Non Patent Literature	BUCHI_Rotovapor_manual_usp	3201325	no	50
		to.pdf	dbfed7bd15d606281bd9a572f405d9bba2 837237		
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7	Non Patent Literature	Certified Translation_FUKUOKA.	1030925	no	5
,	Non Faterit Literature	pdf	914fe24b1316cdd30520cb6428f06c51313 06e95	110	3
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8	Non Patent Literature	pdf	c786df61c5954523a4ad497a21b74ab3222 bf462	no	9
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9	Non Patent Literature	Translation_uspto.pdf	29656443c175797084f343861f01e83fb3ce 5861	no	6
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		Haugsgjerd_Decl_351Patent.	6747221		
10	Non Patent Literature	pdf	ae6ac8409c153468021532be74f9b47965d 16f2a	no	12
Warnings:					
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			32615		
11	Non Patent Literature	FSTA_Shibata_abstract.pdf	e29068a81bef6233f4d9abe88f6c9299b776 4a91	no	1
Warnings:					
Information:					
	Othor Pofore and Patent I American	All Fuldon and Court of 20222	10659074		
12	Other Reference-Patent/App/Search documents	AU_EvidenceinSupport_20132 27998_9-22-2016.pdf	d0fba17e2a2f28c3dcba8bb18109da46dd4 0490e	no	58
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15 Warnings:	Form (SB08)	16pdf	938e6190c630ba17df29196bd2847fff9d41 ecec	no	5
Information Disclosure Statement (ID		14409US12CON_IDS_10-12-20	1036288		
Information:	<u> </u>				
Warnings:					
14	Other Reference-Patent/App/Search documents	AU_ThirdPartyObservation_20 13227998_7-15-2016.pdf	1160150 139d9cdfbc7a8dcb779b830bd0a9b5340d2a 4d825	no	6
Warnings: Information:	:				
***			e8daf		
13	Other Reference-Patent/App/Search documents	AUNoticeofAcceptance_20132 27998_10-5-2016.pdf	164760	no	2

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

#### 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.

U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE
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P	ATENT APPL	ERMINATION TO-875	N RECORD		or Docket Numb /180,431		Filing Date 06/13/2016	To be Mailed		
							ENTITY:	⊠ LA	RGE SMA	LL MICRO
				APPLICA	ATION AS FIL	ED – PAR	ТІ			1
			(Column <sup>-</sup>	1)	(Column 2)					
	FOR	N	UMBER FIL	_ED	NUMBER EXTRA		RATE (\$	S)	F	EE (\$)
	BASIC FEE (37 CFR 1.16(a), (b), o	or (c))	N/A		N/A		N/A			
	SEARCH FEE (37 CFR 1.16(k), (i), c	or (m))	N/A		N/A		N/A			
	EXAMINATION FE (37 CFR 1.16(o), (p), o		N/A		N/A		N/A			
	TAL CLAIMS CFR 1.16(i))		mir	nus 20 = *			X \$ =			
	EPENDENT CLAIM CFR 1.16(h))	S	m	inus 3 = *			X \$ =	=		
	If the specification and drawings exceed 100 sheets of paper, the application size fee due is \$310 (\$155 for small entity) for each additional 50 sheets or fraction thereof. See 35 U.S.C. 41(a)(1)(G) and 37 CFR 1.16(s).					\$155 r				
	MULTIPLE DEPEN	IDENT CLAIM PF	RESENT (3	7 CFR 1.16(j))						
* If t	the difference in colu	ımn 1 is less than	zero, ente	r "0" in column 2.			TOTAL			
		(Column 1)		APPLICATION (Column 2)	ION AS AMEN (Column 3		RT II			
AMENDMENT	10/12/2016	CLAIMS REMAINING AFTER AMENDMENT		HIGHEST NUMBER PREVIOUSLY PAID FOR	PRESENT EX	TRA	RATE (\$	S)	ADDITIO	DNAL FEE (\$)
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	Independent (37 CFR 1.16(h))	* 2	Minus	***3	= 0		x \$420 =			0
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	FIRST PRESEN	ITATION OF MULTI	PLE DEPEN	DENT CLAIM (37 CFF	R 1.16(j))					
						-	TOTAL ADD'	L FEE		0
		(Column 1)		(Column 2)	(Column 3	)				
T		CLAIMS REMAINING AFTER AMENDMENT		HIGHEST NUMBER PREVIOUSLY PAID FOR	PRESENT EX	TR <b>A</b>	RATE (\$	S)	ADDITIO	DNAL FEE (\$)
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AN	FIRST PRESEN	ITATION OF MULTI	PLE DEPEN	DENT CLAIM (37 CFF	R 1.16(j))					
							TOTAL ADD'	L FEE		
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	If the "Highest Numb "Highest Number P	•				ound in the ar	opropriate box in	column	ı 1	

This collection of information is required by 37 CFR 1.16. 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 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

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#### United States Patent and Trademark Office

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APPLICATION NUMBER

15/180,431

FILING OR 371(C) DATE 06/13/2016

FIRST NAMED APPLICANT Inge Bruheim

ATTY. DOCKET NO./TITLE AKBM-14409/US-12/CON

**CONFIRMATION NO. 2763** 

**PUBLICATION NOTICE** 

72960 Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON, WI 53562

Title:BIOEFFECTIVE KRILL OIL COMPOSITIONS

Publication No.US-2016-0281026-A1 Publication Date: 09/29/2016

#### NOTICE OF PUBLICATION OF APPLICATION

The above-identified application will be electronically published as a patent application publication pursuant to 37 CFR 1.211, et seg. The patent application publication number and publication date are set forth above.

The publication may be accessed through the USPTO's publically available Searchable Databases via the Internet at www.uspto.gov. The direct link to access the publication is currently http://www.uspto.gov/patft/.

The publication process established by the Office does not provide for mailing a copy of the publication to applicant. A copy of the publication may be obtained from the Office upon payment of the appropriate fee set forth in 37 CFR 1.19(a)(1). Orders for copies of patent application publications are handled by the USPTO's Office of Public Records. The Office of Public Records can be reached by telephone at (703) 308-9726 or (800) 972-6382, by facsimile at (703) 305-8759, by mail addressed to the United States Patent and Trademark Office, Office of Public Records, Alexandria, VA 22313-1450 or via the Internet.

In addition, information on the status of the application, including the mailing date of Office actions and the dates of receipt of correspondence filed in the Office, may also be accessed via the Internet through the Patent Electronic Business Center at www.uspto.gov using the public side of the Patent Application Information and Retrieval (PAIR) system. The direct link to access this status information is currently http://pair.uspto.gov/. Prior to publication, such status information is confidential and may only be obtained by applicant using the private side of PAIR.

Further assistance in electronically accessing the publication, or about PAIR, is available by calling the Patent Electronic Business Center at 1-866-217-9197.

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UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
15/180,431	06/13/2016	Inge Bruheim	AKBM-14409/US-12/CON	2763	
72960 Casimir Jones, S	7590 07/21/201 S.C.	EXAMINER			
	WAY, SUITE 310		CUTLIFF, YATE KAI RENE		
		ART UNIT	PAPER NUMBER		
			1672		
			NOTIFICATION DATE	DELIVERY MODE	
			07/21/2016	ELECTRONIC	

### 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.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

docketing@casimirjones.com pto.correspondence@casimirjones.com

	Application No. 15/180,431	Applicant(s BRUHEIM E	
Office Action Summary	Examiner YATE' K. CUTLIFF	Art Unit 1672	AIA (First Inventor to File) Status No
The MAILING DATE of this communication app	ears on the cover sheet with the c	orresponden	ce address
Period for Reply  A SHORTENED STATUTORY PERIOD FOR REPLY THIS COMMUNICATION.  - 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).	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	nely filed the mailing date of D (35 U.S.C. § 13	of this communication. 3).
Status			
1) Responsive to communication(s) filed on 6/13/			
A declaration(s)/affidavit(s) under <b>37 CFR 1.1</b>	• • • • • • • • • • • • • • • • • • • •		
<u>'</u>	action is non-final.	a a tifa utla laborit	
3) An election was made by the applicant in responsition requirement and election	·		ng the interview on
; the restriction requirement and election  4) Since this application is in condition for allowar	·		to the merits is
closed in accordance with the practice under E	·		
Disposition of Claims*			
5) Claim(s) 1 - 20 is/are pending in the application	٦.		
5a) Of the above claim(s) is/are withdray	vn from consideration.		
6) Claim(s) is/are allowed.			
7)⊠ Claim(s) <u>1 - 20</u> is/are rejected.			
8) Claim(s) is/are objected to.			
9) Claim(s) are subject to restriction and/or			
* If any claims have been determined allowable, you may be eli			ıway program at a
participating intellectual property office for the corresponding ap	·		
http://www.uspto.gov/patents/init_events/pph/index.jsp or send	an inquiry to <u>PPHfeedback@uspto.c</u>	<u>10V</u> .	
Application Papers			
10) The specification is objected to by the Examine			
11)⊠ The drawing(s) filed on <u>6/13/2016</u> is/are: a)⊠ a			
Applicant may not request that any objection to the	• • • • • • • • • • • • • • • • • • • •		` '
Replacement drawing sheet(s) including the correcti	ion is required if the drawing(s) is ob	jected to. See	3/ CFR 1.121(d).
Priority under 35 U.S.C. § 119			
12) ☐ Acknowledgment is made of a claim for foreign	priority under 35 U.S.C. § 119(a)	)-(d) or (f).	
Certified copies:			
a) ☐ All b) ☐ Some** c) ☐ None of the:			
1. Certified copies of the priority document		tion No	
<ul><li>2. Certified copies of the priority document</li><li>3. Copies of the certified copies of the priority</li></ul>			
application from the International Bureau	- <del>-</del>	cu iii iiiis iva	lional Stage
** See the attached detailed Office action for a list of the certifie			
Attachment(s)			
1) Notice of References Cited (PTO-892)	3) Interview Summary	(PTO-413)	
2) X Information Disclosure Statement(s) (PTO/SB/08a and/or PTO/S	Paper No(s)/Mail Da 3B/08b) 4) Other:	ate	
Paper No(s)/Mail Date See Continuation Sheet.			

Continuation Sheet (PTOL-326)	Application No. 15/180,431
Continuation of Attachment(s) 2). Information Disclosure Statement(s) (PTO/SB/08), Paper No(s)/Mail Date	:6/13/2016,
6/13/2016,6/13/2016, 6/13/2016, 7/13/2016.	

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#### **DETAILED ACTION**

#### Notice of Pre-AIA or AIA Status

1. The present application is being examined under the pre-AIA first to invent provisions.

#### Status of Claims

2. Claims 1 - 20 are pending.

Claims 1 - 20 are rejected.

Claims 13 – 17, 19 and 20 are objected.

#### Information Disclosure Statement

- 3. The information disclosure statements (IDS) submitted on Jun 13, 2016 and July 13, 2016 are in compliance with the provisions of 37 CFR 1.97. Accordingly, the information disclosure statement is being considered by the examiner.
- 4. However, the Information Disclosure Statement of June 13, 2016 comprised of 13 pages May 30, 2008, did not include date information for non-patent literature documents cite no. 32, 40, 43, 44, 45, 46, 48, 49 and 50. It is for this reasons that the reverences a have a line drawn through them. Applicant is required to provide this information. 37 CFR 1.98 (b)(5).

#### Claims

5. Applicant is advised that should claims 3, 4, 5, 6, 7, 9 and 10 be found allowable, claims 13, 14, 15, 16, 17, 19 and 20 will be objected to under 37 CFR 1.75 as being a substantial duplicate thereof. When two claims in an application are duplicates or else are so close in content that they both cover the same thing, despite a slight difference in

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wording, it is proper after allowing one claim to object to the other as being a substantial duplicate of the allowed claim. See MPEP § 706.03(k).

#### Claim Rejections - 35 USC § 112

6. The following is a quotation of 35 U.S.C. 112(b):

(b) CONCLUSION.—The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the inventor or a joint inventor regards as the invention.

The following is a quotation of 35 U.S.C. 112 (pre-AIA), second paragraph: 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.

- 7. Claims 1- 11, 13 17, 19 and 20 are rejected under 35 U.S.C. 112(b) or 35 U.S.C. 112 (pre-AIA), second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which the inventor or a joint inventor, or for pre-AIA the applicant regards as the invention.
- 8. Claims 1 11, 13 17, 19 and 20 are rendered grammatically indefinite for failing to recite –and—after: "24 months" at line 5 of claim 1.
- 9. Claims 2 11, 13 17, 19 and 20 are rejected for being dependent upon a rejected base claim.

#### Double Patenting

10. The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. A nonstatutory double patenting rejection is appropriate where the conflicting claims are not identical, but at least one examined application claim is not patentably distinct from the reference

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claim(s) because the examined application claim is either anticipated by, or would have been obvious over, the reference claim(s). See, e.g., *In re Berg*, 140 F.3d 1428, 46 USPQ2d 1226 (Fed. Cir. 1998); *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970); *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) or 1.321(d) may be used to overcome an actual or provisional rejection based on nonstatutory double patenting provided the reference application or patent either is shown to be commonly owned with the examined application, or claims an invention made as a result of activities undertaken within the scope of a joint research agreement. See MPEP § 717.02 for applications subject to examination under the first inventor to file provisions of the AIA as explained in MPEP § 2159. See MPEP §§ 706.02(l)(1) - 706.02(l)(3) for applications not subject to examination under the first inventor to file provisions of the AIA. A terminal disclaimer must be signed in compliance with 37 CFR 1.321(b).

The USPTO Internet website contains terminal disclaimer forms which may be used. Please visit www.uspto.gov/patent/patents-forms. The filing date of the application in which the form is filed determines what form (e.g., PTO/SB/25, PTO/SB/26, PTO/AIA/25, or PTO/AIA/26) should be used. A web-based eTerminal Disclaimer may be filled out completely online using web-screens. An eTerminal Disclaimer that meets all requirements is auto-processed and approved immediately upon submission. For

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more information about eTerminal Disclaimers, refer to www.uspto.gov/patents/process/file/efs/quidance/eTD-info-I.jsp.

11. Claims 1, 3 – 11, 13 - 17 and 20 are rejected on the ground of nonstatutory double patenting as being unpatentable over claims 1- 4 of U.S. Patent No. 9,034,388 ('388). Although the claims at issue are not identical, they are not patentably distinct from each other because the only difference between the instant claims and copending claims is a matter of scope of the claimed subject matter.

Claims of the instant invention are drawn to a method of producing krill oil from krill. The krill is Euphausia superba of the Antarctic. The krill oil comprises ether phospholipids, astaxanthin esters and phosphatidylcholine. Encapsulating is also claimed. The steps of the instantly claimed invention include denaturing, by heating, whereas the step of claim 1 of '388 include cooking and dying krill. Both methods recite extraction that includes the use of a polar solvent. Claim 1 of the instantly claimed invention broadly recites extraction and as such may include supercritical fluid extraction. The krill oil produced by the method of '388 can be encapsulated.

The claims of the instant invention differ in terms of scope from patent '388.

It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to produce krill oil from krill as claimed herein comprising obtaining a krill oil, and include denaturing, and use of supercritical fluid extraction to provide krill oil and then to encapsulate for oral consumption. One of skill based on a reading of the claims of '388 would have been motivated to provide for the instant claimed method for production of krill oil because each of the steps as instantly

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claimed are taught by the copending subject matter. Obtaining step can clearly include denaturing and treating as the steps are required by the claims of '388. Clearly encapsulating krill oil suggests formulating krill oil for oral consumption. To obtain krill oil by denaturing the krill to provide a denatured krill product is clearly suggested by the claims of '388. The instant claims are, therefore, considered to be prima facie obvious over the claimed subject matter of '388.

12. Claims 1 - 20 are rejected on the ground of nonstatutory double patenting as being unpatentable over claims 1 - 3, 6 - 9, 33 - 38, and 40 - 42 of U.S. Patent No. 9,375,453 ('453). Although the claims at issue are not identical, they are not patentably distinct from each other because the only difference between the instant claims and copending claims is a matter of scope of the claimed subject matter.

Claims of the instant invention are drawn to a method of producing krill oil from denatured krill. The krill is Euphausia superba of the Antarctic. The krill oil comprises ether phospholipids, astaxanthin esters and phosphatidylcholine. Encapsulating is also claimed. The steps of the instantly claimed invention include denaturing of the krill and extraction with a polar solvent. Also, these steps can be conducted on a ship.

The claims of '453 are drawn to a method of production of krill oil from krill comprising obtaining denatured krill oil and extracting the oil using polar solvent. The krill is Euphausia superba. The krill oil comprises astaxanthin esters, ether phospholipids and phosphatidylcholine. The krill oil can be encapsulated and the method of treating, denaturing, the krill can be conducted on a ship.

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It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to produce krill oil from krill as claimed herein comprising obtaining a krill oil, and include denaturing, and use of a polar solvent during extraction to provide krill oil and then to encapsulate for oral consumption. One of skill based on a reading of the claims of '453 would have been motivated to provide for the instant claimed method for production of krill oil because each of the steps as instantly claimed are taught by the '453 subject matter. Obtaining step can clearly include denaturing and treating as the steps are required by the claims of '453.

Clearly encapsulating krill oil suggests formulating krill oil for oral consumption. To obtain krill oil by denaturing the krill to provide a denatured krill product is clearly suggested by the claims of '453. The instant claims are, therefore, considered to be prima facie obvious over the claimed subject matter of '453.

13. Claims 1- 20 rejected on the ground of nonstatutory double patenting as being unpatentable over claims 1, 2, 3, 6 – 12 and 15 - 19 of U.S. Patent No. 9,028,877 ('877). Although the claims at issue are not identical, they are not patentably distinct from each other because the only difference between the instant claims and the claims of '877 is a matter of scope of the claimed subject matter.

Claims of the instant invention are drawn to a method of producing krill oil from denatured krill. The krill is Euphausia superba of the Antarctic. The krill oil comprises ether phospholipids, astaxanthin esters and phosphatidylcholine. Encapsulating is also claimed. The steps of the instantly claimed invention include denaturing of the krill and extraction with a polar solvent. Also, these steps can be conducted on a ship.

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The claims of '877 teach methods of producing of krill oil which include the steps of extraction with a polar solvent and denaturing the krill; and conducing the method on a ship. Also, the method of '877 encapsulates the krill oil.

The difference between the claimed inventions and '877 is that the instantly claimed invention does not teach the invention with particularity so as to amount to anticipation (See M.P.E.P. § 2131: "[t]he identical invention must be shown in as complete detail as is contained in the ... claim." Richardson v. Suzuki Motor Co., 868 F.2d 1226, 1236, 9 USPQ2d 1913, 1920 (Fed. Cir. 1989). The elements must be arranged as required by the claim, but this is not an ipsissimis verbis test, i.e., identity of terminology is not required. In re Bond, 910 F.2d 831, 15 USPQ2d 1566 (Fed. Cir. 1990)). However, based on the above, the instantly claimed invention is consider to be prima facia obvious over '877.

14. Claims 1, 3 – 8, 10, 11, 12 – 18 and 20 are rejected on the ground of nonstatutory double patenting as being unpatentable over claims 15 - 19 of U.S. Patent No. 9,119,864 ('864). Although the claims at issue are not identical, they are not patentably distinct from each other because the only difference between the instant claims and copending claims is a matter of scope of the claimed subject matter.

Claims of the instant invention are drawn to a method of producing krill oil from denatured krill. The krill is Euphausia superba of the Antarctic. The krill oil comprises ether phospholipids, astaxanthin esters and phosphatidylcholine. Encapsulating is also claimed. The steps of the instantly claimed invention include denaturing of the krill and extraction with a polar solvent.

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The claims of '864 teach a method of producing polar krill oil from Euphausia superba krill. The method includes denaturing the Euphausia superba krill, extracting with supercritical fluid and encapsulating the polar krill oil.

It would have been obvious to one of ordinary skill in the art at the time the claimed invention was made to produce krill oil from krill as claimed herein comprising obtaining a krill oil, and include denaturing, and use of a supercritical fluid during extraction to provide krill oil and then to encapsulate for oral consumption. One of skill based on a reading of the claims of '864 would have been motivated to provide for the instant claimed method for production of krill oil because each of the steps as instantly claimed are taught by the '8643 subject matter. Obtaining step can clearly include denaturing and treating as the steps are required by the claims of '864.

Clearly encapsulating krill oil suggests formulating krill oil for oral consumption. To obtain krill oil by denaturing the krill to provide a denatured krill product is clearly suggested by the claims of '864. The instant claims are, therefore, considered to be prima facie obvious over the claimed subject matter of '864.

#### Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to YATE' K. CUTLIFF whose telephone number is (571)272-9067. The examiner can normally be reached on M-F 8:30 a.m. - 5:30 p.m..

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Fereydoun Sajjadi can be reached on (571) 272-3311. The fax phone

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number for the organization where this application or proceeding is assigned is 571-273-8300.

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/YATE' K. CUTLIFF/ Primary Examiner, Art Unit 1672

#### Application No.: 15/180431

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               40 SEA ABB=ON PLU=ON L6 AND PHOSPHOLIPID
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40 SEA ABB=ON PLU=ON L7 AND ASTAXANTHIN
18 SEA ABB=ON PLU=ON L9 AND (DENATURED (3A) KRILL)
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	Application Number			
INFORMATION BIOOLOGUES	Filing Date		2016-06-13	
INFORMATION DISCLOSURE	First Named Inventor Inge Bru		ruheim	
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit			
(Not for Submission under or or N 1.00)	Examiner Name			
	Attorney Docket Number		AKBM-14409/US-12/CON	

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First Named Inventor Inge E		Bruheim
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#### INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

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Examiner Name		
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	RMATION DISCLOSURE First Named Inventor Inge Bruheim Art Unit 1672 Examiner Name CUTLIFF, YATE KAI RENE	Filing Date		2016-06-13	
	First Named Inventor Inge E		ge Bruheim		
	Art Unit		1672		
INFORMATION DISCLOSURE STATEMENT BY APPLICANT ( Not for submission under 37 CFR 1.99)	Examiner Name CUTL		LIFF, YATE KAI RENE		
	Attorney Docket Number	er	AKBM-14409/US-12/CON		

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2 20100226977 2010-09-09 TILSETH SNORRE et al.  3 20140274968 2014-09-18 BERGE KJETIL et al.	
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1 2014/013335 WO 2014-01-23 HOEM, Nils et al.	

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	3	2909508	JP	1999-06-23	TAIYO FISHERY CO LTD.		$\boxtimes$		
	4	2010/097701	wo	2010-09-02	AKER BIOMARINE ASA				
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	1	International Search Report, International Patent Application No. PCT/IB2016/000208, mailed May 13, 2016, five pages							
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	1	9119864		2015-09-01	AKER BIOMARINE ANTARCTIC AS		
	2	9072752		2015-07-07	AKER BIOMARINE ANTARCTIC AS		
	3	9034388		2015-05-19	Inge Bruheim et al.		
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	6	8372812		2013-02-12	Snorre Tilseth et al.		
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	1	20050003073		2005-01	-06	PIVOVAROV et al.					
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	1	Declaration of Bjorn Ole Haugsgjerd submitted during inter partes reexamination of parent pater ("Haugsgjerd '348 Decl.")	Declaration of Bjorn Ole Haugsgjerd submitted during inter partes reexamination of parent patent U.S. 8,030,348 "Haugsgjerd '348 Decl.")					
	2	Declaration of Dr. Albert Lee in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Lee")						
	3	Declaration of Dr. Albert Lee in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Lee'	")					
	4	Declaration of Dr. Chong Lee submitted during inter partes reexamination of parent patent U.S.: Reexam Decl.")	8,030,348	("Yeboah				
	5	Declaration of Dr. Earl White submitted during prosecution of parent patent U.S. 8,030,348 ("20"	11 White D	ecl.")				
	6	Declaration of Dr. Ivar Storrø in support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Stor	ггø")					
	7	Declaration of Dr. Ivar Storrø in support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Sto	ггø")					
	8	Declaration of Dr. Jacek Jaczynski from inter partes reexamination of the parent patent U.S. 8,0: Reexam. Decl.")	30,348 ("Ja	aczynski				
	9	Declaration of Dr. Jaczynski submitted during prosecution of parent patent U.S. 8,278,351 (Jacz	ynski '351	Decl.")				

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10	0	Declaration of Dr. Jeff Moore in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Moore")
1	1	Declaration of Dr. Jeff Moore in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Moore")
12	2	Declaration of Dr. Richard van Breemen in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Van Breemen")
13	3	Declaration of Dr. Richard van Breemen in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Van Breemen")
14		Declaration of Dr. Shahidi submitted during inter partes reexamination of parent patent U.S. 8,030,348 (Shahidi Reexam. Decl.")
15	5	Declaration of Dr. Shahidi submitted during prosecution of parent patent U.S. 8,278,351 (Shahidi '351 Decl.")
16	6	Declaration of Dr. Suzanne Budge in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Budge")
17	7	Declaration of Dr. Suzanne Budge in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Budge")
18	8	Declaration of Dr. Thomas Brenna in support of Inter Partes Review of U.S. Pat. No. 8,278,351
15	9	Declaration of Dr. Thomas Brenna in support of Inter Partes Review of U.S. Pat. No. 8,383,675
20	20	Declaration of Dr. Thomas Gundersen submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("Gundersen Decl.")
15	9	Declaration of Dr. Thomas Brenna in support of Inter Partes Review of U.S. Pat. No. 8,383,675  Declaration of Dr. Thomas Gundersen submitted during inter partes reexamination of parent patent U.S. 8,030,348

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21	Declaration of Dr. Tina Sampalis submitted during inter partes reexamination of parent patent U.S. 8,030,348 (Sampalis")	
22	Declaration of Dr. Van Breemen submitted during Ex parte Reexamination of the '351 patent (Van Breemen '351 Reexam. Decl."	
23	Declaration of Dr. Van Breemen submitted during Inter partes Reexamination of the '348 patent (Van Breemen '348 Reexam Decl."	
24	Declaration of Dr. Yeboah submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("Yeboah Reexam Decl.")	
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	43	Provisional Application No. 60/307,842 (Priority document for the '351 patent), available in PAIR						
	44	Supplemental Declaration of Bjorn Ole Haugsgjerd submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("Haugsgjerd '348 Supp. Decl.")						
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S2	27	(("TILSETH") near3 ("Snorre")).I <b>NV</b> .	US- PGPUB; USPAT; USOCR	OR	OFF	2016/07/12 16:47
S3	20	(("MANCINELLI") near3 ("Daniele")).INV.	US- PGPUB; USPAT; USOCR	OR	OFF	2016/07/12 16:47
S4	7	("20030044495"   "20040241249"   "20080166419"   "4119619"   "5266564"   "5434183"   "6537787"   "6800299"   "7666447"   "8030348").PN.	USPAT	OR	OFF	2016/07/13 13:44
S5	19	("20030044495"   "20040241249"   "20080166419"   "4119619"   "5266564"   "5434183"   "6537787"   "6800299"   "7666447"   "8030348").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:44
S6	5	("4714571"   "8278351"   "8383675").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:44
S7	28	("2652235"   "4036993"   "4251557"   "4505936"   "5006281"   "6214396"   "6346276").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:45
S8	21	("20020076468"   "20030113432"   "20060078625"   "20080166419"   "20080166420"   "20100143571"   "20100160659"   "20110130458"   "4133077"   "4749522"   "4814111"   "7488503"   "8697138").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:46
S9	1	("20060078625"   "8057825").PN.	USPAT	OR	OFF	2016/07/13 13:46
S10	3	("20060078625"   "8057825").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:46
S11	10	("20050003073"   "20110160161"   "20110256216"   "4038722"   "8586567").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 13:47
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S12	0	("5266564," "8030348," "7666447," "4714571," "8278351," "8383675," "2652235," " 5006281").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:11
S13	0	("5266564," "8030348," "7666447," "4714571," "8278351," "8383675," "2652235," "5006281").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:11
S14	24	("5266564," "8030348," "7666447," "4714571," "8278351," "8383675," "2652235," "5006281").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:12
S15	22	("4251557," "4505936," "6214396," "4036993," "6346276," "8697138," "7488503," "4749522," "4133077").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:14
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S17	9	("9119864," "9072752," "9034388," "9028877," "9078905," "8372812," "8697138").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:18
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S19	22	("20140274968," "20150030718," "20140088043," "20140088047," "20140080791," "20150164841," "20140363517," "20100226977," "20150050403," "20140005421," "20140107072," "20090061067," "20140010888").pn.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:21
S20	10	(S5 or S6 or S7 or S8 or S10) and krill	USPAT	OR	OFF	2016/07/13 14:22
S21	15	(S11 or S14 or S15 or S16 or S17 or S19) and krill	USPAT	OR	OFF	2016/07/13 14:23
S22	16	(S20 or S21) and (extraction or extract or extracting) and oil	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:24
S23	20	(S5 or S6 or S7 or S8 or S10) and krill	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:24
S24	40	(S11 or S14 or S15 or S16 or S17 or S19) and krill	US- PGPUB;	OR OST EX	OFF	2016/07/13 14:25 <b>1111 P</b> a

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S25	42	(S23 or S24) and (extraction or extract or extracting) and oil	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:25
S26	10	S25 and (polar near2 solvent)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/13 14:27
S27	3	("20100226977"   "20140274968"   "20140370115").PN.	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:47
S28	0	S27 and krill	USPAT	OR	OFF	2016/07/14 09:47
S29	0	S27 and (extraction or extracting or extract) and pholpholipid?	USPAT	OR	OFF	2016/07/14 09:48
S30	0	S27 and (extraction or extracting or extract)	USPAT	OR	OFF	2016/07/14 09:48
S31	3	S27 and (extraction or extracting or extract)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:49
S32	3	S27 and (extraction or extracting or extract) and ((polar near2 solvent) or ethanol)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:50
S33	3	S27 and krill	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 09:53
S34	696	krill and oil and astaxanthin	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 10:55
S35	201	(krill and oil and astaxanthin) and phosphatidylcholine	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 10:56
S36	13	denature near3 lipase	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 11:17
S37	21	denature near3 lipases	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 11:17

31	`F	N	30	7	:	31
S38		S34 and S37	USPAT	OR	OFF	2016/07/14 11:17
S39	52	(krill adj oil) and (extraction or extracting or extract) and (ship or boat)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 13:36
S40	0	S39 and @ay<="2007"	USPAT	OR	OFF	2016/07/14 13:36
S41	1	S39 and @ay<="2008"	USPAT	OR	OFF	2016/07/14 13:36
S42	50	((krill adj oil) same (extraction or extracting or extract)) and (ship or boat)	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 13:37
S43	179	c11b3/006.cpc.	USPAT	OR	OFF	2016/07/14 15:55
S44	0	c11b1/02,16,cpc.	USPAT	OR	OFF	2016/07/14 15:56
S45	133	c11b1/02,16.cpc.	USPAT	OR	OFF	2016/07/14 15:57
S46	664	c07c51/48.cpc.	USPAT	OR	OFF	2016/07/14 15:57
S47	1226	a61k31/122,685.cpc.	USPAT	OR	OFF	2016/07/14 15:57
S48	236	a61k9/48.cpc.	USPAT	OR	OFF	2016/07/14 15:58
S49	201	(krill and oil and astaxanthin) and phosphatidylcholine	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 16:01
S50	9	S43 and S49	US- PGPUB; USPAT; USOCR; FPRS	OR	OFF	2016/07/14 16:01
S51	6	S45 and S49	USPAT	OR	OFF	2016/07/14 16:03
S52	7	S45 and (krill near2 oil)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:04
S53	1	S46 and (krill near2 oil)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:08
S54	55	S47 and (krill near2 oil)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:09
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S55	7	S47 and (krill near2 oil) and (denature near3 lipases)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:09
S56	7	S48 and (krill near2 oil) and (denature near3 lipases)	US- PGPUB; USPAT; USOCR; FPRS; EPO	OR	OFF	2016/07/14 16:10

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#### Search Notes



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CPC- SEARCHED		
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C11B 1/02, 16: see search history	7/14/2016	ykc
C11b 3/006: see search history	7/14/2016	ykc
C07C 51/48: see search history	7/14/2016	ykc
A61K 31/122, 685: see search history	7/14/2016	ykc

CPC COMBINATION SETS - SEAR	CHED	
Symbol	Date	Examiner

	US CLASSIFICATION SE	ARCHED	
Class	Subclass	Date	Examiner

SEARCH NOTES		
Search Notes	Date	Examiner
Palm Inventor Search	7/14/2016	ykc
East	7/14/2016	ykc
STN: caplus, agricola USpat2, US patfull, IFlall	7/14/2016	ykc

	INTERFERENCE SEARCH		
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INFORMATION PIGGL COURT	Filing Date		2016-06-13	
INFORMATION DISCLOSURE	First Named Inventor Inge Br		Bruheim	
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit			
(Not for Submission under or or N 1.00)	Examiner Name			
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	1	4119619		1978-10-10	0 ROGOZHIN SERGEI VASILIEVICH et al.			
	2	5434183		1997-07-18	LARSSON-BACKSTROM			
	3 6537787			2003-03-25	BRETON			
	4 6800299			2004-10-05	BEAUDOIN & MARTIN			
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	2	1098900	CA		1981-04-07	ROGOZHIN, et al		
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	5	1406641	EΡ		2004-04-14	NEPTUNE TECHNOLOGIES & BIORESSOURCES INC.		
	6	0670306	EP		1995-06-09	NIPPON OIL CO. LTD		
	7	2097014	GB		1982-10-27	BAIKOFF		
	8	921537	GB		1999-06-09	PICKER NORDSTAR INC.		
	9	02049091	JP		1990-02-19	SUNTORY LTD		
	10	2215351	JP		1990-08-28	TAIYO FISHERY CO LTD		

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12	2963152	JP	1992-02-25	CHLORINE ENG CORP LTD
13	3081692	JP	1994-07-19	CHLORINE ENG CORP LTD
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15	3467794	JP	2003-09-05	NIPPON OIL & FATS CO LTD
16	3486778	JP	2003-10-31	GREEN CROSS CORP
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18	3678317	JP	2005-05-20	CHLORINE ENG CORP LTD
19	4012665	JP	1992-01-17	MATSUSHITA ELECTRIC IND CO LTD
20	61281159	JP	1986-12-11	SHISEIDO CO LTD; NIPPON SUISAN KAISHA LTD.
21	2001-158736	JP	2001-06-12	SNOW BRAND MILK PROD CO LTD

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23	2003-048831	JP	2003-02-21	SUNTORY LTD
24	2003-146883	JP	2003-05-21	SNOW BRAND MILK PROD CO LTD
25	2005-245379	JP	2005-09-15	NIPPON SUISAN KAISHA LTD
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29	2006-316073	JP	2006-11-24	IBR ISRAELI BIOTECHNOLOGY RESEARCH LTD
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31	2007-126455	JP	2007-05-24	FUJI CHEM IND CO LTD
32	2007-246404	JP	2007-09-27	SNOW BRAND MILK PROD CO LTD

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34	1986/06082	wo	1986-10-23	MAT-CON RADGIVENDE INGENIØRFIRMA A/S
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38	1997/39759	WO	1997-10-30	BRIGHAM AND WOMEN'S HOSPITAL
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41	2000/23546	WO	2000-04-27	UNIV SHERBROOKE
42	2 2000/25608	WO	2000-05-11	NIPPON SUISAN KAISHA, LTD
43	2000/38708	wo	2000-07-06	PHAIRSON MEDICAL INC.

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	44	2002/102394	wo		2002-12-27	NEPTUNE TECHNOLOGIES & BIORESS		
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	48	2007/080514	wo		2007-07-19	KRILL A/S		
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	50	2007/108702	wo		2007-09-27	AKER SEAFOODS HOLDING AS		
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# INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

Application Number		15180431 - GAU: 1672					
Filing Date		2016-06-13					
First Named Inventor	Inge E	Bruheim					
Art Unit							
Examiner Name							
Attorney Docket Number		AKBM-14409/US-12/CON					

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Examiner	Signat	ture /YATE' K CUTLIFF/	Date Considered	07/13/2016							
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Application Number			12100421 -	GAU:	10/2
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First Named Inventor Inge B		Bruheim			
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Attorney Docket Numb	er	AKBM-14409/US-12/CO	N		

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Signature	/J. Mitchell Jones/	Date (YYYY-MM-DD)	2016-06-13
Name/Print	J. Mitchell Jones	Registration Number	44174

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INVENTORS Inge Bruheim, Volda, NORWAY; Snorre Tilseth, Bergen, NORWAY; Daniele Mancinelli, Orsta, NORWAY;												
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	Application Number		15180431		
	Filing Date		2016-06-13		
INFORMATION DISCLOSURE	First Named Inventor Inge Br		Bruheim		
STATEMENT BY APPLICANT ( Not for submission under 37 CFR 1.99)	Art Unit		1672		
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	Attorney Docket Number	er	AKBM-14409/US-12/CON		

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Application Number		15180431	
Filing Date		2016-06-13	
First Named Inventor	Inge Bruheim		
Art Unit		1672	
Examiner Name	CUTLIFF, YATE KAI RENE		
Attorney Docket Number		AKBM-14409/US-12/CON	

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(72) Inventors; and

(71) Applicants (for US only): HOEM, Nils [NO/NO]; Skovveien 6a, N-0257 Oslo (NO). TILSETH, Suorre Published: [NO/NO]; Fantoltasen 27A, N-5072 Bergen (NO).

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(54) Title: CONCENTRATION OF OMEGA-3 POLYUNSATURATED FATTY ACIDS IN KRILL OIL

(57) Abstract: The present invention relates to krill oil, and in particular to krill oil with elevated levels of omega-3 fatty acids and decreased levels of saturated fatty acids.

#### Concentration of Omega-3 Polyunsaturated Fatty Acids in Krill Oil

#### Field of the Invention

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The present invention relates to krill oil, and in particular to krill oil with elevated levels of omega-3 fatty acids and decreased levels of saturated fatty acids.

#### **Background of the Invention**

Fish oils that are abundant in omega-3 polyunsaturated fatty acids (PUFA) have traditionally been used as the raw material for preparation of omega-3 PUFA concentrate. Since fish oils are complex mixtures of triglycerides containing fatty acids with varying chain lengths and degrees of unsaturation, separation of individual fatty acids is difficult for production of concentrated omega-3 components. Therefore, commercial production of marine oil concentrates with enhanced percentages of EPA and DHA has been a major challenge for food scientists and biotechnologists engaged in research in this area.

Methods for concentration of omega-3 PUFA are numerous, but only few are suitable for large-scale production. Distillation has been used for partial separation of mixtures of fatty acid esters. This method takes advantage of differences in the boiling point and molecular weight of fatty acids under reduced pressure. This technique requires high temperatures of approximately 250 C. Berger, R. and McPherson, W. (1979) 'Fractional Distillation' in J. Am. Oil Chem. Soc. 56, 743A-746A. Short-path distillation or molecular distillation uses lower temperatures and short heating intervals. However, fractionation of fish oil esters is difficult since separation of these components becomes less effective with increasing molecular weight. Weitkamp, A.W. (1955) 'Distillation' in J. Am. Oil Chem. Soc. 32, 640-646; Brevik, H. (1992) 'N-3 Concentrates: A Scandinavian View-point' in AOCS Short Courses, Modern Application of Marine Oils, 7-8 May, Toronto, ON, Canada.

The most widely used distillation procedure is fractional distillation of methyl esters under reduced pressure (0.1-1.0 mmHg). Even under these conditions, moderately high temperatures are required; the more highly unsaturated acids, especially omega-3 PUFA are more prone to oxidation, polymerization and isomerization of double bonds. Distillation at still lower pressures has been used in the isolation of some highly unsaturated acids, and is particularly valuable in polymerization studies to separate monomeric, dimeric and

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polymeric materials and in the separation of monoacylglycerols from di- and triacylglycerol mixtures.

Another method for making fish oil concentrates is via enzymatic processing, such a lipase catalyzed hydrolysis. The presence of cis carbon-carbon double-bonds in the fatty acids results in bending of the chains. Therefore, the terminal methyl group of the fatty acid lies close to the ester bond which may cause a steric hinderance effect on lipases. The high bending effect of EPA and DHA due to the presence of the 5 and 6 double-bonds, respectively, enhances the steric hinderance effect; therefore, lipases cannot reach the ester-linkage between these fatty acids and glycerol. However, saturated and monounsaturated fatty acids do not present any barriers to lipases and they could be easily hydrolyzed. Therefore, fatty acid selectivity of a lipase for EPA and DHA allows separation and concentration of these fatty acids from others in the remaining portion of marine oils. In addition, lipases have been frequently used to discriminate between EPA and DHA in concentrates containing both of these fatty acids. See, e.g., Bottino, N.R., Vandenberg, G.A. and Reiser, R. (1967) 'Resistance of Certain Long-chain Polyunsaturated Fatty Acids of Marine Oils to Pancreatic Lipase Hydrolysis' in Lipids 2, 489-493. In most commercial processes, the hydrolysis is performed on esters produced from fish oils. The end product is accordingly an ester concentrate.

Other methods used in the art for processing fish oils include low temperature crystallization and treatment with solvents (see, e.g., Brown, L.B. and Kolb, D.X. (1955) `Application of Low Temperature Crystallization in the Separation of the Fatty Acids and their Compounds' in Prog. Chem. Fats Lipids 3, 57-94; WO91/13957) and supercritical fluid extraction (see, e.g, Mishra, V.K., Temelli, F. and Ooraikul, B. (1993) `Extraction and Purification of Omega 3-Fatty Acids with an Emphasis on Supercritical Fluid Extraction, a Review' in Food Res. Inter. 26, 217-226).

The concentrated esters produced by these processes can be encapsulated and sold, or the esters can be used to make triglycerides. The TAG form of PUFA is considered to be nutritionally more favorable than methyl or ethyl esters of fatty acids because experimental results have shown impaired intestinal absorption of methyl or ethyl esters of omega-3 PUFA in laboratory animals. Hamazaki, T., Hirai, A., Terano, T., Sajiki, J., Kondo, S., Fujita, T., Tamura, Y. and Kumagai, A. (1982) `E ect of Orally Administrated Ethyl Ester of Eicosapentaenoic Acid on PGI-like Substance Production by Rat Aorta' in Prostaglandins 23,

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557-567; El-Boustani, S., Colette, C, Monnier, L., Descomps, B., Paulet, C.A. and Mendey, F. (1987) `Eternal Absorption in Man of Eicosapentaenoic Acid in Different Chemical Forms' in Lipids 22, 711-714; Lawson, L.D. and Hughes, B.G. (1988) `Human Absorption of Fish Oil Fatty Acids as Triacylglycerols, Free Fatty Acids or Ethyl Esters' in Biochem. Biophys. Res.

Comm. 152, 328±335. Yang et al. have shown that methyl and ethyl esters are hydrolysed slower than their corresponding TAG. Yang, L.Y., Kuksis, A. and Myher, J.J. (1989) `Lumenal Hydrolysis of Menhaden and Rapeseed Oils and their Fatty Acid Methyl and Ethyl Esters in the Rats' in Biochem. Cell Biol. 67, 192-204. From a marketing point of view, triacylglycerols of PUFA are often promoted as being more `natural' than other fatty

acid derivatives. For these reasons, the esters produced by the processes described above are converted into fatty acids and then incorporated into glycerol by direct esterification or incorporated into glycerides by transesterification. See, e.g., Osada, K., Nakamura, M., Nonaka, M. and Hatano, M. (1992) Esterification of Glycerol with EPA and DHA by Chromobacterium viscosum and Candida cylindracea Lipases' in J. Jpn. Oil Chem. Soc. 41, 39-

43; He, Y. and Shahidi, F. (1997) `Enzymatic Esterification of Omega-3-fatty Acid
Concentrates from Seal Blubber Oil with Glycerol' in J. Am. Oil Chem. Soc. 74, 1133-1136;
Akoh, C.C., Jennings, B.H. and Lillard, D.A. (1996) `Enzymatic
Modification of Evening Primrose Oil: Incorporation of n-3 Polyunsaturated Fatty Acids' in J.

Am. Oil Chem. Soc. 73, 1059-1062; Akoh, C.C., Jennings, B.H. and Lillard, D.A. (1995)

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Modification of Triolein: Incorporation of n-3 Polyunsaturated Fatty Acids' in J. Am. Oil Chem. Soc. 72, 1317-1321; Sridar, R. and Laksminarayana, G. (1992) 'Incorporation of Eicosapentaenoic and Docosahexaenoic Acids into Ground Nut Oil by Lipase-catalyzed Ester Exchange' in J. Am. Oil Chem. Soc. 69, 1041-1042. Basheer, S., Mogi, K. and Nakajuma, M. (1995) 'Interesterification Kinetics of Triacylglycerides and Fatty Acids with Modified Lipase in n-hexane' in J. Am. Oil Chem. Soc. 72, 511-518.

Krill oil differs from fish oil in that krill oil comprises high amounts of phospholipids. See e.g., WO 2008/117062; US PUBL. NO. 20080274203. One of the main advantages of krill oil as compared to fish oil is increased bioavailability of omega-3 PUFA in the form of a phospholipid. However, the enzymatic processes described above are not amenable for use with phospholipids. In particular, conversion of esterified or non-esterified omega-3 PUFA

back to the phospholipid form is not trivial. Thus, the methods that have been developed for production of fish oil concentrates are not easily transferred to krill oil processing.

Oils with increased amounts of EPA and DHA are desirable because a lower dose is needed to provide the same amount of DHA and EPA. Krill oil concentrates containing increased amounts of omega-3 PUFA in the phospholipid form as compared to other fatty acids in the krill oil compositions have not been developed due to the problems described above. This has been a disadvantage in the market because the fish oil concentrates contain higher amounts of omega-3 PUFA, in particular EPA and DHA, than commercial available krill oil.

Accordingly, what is needed in the art are krill oil concentrates comprising higher amounts of omega-3 PUFA as compared to commercially available krill oils.

#### Summary of the Invention

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The present invention relates to krill oil, and in particular to krill oil with elevated levels of omega-3 fatty acids and decreased levels of saturated fatty acids.

In some embodiments, the present invention provides a krill oil, such as a krill oil concentrate, comprising greater than about 22% EPA (w/w total fatty acids), greater than about 10% DHA (w/w total fatty acids), from 4% to 8% myristic acid (w/w total fatty acids), from 3% to 9% c9 oleic acid (w/w total fatty acids), and 20 to 4000 ppm astaxanthin. In some embodiments, the krill oil further comprises about 22% to 30% EPA. In some embodiments, the krill oil further comprises about 10% to 15% DHA. In some embodiments, the krill oil is extracted from Euphausia superba. In some embodiments, the ratio of DHA and EPA: omega 6 (w/w total fatty acids) is from about 10:1 to 14:1. In some embodiments, the ratio of DHA and EPA: c9 oleic acid (w/w total fatty acids) is from about 4:1 to 8:1. In some embodiments, the ratio of DHA and EPA: myristic acid (w/w total fatty acids) is from about 4:1 to 8:1. In some embodiments, the ratio of DHA and EPA: myristic acid and c9 oleic acid (w/w total fatty acids) is from about 2:1 to 4:1. In some embodiments, the ratio omega 3: omega 6 (w/w total fatty acids) is from about 11:1 to 15:1. In some embodiments, the ratio of omega 3: c9 oleic acid (w/w total fatty acids) is from about 5:1 to 9:1. In some embodiments, the ratio of omega 3: myristic acid (w/w total fatty acids) is from about 5:1 to 9:1. In some embodiments, the ratio of omega 3: myristic acid and c9 oleic acid (w/w total fatty acids) is from about 2.5:1 to 4.5:1.

In some embodiments, the present invention provides a capsule containing a krill oil as described above. In some embodiments, the present invention provides a food product containing a krill oil as described above. In some embodiments, the present invention provides a dietary supplement containing a krill oil as described above. In some embodiments, the present invention provides a oil in water emulsion containing a krill oil as described above.

In some embodiments, the krill oils are used for oil administration to a subject. In some embodiments, the krill oil are used for treatment of a condition for which omega-3 is effective.

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#### Definitions

As used herein, "krill oil" refers to an oil extracted from *Euphausia sp.*, for example, *Euphausia superba*.

As used herein, "phospholipid" refers to an organic compound having the following general structure:

wherein R1 is a fatty acid residue, R2 is a fatty acid residue or –OH, and R3 is a –H or nitrogen containing compound choline (HOCH<sub>2</sub>CH<sub>2</sub>N<sup>+</sup>(CH<sub>3</sub>)<sub>3</sub>OH<sup>-</sup>), ethanolamine (HOCH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub>), inositol or serine. R1 and R2 cannot simultaneously be OH. When R3 is an –OH, the compound is a diacylglycerophosphate, while when R3 is a nitrogen-containing compound, the compound is a phosphatide such as lecithin, cephalin, phosphatidyl serine or plasmalogen.

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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, 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).

As used herein, the term omega-6 fatty acid refers to polyunsaturated fatty acids that have the final double bond in the hydrocarbon chain between the sixth and seventh carbon atoms from the methyl end of the molecule.

As ued herein, astaxanthin refers to the following chemical structure:

As used herein, astaxanthin esters refer to the fatty acids esterified to OH group in the astaxanthin molecule.

As used herein, the term w/w (weight/weight) refers to the amount of a given substance in a composition on weight basis and can be expressed as a percentage. For example, a composition comprising 50% w/w phospholipids means that the mass of the phospholipids is 50% of the total mass of the composition (i.e., 50 grams of phospholipids in 100 grams of the composition, such as an oil). The w/w may also be used to refer to the amount, on a weight basis, of one member of a class of molecules in a composition as compared to all members of the class of molecules. For example, the amount of a particular fatty acid (or class of fatty acids such as omega 3 fatty acids) may be expressed as a percentage of all other fatty acids in the composition on a weight/weight basis, i.e., the weight of the specific fatty acids as a percentage of the total weight of fatty acids in the composition.

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#### **Description of the Invention**

The present invention relates to krill oil, and in particular to krill oil with elevated levels of omega-3 fatty acids and decreased levels of saturated fatty acids. While it was previously known that krill oil contains both phospholipid and triglyceride fractions, the inventors have discovered that krill oil is a multiphase dispersion of these fractions. This property of krill oil has not been previously described. The present inventors have taken advantage of this novel observation to develop processes for separation of the phases based on solubility of phospholipids in a polar solvent. By these processes, it is possible to separate, at least partially, the triglyceride and phospholipid phases. Unexpectedly, analysis of the phospholipid phase has revealed that the phospholipid phase has a higher content of omega-3 PUFA as compared to the triglyceride phase or to commercially available krill oil. The novel krill oil compositions and processes are described below.

#### 1. Krill oil compositions

In some embodiments, the present invention provides novel krill oil compositions. The novel krill oil compositions are preferably defined by the amount or ratio of total omega-3 PUFA, the amount of EPA and DHA alone or combined, the amount of c9 oleic acid, and/or the amount of myristic acid as compared to previously described krill oils. In some embodiments, the krill oil is produced in whole or in part from *Euphausia superba*.

In some embodiments, the krill oil comprises greater than about 40% total omega-3 PUFA w/w total fatty acids. By this it is meant that the total amount of omega-3 fatty acids in the krill oil is greater than about 40% of the total fatty acid content of the krill oil on a weight basis. In some of the embodiments described herein, the amounts of particular fatty acids in the krill oil are preferably determined by gas chromatography of a fatty acid methyl esters prepared from the krill oil. In some embodiments, the krill oil comprises greater than about 41%, 42% or 43% total omega-3 PUFA w/w total fatty acids. In some embodiments, the krill oil comprises greater than about 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42% or 43% total omega-3 PUFA w/w total fatty acids up to about 45% total omega-3 PUFA w/w total fatty acids up to about 45% total omega-3 PUFA w/w total fatty acids up to about 46% total omega-3 PUFA w/w total fatty acids up to about 46% total omega-3 PUFA w/w total fatty acids up to about 46% total omega-3 PUFA w/w total fatty acids. In some embodiments, the krill oil

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comprises greater than about 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42% or 43% total omega-3 PUFA w/w total fatty acids up to about 47% total omega-3 PUFA w/w total fatty acids. In some embodiments, the krill oil comprises greater than about 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42% or 43% total omega-3 PUFA w/w total fatty acids up to about 48% total omega-3 PUFA w/w total fatty acids. In some embodiments, the krill oil comprises greater than about 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42% or 43% total omega-3 PUFA w/w total fatty acids up to about 49% total omega-3 PUFA w/w total fatty acids. In some embodiments, the krill oil comprises greater than about 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42% or 43% total omega-3 PUFA w/w total fatty acids up to about 50% total omega-3 PUFA w/w total fatty acids. In some embodiments, the krill oil comprises greater than about 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42% or 43% total omega-3 PUFA w/w total fatty acids up to about 55% total omega-3 PUFA w/w total fatty acids. In some embodiments, the rill oil comprises astaxanthin. In some embodiments, the krill comprises from about 10, 20, 30, 40, 50, 60, 70, 80 or 100 ppm astaxanthin up to about 200, 400, 600, 800, 1000, 1500, 2000, or 4000 ppm astaxanthin. In some embodiments, the krill oil comprises from about 4% to about 8% myristic acid (w/w total fatty acids), from 5% to 7% myristic acid (w/w total fatty acids), or from about 6.0% to 6.6% myristic acid. In some embodiments, the krill oil comprises from about 3% to about 9% c9 oleic acid (w/w total fatty acids), from about 4% to about 8% c9 oleic acid (w/w total fatty acids), from about 5% to about 7% c9 oleic acid (w/w total fatty acids), or from about 6.0% to about 6.6% c9 oleic acid (w/w total fatty acids). In some embodiments, the krill oil comprises less than about 5.0%, 4.5%, 4.0%, 3.8% or 3.6% palmitoleic acid (C16:1; w/w total fatty acids). In some embodiment, the krill oil comprises from about 1% to about 5.0%, about 1.5% to about 4.5%, about 2% to about 4.0%, or about 2.5% to about 3.8% palmitoleic acid (C16:1; w/w total fatty acids).

In some embodiments, the krill oil of the present invention comprises greater than about 22% EPA (w/w total fatty acids), greater than about 10% DHA (w/w total fatty acids), or greater than about 32% EPA and DHA (w/w total fatty acids). In some embodiments, the krill oil of the present invention comprises greater than about 23% EPA (w/w total fatty acids), greater than about 11% DHA (w/w total fatty acids), or greater than about 34% EPA and DHA (w/w total fatty acids). In some embodiments, the krill oil of the present invention comprises greater than about 24% EPA (w/w total fatty acids), greater than about 12% DHA (w/w total fatty acids), or greater than about 36% EPA and DHA (w/w total fatty acids). In

some embodiments, the krill oil of the present invention comprises greater than about 25% EPA (w/w total fatty acids), greater than about 12.3% DHA (w/w total fatty acids), or greater than about 37.3% EPA and DHA (w/w total fatty acids). In some embodiments, the krill oil of the present invention comprises greater than about 25.5% EPA (w/w total fatty acids), greater than about 12.5% DHA (w/w total fatty acids), or greater than about 38% EPA and DHA (w/w total fatty acids). In some embodiments, the krill oil comprises an upper limit of 27% EPA and 13% DHA (w/w total fatty acids; total of 40% EPA and DHA), 28% EPA and 14% DHA (w/w total fatty acids; total of 42% EPA and DHA), 30% EPA and 16% DHA (w/w total fatty acids; total of 46% EPA and DHA), 32% EPA and 18% DHA (w/w total fatty acids; total of 50% EPA and DHA), or 37% EPA and 23% DHA (w/w total fatty acids; total of 60% EPA and DHA). In some embodiments, the rill oil comprises astaxanthin. In some embodiments, the krill comprises from about 10, 20, 30, 40, 50, 60, 70, 80 or 100 ppm astaxanthin up to about 200, 400, 600, 800, 1000, 1500 or 2000 ppm astaxanthin. In some embodiments, the krill oil comprises from about 4% to about 8% myristic acid (w/w total fatty acids), from 5% to 7% myristic acid (w/w total fatty acids), or from about 6.0% to 6.6% myristic acid. In some embodiments, the krill oil comprises from about 3% to about 9% c9 oleic acid (w/w total fatty acids), from about 4% to about 8% c9 oleic acid (w/w total fatty acids), from about 5% to about 7% c9 oleic acid (w/w total fatty acids), or from about 6.0% to about 6.6% c9 oleic acid (w/w total fatty acids).

In some embodiments, the krill oil has a ratio of DHA and EPA: omega 6 PUFAs (w/w total fatty acids) of from about 10:1 to 14:1, 11:1 to 13:1, 11.3:1 to 12.1:1, or 11.5:1 to 11.9:1. In some embodiments, the krill oil has a ratio of DHA and EPA: c9 oleic acid (w/w total fatty acids) of from about 4:1 to 8:1, 5:1 to 7:1, 5.7:1 to 6.9:1, or 6.0:1 to 6.6:1. In some embodiments, the krill oil has a ratio of DHA and EPA: myristic acid (w/w total fatty acids) of from about 4:1 to 8:1, 5:1 to 7:1, 5.7:1 to 6.9:1, or 6.0:1 to 6.6:1. In some embodiments, the krill oil has a ratio of DHA and EPA: myristic acid and c9 oleic acid (w/w total fatty acids) of from about 2:1 to 4:1, 2.4:1 to 3.5:1, 2.7:1 to 3.5:1, or 2.9:1 to 3.3:1.

In some embodiments, the krill oil has a ratio omega-3 PUFAs: omega 6 PUFAs (w/w total fatty acids) of from about 11:1 to 15:1, 12:1 to 14:1, 12.5:1 to 13.5:1, or 12.8:1 to 13.2:1. In some embodiments, the krill oil has a ratio omega-3 PUFAs: c9 oleic acid (w/w total fatty acids) of from about 5:1 to 9:1, 6:1 to 8:1, 6.2:1 to 7.4:1, or 6.4:1 to 7.2:1. In some embodiments, the krill oil has a ratio omega-3 PUFAs: myristic acid (w/w total fatty

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acids) of from about 5:1 to 9:1, 6:1 to 8:1, 6.2:1 to 7.4:1, or 6.4:1 to 7.2:1. In some embodiments, the krill oil has a ratio omega-3 PUFAs: myristic acid and c9 oleic acid (w/w total fatty acids) of from about 2:1 to 5:1, 2.5:1 to 4.5:1, 3:1 to 3.9:1, or 3.2:1 to 3.6:1.

In some embodiments, the krill oil of this invention is formulated with acceptable excipients and/or carriers for oral consumption. The actual form of the carrier, and thus, the composition itself, is not critical. The carrier may be a liquid, gel, gelcap, capsule, powder, solid tablet (coated or non-coated), tea, or the like. The composition is preferably in the form of a tablet or capsule and most preferably in the form of a soft gel capsule. Suitable excipient and/or carriers include maltodextrin, calcium carbonate, dicalcium phosphate, tricalcium phosphate, microcrystalline cellulose, dextrose, rice flour, magnesium stearate, stearic acid, croscarmellose sodium, sodium starch glycolate, crospovidone, sucrose, vegetable gums, lactose, methylcellulose, povidone, carboxymethylcellulose, corn starch, and the like (including mixtures thereof). Preferred carriers include calcium carbonate, magnesium stearate, maltodextrin, and mixtures thereof. The various ingredients and the excipient and/or carrier are mixed and formed into the desired form using conventional techniques. The tablet or capsule of the present invention may be coated with an enteric coating that dissolves at a pH of about 6.0 to 7.0. A suitable enteric coating that dissolves in the small intestine but not in the stomach is cellulose acetate phthalate. Further details on techniques for formulation for and administration may be found in the latest edition of Remington's Pharmaceutical Sciences (Maack Publishing Co., Easton, PA).

In some embodiments, dietary supplements of the present invention comprise krill oil as described above and one or more inert ingredients, especially if it is desirable to limit the number of calories added to the diet by the dietary supplement. The dietary supplement of the present invention may also contain optional ingredients including, for example, herbs, vitamins, minerals, enhancers, colorants, sweeteners, flavorants, inert ingredients, and the like. For example, the dietary supplement of the present invention may contain one or more of the following: ascorbates (ascorbic acid, mineral ascorbate salts, rose hips, acerola, and the like), dehydroepiandosterone (DHEA), Fo-Ti or Ho Shu Wu (herb common to traditional Asian treatments), Cat's Claw (ancient herbal ingredient), green tea (polyphenols), inositol, kelp, dulse, bioflavinoids, maltodextrin, nettles, niacin, niacinamide, rosemary, selenium, silica (silicon dioxide, silica gel, horsetail, shavegrass, and the like),

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spirulina, zinc, and the like. Such optional ingredients may be either naturally occurring or concentrated forms.

In some embodiments, the dietary supplements further comprise vitamins and minerals including, but not limited to, calcium phosphate or acetate, tribasic; potassium phosphate, dibasic; magnesium sulfate or oxide; salt (sodium chloride); potassium chloride or acetate; ascorbic acid; ferric orthophosphate; niacinamide; zinc sulfate or oxide; calcium pantothenate; copper gluconate; riboflavin; beta-carotene; pyridoxine hydrochloride; thiamin mononitrate; folic acid; biotin; chromium chloride or picolonate; potassium iodide; sodium selenate; sodium molybdate; phylloquinone; vitamin D3; cyanocobalamin; sodium selenite; copper sulfate; vitamin A; vitamin C; inositol; potassium iodide. Suitable dosages for vitamins and minerals may be obtained, for example, by consulting the U.S. RDA guidelines.

In further embodiments, the compositions comprise at least one food flavoring such as acetaldehyde (ethanal), acetoin (acetyl methylcarbinol), anethole (parapropenyl anisole), benzaldehyde (benzoic aldehyde), N butyric acid (butanoic acid), d or I carvone (carvol), cinnamaldehyde (cinnamic aldehyde), citral (2,6 dimethyloctadien 2,6 al 8, gera nial, neral), decanal (N decylaldehyde, capraldehyde, capric aldehyde, caprinaldehyde, aldehyde C 10), ethyl acetate, ethyl butyrate, 3 methyl 3 phenyl glycidic acid ethyl ester (ethyl methyl phenyl glycidate, strawberry aldehyde, C 16 aldehyde), ethyl vanillin, geraniol (3,7 dimethyl 2,6 and 3,6 octadien 1 ol), geranyl acetate (geraniol acetate), limonene (d , l , and dl ), linalool (linalol, 3,7 dimethyl 1,6 octadien 3 ol), linalyl acetate (bergamol), methyl anthranilate (methyl 2 aminobenzoate), piperonal (3,4 methylenedioxy benzaldehyde, heliotropin), vanillin, alfalfa (Medicago sativa L.), allspice (Pimenta officinalis), ambrette seed (Hibiscus abelmoschus), angelic (Angelica archangelica), Angostura (Galipea officinalis), anise (Pimpinella anisum), star anise (Illicium verum), balm (Melissa officinalis), basil (Ocimum basilicum), bay (Laurus nobilis), calendula (Calendula officinalis), (Anthemis nobilis), capsicum (Capsicum frutescens), caraway (Carum carvi), cardamom (Elettaria cardamomum), cassia, (Cinnamomum cassia), cayenne pepper (Capsicum frutescens), Celery seed (Apium graveolens), chervil (Anthriscus cerefolium), chives (Allium schoenoprasum), coriander (Coriandrum sativum), cumin (Cuminum cyminum), elder flowers (Sambucus canadensis), fennel (Foeniculum vulgare), fenugreek (Trigonella foenum graecum), ginger (Zingiber officinale), horehound (Marrubium vulgare), horseradish (Armoracia lapathifolia),

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hyssop (Hyssopus officinalis), lavender (Lavandula officinalis), mace (Myristica fragrans), marjoram (Majorana hortensis), mustard (Brassica nigra, Brassica juncea, Brassica hirta), nutmeg (Myristica fragrans), paprika (Capsicum annuum), black pepper (Piper nigrum), peppermint (Mentha piperita), poppy seed (Papayer somniferum), rosemary (Rosmarinus officinalis), saffron (Crocus sativus), sage (Salvia officinalis), savory (Satureia hortensis, Satureia montana), sesame (Sesamum indicum), spearmint (Mentha spicata), tarragon (Artemisia dracunculus), thyme (Thymus vulgaris, Thymus serpyllum), turmeric (Curcuma longa), vanilla (Vanilla planifolia), zedoary (Curcuma zedoaria), sucrose, glucose, saccharin, sorbitol, mannitol, aspartame. Other suitable flavoring are disclosed in such references as Remington's Pharmaceutical Sciences, 18th Edition, Mack Publishing, p. 1288-1300 (1990), and Furia and Pellanca, Fenaroli's Handbook of Flavor Ingredients, The Chemical Rubber Company, Cleveland, Ohio, (1971), known to those skilled in the art.

In other embodiments, the compositions comprise at least one synthetic or natural food coloring (e.g., annatto extract, astaxanthin, beet powder, ultramarine blue, canthaxanthin, caramel, carotenal, beta carotene, carmine, toasted cottonseed flour, ferrous gluconate, ferrous lactate, grape color extract, grape skin extract, iron oxide, fruit juice, vegetable juice, dried algae meal, tagetes meal, carrot oil, corn endosperm oil, paprika, paprika oleoresin, riboflavin, saffron, tumeric, tumeric and oleoresin).

In still further embodiments, the compositions comprise at least one phytonutrient (e.g., soy isoflavonoids, oligomeric proanthcyanidins, indol 3 carbinol, sulforaphone, fibrous ligands, plant phytosterols, ferulic acid, anthocyanocides, triterpenes, omega 3/6 fatty acids, conjugated fatty acids such as conjugated linoleic acid and conjugated linolenic acid, polyacetylene, quinones, terpenes, cathechins, gallates, and quercitin). Sources of plant phytonutrients include, but are not limited to, soy lecithin, soy isoflavones, brown rice germ, royal jelly, bee propolis, acerola berry juice powder, Japanese green tea, grape seed extract, grape skin extract, carrot juice, bilberry, flaxseed meal, bee pollen, ginkgo biloba, primrose (evening primrose oil), red clover, burdock root, dandelion, parsley, rose hips, milk thistle, ginger, Siberian ginseng, rosemary, curcumin, garlic, lycopene, grapefruit seed extract, spinach, and broccoli.

In still other embodiments, the compositions comprise at least one vitamin (e.g., vitamin A, thiamin (B1), riboflavin (B2), pyridoxine (B6), cyanocobalamin (B12), biotin, ascorbic acid (vitamin C), retinoic acid (vitamin D), vitamin E, folic acid and other folates,

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vitamin K, niacin, and pantothenic acid). In some embodiments, the particles comprise at least one mineral (e.g., sodium, potassium, magnesium, calcium, phosphorus, chlorine, iron, zinc, manganese, flourine, copper, molybdenum, chromium, selenium, and iodine). In some particularly preferred embodiments, a dosage of a plurality of particles includes vitamins or minerals in the range of the recommended daily allowance (RDA) as specified by amino acid supplement formula in which at least one amino acid is included (e.g., I-carnitine or tryptophan).

In some embodiments, the present invention provides functional food products containing krill oil as described above. Examples of functional foods include, but are not limited to dairy products such yogurt, milk and cheese, cereals, beverages, shakes, powdered supplements, and the like.

#### 2. Processes for making krill oil

The processes of the present invention are useful with krill oil produced by a variety of processes. Suitable processes for producing krill oil include extraction with polar solvents such as ethanol, supercritical fluid extraction, extraction with non-polar organic solvents such as acetone, cold pressing, etc. See, e.g., WO2009/027692, WO2008/117062, WO2003/011873, all of which are incorporated herein by reference. The processes of the present invention may also be performed on commercially available krill oils such as those supplied by Aker Biomarine, Neptune Bioressources, and Enzymotec.

As described above, the present inventors have discovered that krill oil is a multiphase dispersion. The present invention provides processes for separating the multiphase dispersion into two or more phases that can be separated. In some embodiments, krill oil in the multiphase dispersion state is further processed by mixing the krill oil with a polar solvent and incubating the mixture for a period of time (the incubation period) sufficient for the formation of least two phases in the mixture. The upper phase, or phospholipid phase, comprises the krill oil of the present invention.

In preferred embodiments, the phospholipid phase is separated from any other phases formed during the incubation phase, for example, by decanting the phospholipid phase. In some embodiments, the incubation phase is from about 0.5 hours to about 48 hours, 0.5 hours to 24 hours, 0.5 hours to 12 hours, 0.5 hours to 6 hours, 0.5 hours to 4 hours, or 1 hour to 4 hours. In some embodiments, the incubation is conducted at from

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about 0 C to about 25 C. In some embodiments, the incubation is conducted at from about 4 C to about 10 C, about 4 C to about 20 C, about 4 C to about 25 C, about 10 C to about 20 C, about 10 C to about 25 C, or about 15 C to 25 C.

In some embodiments, the polar solvent is an alcohol, such as a monohydric alchohol. Suitable monohydric alcohols include, but are not limited to, methanol, ethanol, propanol and isopropanol. Other polar solvents include dimethyl sulfoxide (DMSO), formamide, acetonitrile, N,N-dimethylformamide (DNF) and other solvents with a dielectric constant of higher than 15 or 20. In some embodiments, the krill oil is diluted with the polar solvent at a ratio of krill oil: polar solvent of 1:0.5 to 1:10, 1:1 to 1:5, 1:1 to 1:3, 1:2 to 1:5, 1:2 to 1:4, or 1:2 to 1:3. In some embodiments, the polarity of the solvent is adjusted by adding water. In some embodiments, the ratio of polar solvent, for example ethanol, to water is from about 1:1 to 100:1, 2:1 to 100:1, 2:1 to 20:1, 3:1 to 20:1, 4:1 to 20:1, 5:1 to 20:1, or 10:1 to 20:1.

#### 15 3. Uses of krill oil

The krill oil of the present invention is useful for treatment of any disease, disorder or condition in which omega-3 PUFAs have been shown to be effective. Diseases and disorders that may be treated with the omega-3 fatty acid formulations described herein include alopecia, Alzheimer's dementia, angina, anxiety disorders, asthma, attention deficit disorder, attention-deficit hyperactivity disorder, atopic dermatitis, autism, bipolar disorder, borderline personality disorder, cardiovascular disease, chronic fatigue syndrome, chronic pain, chronic polyarthritis, cognitive disorders, communication disorders, colitis, Crohn's disease, cystic fibrosis, dementia, depression, diabetes (of the non-insulin dependent or insulin dependent forms), diabetes-related sequelae, diabetic neuropathy, dry eyes and other inflammatory eye disorders, dry skin, dysmenorrhea, eating disorders (such as anorexia nervosa or bulimia nervosa and obesity), eczema, fibromyalgia, gout, learning disorders (e.g. reading, spelling, mathematics, receptive, and expressive language, and motor skills disorders), lupus, male infertility, metabolic syndrome, melanoma, mild cognitive impairment, migraine, mood disorders, multiple sclerosis, obsessive-compulsive disorder, oppositional-defiant disorder, osteoarthritis, osteoporosis, pervasive developmental disorders, 'polyarteritis nodosa, psoriasis, psoriatic arthritis, rheumatoid arthritis, schizophrenia, sclerodermia, self-injurious behavior, sickle cell anemia, tic

disorders, tinnitus, ulcerative colitis, or vasculitic disorders (such as polyarteritis nodosa and temporal arthritis. Cardiovascular disease and disorders that can be treated with the omega-3 fatty acid formulations described herein include angina, atherosclerosis, hypercholesterolemia, hypertriglyceridemia, low HDL, high blood pressure, Raynaud's disease, and cardiac arrhythmias. Methods of treatment with the omega-3 fatty acid formulations described herein include prophylaxis with Omega-3 formulations to prevent post-cardiotomy (including but not limited to coronary artery bypass graft surgery and valve surgery) complications (including but not limited to depression, neuro-cognitive decline, congestive heart failure and infarction, clotting events, and arrhythmias) as well as for the treatment for such complications.

#### **Experimental**

#### Example 1

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Krill oil is extracted from krill meal (Aker Biomarine) by ethanol extraction. Briefly, krill meal is extracted with ethanol for 1 hour at 15-30 C. The liquid fraction is separated by filtration. The liquid fraction is concentrated by evaporating the ethanol under a vacuum at about 50 C until the concentration of ethanol is reduced to about 20%. The concentrated liquid phase is then centrifuged to remove any remaining solids or precipitates and then evaporated under a vacuum to a final concentration of less than 0.5% ethanol. The extracted krill oil is a multiphase dispersion. An exemplary batch of krill oil prepared by this process had the following composition.

	Triacylgycerol	g/100g oil	30
•	Diacylglycerol	g/100g oil	0.7
25	Monoacylglycerol	g/100g oil	<1
	Free fatty acids	g/100g oil	4.8
	Cholesterol	g/100g oil	1.2
	Cholesterol esters	g/100g oil	<0.5
	Phosphatidylethanolamine	g/100g oil	1.5
30	Phosphatidylinositol	g/100g oil	<1
	Phosphatidylserine	g/100g oil	<1
	Phosphatidylcholine	g/100g oil	40

Lyso-phosphatidylcholine	g/100g oil	3.3
Total polar lipids	g/100g oil	44.6
Total neutral lipids	g/100g oil	36.9
Total sum lipids	g/100g oil	81.5

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#### **Example 2**

This example describes attempts to concentrate omega-3 PUFA in krill oil by lowering temperature. It was thought that by lowering the temperature, highly saturated fat would turn solid at a faster rate and sediment or in another way form a layer in the column of krill oil. The experiment tested whether triacylglycerol (TAG) and phospholipids (PL) with saturated fatty acyl chains would form a layer or if TAG and PL would form layers independent of the fatty acyl chain type. Krill oil described in Example 1 was diluted with 20% absolute ethanol and a glass column was filled. The column was placed in a refrigerator overnight. The oil was examined for the formation of layers. The oil turned very viscous, paler but not white. No layers formed. The oil was then taken to room temperature. Again, there was no layer formation. It was not possible to determine if some portion of the fat turns solid faster than other parts. The lack of layer formation can possibly be explained by the fact that the viscosity of the oil is too high to allow vertical movement of fat with higher or lower density. Dilution with 50% absolute ethanol had the same result of no layer formation even though the solution was less viscous.

#### Example 3

This example describes the concentration of omega-3 PUFA in krill oil. Krill oil described in Example 1 was diluted 1:1 with absolute ethanol (B), 1:2 with absolute ethanol (C), 1:3 with absolute ethanol (D), and 1:3 with 95% ethanol (E) in 15 ml polypropylene vials. No phase separation was seen in B, C, or D. E was slightly opaque and after five minutes a layer formed at the bottom. All four vials were then stored at -30 C for two hours. At this time, the oil was solidified and appeared as a white solid. No visible layers were observed in the solid form. After thawing at room temperature, the solid oil melted and drops formed that sedimented in the vials C, D, and E. The volume of the bottom layer increased with increasing ethanol dilution. Vial E had a larger lower phase than vial D. The bottom layer was darker than the top layer in all vials but C.

The solubility of the lower phase was examined. The lower phase was not soluble in ethanol or water. The lower phase exhibited good solubility in hexane and the color was more brown than the top layer.

The UV spectra of the top and bottom layers was examined. The UV spectra of the lower phase indicated the presence of astaxanthin together with components absorbing at lower wavelengths. The UV spectra of the upper phase indicated a similar pattern with astaxanthin less prominent compared to lower wavelengths.

The upper and lower phases were examined by thin layer chromatography (TLC). The TLC data indicates that after separation of krill oil into two phases is a higher portion of TAG in the lower phase than in the upper phase. It also appears that there is more PL in the upper phase.

The upper and lower phases were next examined by GC-FID. The results are provided below.

15	Fraction	Upper	Upper	Lower	Lower
	ID	1 (D)	2 (E)	3 (D)	4 (E)
	C12:0	0.1	0.2	0.4	0.4
	C14:0	6.3	6.4	17.4	17.6
	C15:0	0.4	0.3	0.5	0.5
20	C16:0	19.9	20.1	19.6	19.8
	C16:1	3.5	3.5	7.8	7.8
	C18:0	0.9	0.9	1.3	1.3
	C18:1, t6-11	0.3	0.3	0.6	0.6
	C18:1, c9	6.3	6.4	13.2	13.2
25	C18:1, c11	5.7	5.7	6.8	6.7
	C18:2, n-6	1.6	1.6	1.8	1.8
	C20;0	0.1	0.1	0.1	0.1
	C18:3, n-6	0.3	0.2	0.2	0.2
	C18:3, n-3	1.0	1.0	0.8	8.0
30	C20:1, n-9	0.5	0.5	0.8	8.0
	C18:4, n-3	2.4	2.4	3.2	3.2
	C20:2, n-6	0.1	0.1	0.1	0.1

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	C22:0	0.2	0.2	0.2	0.2
	C20:3, n-6	0.1	0.1	0.1	0.1
	C20:3, n-3	0.1	0.1	0.1	0.1
	C20:4, n-6/C22:1, n-9	1.3	1.3	0.5	0.5
5	C20:4, n-3	0.4	0.4	0.2	0.2
	C20:5, n-3	26.0	26.0	7.7	7.7
	C22:5, n-3	0.6	0.6	0.3	0.3
	C22:6, n-3	12.7	12.7	3.8	3.7
	SFA	27.8	28.2	39.6	40.0
10	MUFA	16.3	16.4	29.2	29.2
	PUFA (n-6)	3.3	3.3	2.5	2.6
	PUFA (n-3)	43.1	43.2	16.1	16.0
	Total PUFA	46.4	46.4	18.6	18.6

This analysis indicates that there is a higher relative portion of omega-3 PUFA in the upper phase compared to the lower phase. The upper phase has 43 % total omega-3 PUFA. EPA and DHA is equally concentrated. In the lower phase the fatty acids 14:0 (myristic acid) and 18:1 (c9 oleic acid) are concentrated. There are no changes in 16:0.

This data describes the separation of a krill oil with approximately 40 % nonpolar lipids (30 % TAG) into two phases by adding ethanol/water and lowering the temperature. The layers form by passive sedimentation. TLC data indicates that PL is high in the upper phase and that TAG is high in the lower phase. GC-FID data shows that omega-3 lipids is high in the upper phase (43 %, 43 g/100 g FAME) and 18 % in the lower phase. An intriguing observation is that the particularly unhealthy fatty acid 14:0 is lowered in the upper phase.

The present invention is not limited to any particular mechanism. Nevertheless, the inventors have discovered that krill oil produced by ethanol extraction is a multiphase dispersion. Unsoluble components such as triglycerides are entrapped in the soluble phase (polar lipids) and coextracted. The processes describe above take advantage of this fact to provide krill oil compositions with concentrated amounts of desirable omega-3 fatty acids.

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#### Claims

What is claimed is:

- 5 1. Krill oil comprising greater than about 22% EPA (w/w total fatty acids), greater than about 10% DHA (w/w total fatty acids), from 4% to 8% myristic acid (w/w total fatty acids), from 3% to 9% c9 oleic acid (w/w total fatty acids), and 20 to 4000 ppm astaxanthin.
  - 2. Krill oil of Claim 1, further comprising about 22% to 30% EPA.

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- 3. Krill oil of Claim 1, further comprising about 10% to 15% DHA.
- 4. Krill oil of Claim 1, wherein said krill oil is extracted from Euphausia superba.
- 15 5. Krill oil of Claim 1, wherein the ratio of DHA and EPA: omega 6 (w/w total fatty acids) is from about 10:1 to 14:1.
  - 6. Krill oil of Claim 1, wherein the ratio of DHA and EPA: c9 oleic acid (w/w total fatty acids) is from about 4:1 to 8:1.

- 7. Krill oil of Claim 1, wherein the ratio of DHA and EPA: myristic acid (w/w total fatty acids) is from about 4:1 to 8:1.
- 8. Krill oil of Claim 1, wherein the ratio of DHA and EPA: myristic acid and c9 oleic acid (w/w total fatty acids) is from about 2:1 to 4:1.
  - 9. Krill oil of Claim 1, wherein the ratio omega 3: omega 6 (w/w total fatty acids) is from about 11:1 to 15:1.
- 30 10. Krill oil of Claim 1, wherein the ratio of omega 3: c9 oleic acid (w/w total fatty acids) is from about 5:1 to 9:1.

- 11. Krill oil of Claim 1, wherein the ratio of omega 3: myristic acid (w/w total fatty acids) is from about 5:1 to 9:1.
- 12. Krill oil of Claim 1, wherein the ratio of omega 3: myristic acid and c9 oleic acid (w/w total fatty acids) is from about 2.5:1 to 4.5:1.
  - 13. A capsule containing the krill oil of Claims 1 to 14.
  - 14. A food product containing the krill oil of Claims 1 to 14.
  - 15. A dietary supplement containing the krill oil of Claims 1 to 14.
  - 16. An oil in water emulsion containing the krill oil of Claims 1 to 14.
- 15 17. Use of the krill oil of Claims 1 to 14 for oral administration.
  - 18. Use of the krill oil of Claims 1 to 14 for treatment of a condition for which omega-3 is effective.

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(54) Title: CONCENTRATION OF OMEGA-3 POLYUNSATURATED FATTY ACIDS IN KRILL OIL

(57) Abstract: The present invention relates to krill oil, and in particular to krill oil with elevated levels of omega-3 fatty acids and decreased levels of saturated fatty acids.

#### INTERNATIONAL SEARCH REPORT

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PCT/IB2013/001959 A, CLASSIFICATION OF SUBJECT MATTER
INV. A23D9/00 A23D9 A23D9/013 A23D9/04 A61K35/56 ADD. According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) A23L A23D A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) EPO-Internal, BIOSIS, FSTA, PAJ, WPI Data C. DOCUMENTS CONSIDERED TO BE RELEVANT Category\* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. 1-18 Х US 2008/274203 A1 (BRUHEIM INGE [NO] ET AL) 6 November 2008 (2008-11-06) paragraphs [0002], [0009] - [0017], [0026], [0028], [0031], [0036] tables 9,13,14 Χ US 2011/223246 A1 (OPHEIM JOAR [US]) 1-18 15 September 2011 (2011-09-15) paragraphs [0012], [0066], [0067], [0085] - [0091] paragraph [0094] US 2011/130458 A1 (BREIVIK HARALD [NO] ET 1 - 18Х AL) 2 June 2011 (2011-06-02) paragraphs [0002], [0009], [0066] - [0069] tables 1,2,5 -/--Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier application or patent but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination special reason (as specified) document referring to an oral disclosure, use, exhibition or other being obvious to a person skilled in the art document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 17 December 2013 07/01/2014 Authorized officer Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Couzy, François Fax: (+31-70) 340-3016

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A,P	WO 2012/139588 A2 (TRIPLENINE PHARMA AS [DK]; SOERENSEN HANS OTTO [DK]; JENSEN NILS CHRIS) 18 October 2012 (2012-10-18) page 6, line 21 - page 7, line 8 example 2 claims 1,2,11,16		1-18

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## Espacenet

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Method for enriching phosphatidyl inositol from antarctic krill

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Abstract of CN102746941 (A)

The invention discloses a method for enriching phosphatidyl inositol from antarctic krill. The method includes the steps: (1) adding 1 kilogram of frozen antarctic krill to 1-1.3 liters of ethanol solution with the volume fraction of 90-100%, stirring to extract for 4-6 times under natural conditions, keeping extraction each time for 1-3h, and filtering and combining extracting solutions so as to obtain filtrate; (2) concentrating the obtained filtrate under the pressure of negative 0.07-negative 0.09Mpa at the temperature of 60-65 DEG C until 8-10% of the volume of the filtrate remains so as to obtain concentrated extracting solution; (3) adding isometric normal hexane into the obtained concentrated extracting solution, uniformly mixing and statically layering so as to obtain transparent reddish bottom solution; and (4) removing the normal hexane from the obtained bottom solution by vaporizing under the pressure of negative 0.07-negative 0.09Mpa at the temperature of 50-55 DEG C so as to obtain oily solution, namely, a phosphatidylinositol-enriched product, wherein the product comprises 2.70-2.76% of phospholipid, and the phosphatidyl inositol accounts for 50.5-51.92% of the phospholipid. Reextraction, separation, purification and the like can be further performed on the basis so as to obtain the phosphatidyl inositol.

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权利要求书1页 说明书3页

## (54) 发明名称

一种从南极磷虾中富集磷脂酰肌醇的方法

### (57) 摘要

本发明公开了一种从南极磷虾中富集磷脂酰肌醇的方法,步骤如下:(1)取1公斤冰冻的南极磷虾,加入到1~1.3升的体积分数为90~100%的乙醇液中,自然条件下搅拌提取4~6次,每次1~3h,提取液过滤合并,得滤液:(2)将上述得到的滤液于60~65°C和-0.07~-0.09MPa条件下浓缩至滤液体积的8~10%,得浓缩提取液:(3)向上述得到的浓缩提取液中加入等体积的正己烷,混匀,静止分层后,取透明微红的最下层液:(4)将上述得到的最下层液于50~55°C和-0.07~-0.09MPa条件下蒸发除去正己烷,得油状液,即为富集了磷脂酰肌醇的产品,该产品中含磷脂2.70~2.76%,其中PI占磷脂的50.5~51.92%,可以在此基础上对磷脂酰肌醇进行进一步的再提取、分离、纯化等。

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- 1. 一种从南极磷虾中富集磷脂酰肌醇的方法,步骤如下;称冰冻的南极磷虾500g,加入95%乙醇600mL,在室温下搅拌3h,倒出上清液并过滤;沉淀再反复提取4次,共提取5次,提取液过滤合并得滤液;将滤液在60℃,-0.07MPa条件下蒸发浓缩至合并滤液体积的10%,得到浓缩提取液;向浓缩提取液中加入同体积的正己烷,摇匀静置分层,取透明微红色的最下层液,在52℃,-0.08MPa条件下蒸发除去正己烷,得含有磷脂酰肌醇的油状液。
- 2. 一种从南极磷虾中富集磷脂酰肌醇的方法,步骤如下:称冰冻的南极磷虾 500g,加入 90% 乙醇 550mL;在室温下搅拌提取 3h,倒出上清液并过滤;沉淀再重复提取 3次,共提取 4次,提取液过滤合并得滤液;将滤液在 65℃, -0.09MPa 条件下浓缩至合并滤液体积的 8%,得到浓缩提取液;向浓缩液中加入同体积的正己烷,摇匀静置分层,取透明微红的最下层液,在 55℃, -0.09MPa 条件下蒸除溶剂,得含有磷脂酰肌醇的油状液。
- 3. 一种从南极磷虾中富集磷脂酰肌醇的方法,步骤如下:称冰冻的南极磷虾500g,加入无水乙醇650mL,在室温下搅拌提取1h,倒出上清液并过滤;沉淀再重复提取5次,共提取6次;提取液过滤合并得滤液;将滤液在60℃,-0.08MPa条件下浓缩至合并滤液体积的10%,得到浓缩提取液;向浓缩液中加入同体积的正己烷,摇匀静置分层,取透明微红的最下层液,在50℃,-0.07MPa下蒸除溶剂,得含有磷脂酰肌醇的油状液。
- 4. 一种从南极磷虾中富集磷脂酰肌醇的方法,步骤如下:称冰冻的南极磷虾500g,加入95%乙醇500mL,在室温条件下搅拌提取2h,倒出上清液并过滤;沉淀再重复提取4次,共提取5次,提取液过滤合并得滤液;将滤液在65℃,一0.09MPa条件下浓缩至合并滤液体积的9%,得到浓缩提取液;向该浓缩液中加入同体积的正己烷,摇匀静置分层,取透明微红的最下层液,在50℃,一0.09MPa条件下蒸发溶剂,得含有磷脂酰肌醇的油状液。

## 一种从南极磷虾中富集磷脂酰肌醇的方法

### 技术领域

[0001] 本发明涉及一种从南极磷虾中富集磷脂酰肌醇的方法。

### 背景技术

[0002] 南极磷虾是一种生活在南冰洋的南极洲水域的磷虾,以群集方式生活,有时密度高达1万-3万只/M³,它们可能是地球上最成功的动物物种,据统计大约有5-50亿吨(因统计方法不一样有较大差异)。

[0003] 目前南极磷虾渔业每年约 10 万吨,其中 80% 由日本捕获,由于南大洋以磷虾为主食物的须鲸也被歼捕殆尽。有关科学家曾根据旺盛时期鲸类对磷虾类的摄食量估计大约每年有 1.5 亿吨由于鲸类资源的衰退而剩余的磷虾可供捕获。世界上每年的全部的渔业产量为 0.99 亿吨,这标志着南极磷虾有较大的可利用空间。我国正在试探捕南极磷虾以完善南极磷虾的捕捞和相关海上技术。南极磷虾的大量捕捞标志着深加工技术的开始。在日本南极磷虾大多作为料理,而在其他国家则主要作为渔业的饲料和鱼的饵料。南极磷虾高附加值的研究是其深加工的研究核心。

[0004] 南极磷虾含有较为丰富的油脂,含有不饱和脂肪酸、虾青素和磷脂。磷脂酰肌醇 (phosphatidylinositol;PI)是细胞信号的通路成分,称为第三信使,信号通路的研究是当前生命科学研究的热点,也是生物制剂争相开发利用的重大课题。已知的信使如第一信使 脂类激素和含氮激素,第二信使 cAMP 等均或为药物,或为保健成分。第三信使 PI 在人体、细胞代谢中起着极为重要的生理生化作用。是细胞快速应答和缓慢应答的启点、交叉点、关键点。PI 就像通信网络中控制着信息传递方向的接线员,控制着人体生理生化、细胞生与死的进展方向。

[0005] PI 的应用开发备受世界各国关注,2001 年美国开发出了世界上第一个以 PI 作为中间体的抗艾滋病药物(US 6316424.B)。科学家在 PI 的肌醇环上接上磺酰基基团(-SO<sub>3</sub>),当这种药物插入细胞膜后,暴露在膜外带有 1-几个磺酰基团的肌醇环有极强的杀病菌、病毒作用,就像细胞全身穿上了盔甲。PI 有望成为抗其他病毒药物的中间体。在流感病毒、SARS、霍乱等菌毒肆虐世界的今天,PI 的应用具有极其广阔的前景。日本等国把 PI 也用于化妆品,作为药物脂质体和日常保健品的原料、食品和药物的乳化剂(JP 05097873 A<sub>2</sub>)海产品的保鲜剂;PI 还是心、肝、肾等人体重要器官移植前保存液的主要成分,有抗细胞凋亡(WO 99/47101),也有很好的免疫效果(WO 03/013513 A1)。另外实验证明 PI 有抗动脉粥样硬化的作用,有利尿和治疗便秘的作用等。

[0006] PI 因 sn-1 和 sn-2 位所联接的脂肪酸不一而分子结构和分子量有很大差异。不同动物体 PI 分子群中各种 PI 分子种类及其比例有很大差异,即 PI 是一个分子群。不同的动物体中磷脂中的卵磷脂(PC)、脑磷脂(PE)、PI 和磷脂酸,磷酯酰丝氨酸(PS)等比例也有很大差异。南极磷虾中虾油(磷脂是油的伴随物)的磷脂中 PC 占 80%以上,而 PS、PE、PA 以及溶血性磷脂的总含量低于 20%,而 PI 量很少,因此制备南极磷虾中的 PI,最重要的第一步是 PI 的富集,然后才能在此基础上进一步再提取,分离,纯化。至今,人们还尚未对南极磷

虾中的 PI 进行过研究报道。

### 发明内容

[0007] 针对上述现有技术,本发明提供了一种从南极磷虾中富集磷脂酰肌醇的方法,为南极磷虾中磷脂酰肌醇的深入研究奠定了基础。

[0008] 本发明是通过以下技术方案实现的:

[0009] 一种从南极磷虾中富集磷脂酰肌醇的方法,步骤如下:

[0010] (1)取 1 公斤冰冻的南极磷虾,加入到 1  $\sim$  1.3 升的体积分数为 90  $\sim$  100%的乙醇液中,自然条件下搅拌提取 4  $\sim$  6 次,每次 1  $\sim$  3h,提取液过滤合并,得滤液;

[0011] (2)将上述得到的滤液于  $60 \sim 65$ ° C和  $-0.07 \sim -0.09$ MPa 条件下浓缩至滤液体积的  $8 \sim 10\%$ , 得浓缩提取液;

[0012] (3)向上述得到的浓缩提取液中加入等体积的正己烷,混匀,静止分层后,取透明微红的最下层液;

[0013] (4)将上述得到的最下层液于  $50 \sim 55^{\circ}$  C 和  $-0.07 \sim -0.09$ MPa 条件下蒸发除去正己烷,得油状液,即为富集了磷脂酰肌醇的产品,该产品中含磷脂  $2.70 \sim 2.76\%$  (质量分数),其中 PI 占磷脂的  $50.5 \sim 51.92\%$  (质量分数)。

[0014] 本发明的从南极磷虾中富集磷脂酰肌醇的方法,在低温下进行,能够保证南极磷虾中的磷脂、DHA、EPA、虾红素、动物类黄酮素、不饱和脂肪酸、维生素 A、维生素 E 和微量元素等成分不受到破坏,且富集过程中的上层溶液可以回收再利用,提取后的南极磷虾可以再用于生产其它产品。经本发明的方法得到的产品,磷脂酰肌醇的富集效率高,产品中含磷脂 2.70~2.76%,其中磷脂酰肌醇占磷脂的 50.5~51.92%,可以在此基础上对磷脂酰肌醇进行进一步的再提取、分离、纯化等。

### 具体实施方式

[0015] 下面结合实施例对本发明作进一步的说明。

[0016] 实施例 1

[0017] 称冰冻的南极磷虾 500g(含水 80%,虾干物质占 20%,下同),加入 95% 乙醇 600mL,在室温下搅拌 3h,倒出上清液并过滤。沉淀再反复提取 4次,共提取 5次,提取液过滤合并得滤液;将滤液在 60℃, -0.07MPa 条件下蒸发浓缩至合并滤液体积的 10%,得到浓缩提取液;向浓缩提取液中加入同体积的正己烷,摇匀静置分层,取透明微红色的最下层液,在52° C, -0.08MPa 条件下蒸发除去正己烷,得 13g 油状液。该液体中磷脂含量为 2.72%,其中 PI 占磷脂的 51.92%。

[0018] 实施例 2

[0019] 称冰冻的南极磷虾 500g, 加入 90% 乙醇 550mL。在室温下搅拌提取 3h, 倒出上清液并过滤。沉淀再重复提取 3次, 共提取 4次, 提取液过滤合并得滤液; 将滤液在65° C, -0.09MPa条件下浓缩至合并滤液体积的 8%, 得到浓缩提取液; 向浓缩液中加入同体积的正己烷, 摇匀静置分层, 取透明微红的最下层液, 在55° C, -0.09MPa条件下蒸除溶剂, 得 11g 油状液。该液体中磷脂含量为 2.70%, 其中 PI 占磷脂的 50.5%。

[0020] 实施例3

[0021] 称冰冻的南极磷虾 500g, 加入无水乙醇 650mL, 在室温下搅拌提取 1h, 倒出上清液并过滤。沉淀再重复提取 5次, 共提取 6次。提取液过滤合并得滤液; 将滤液在 60° C, -0.08MPa 条件下浓缩至合并滤液体积的 10%, 得到浓缩提取液; 向浓缩液中加入同体积的正己烷, 摇匀静置分层, 取透明微红的最下层液, 在 50° C, -0.07MPa 下蒸除溶剂, 得 12g 油状液。该液体中磷脂含量为 2.76%, 其中 PI 占磷脂的 51.7%。

[0022] 实施例 4

[0023] 称冰冻的南极磷虾 500g, 加入 95% 乙醇 500mL, 在室温条件下搅拌提取 2h, 倒出上清液并过滤。沉淀再重复提取 4 次, 共提取 5 次, 提取液过滤合并得滤液; 将滤液在 65° C, -0.09MPa 条件下浓缩至合并滤液体积的 9%, 得到浓缩提取液; 向该浓缩液中加入同体积的正己烷, 摇匀静置分层, 取透明微红的最下层液, 在 50° C, -0.09MPa 条件下蒸发溶剂, 得 12g 油状液, 该液体中磷脂含量为 2.74%, 其中 PI 占磷脂的 51.9%。



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METHOD FOR COLLECTING KRILL PHOSPHOLIPID AND FUNCTIONAL FOOD AND NERVE FUNCTION IMPROVING AGENT HAVING NERVE FUNCTION IMPROVING EFFECT

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MASAZUMI, ; KIMURA SEIJI, ; NONAKA MICHIO)

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PURPOSE: To obtain an useful phospholipid in high purity by fractionating an ethanol extracted total lipid of fresh krill dehydrated by vacuum freeze drying method to specific two ingredients using an absorption column chromatography and further isolating these ingredients using a fraction collector. CONSTITUTION: A fresh krill is dehydrated to <=6% water content using a vacuum freeze drying method. Then the dried krill is homogenized with ethanol to extract total lipid. The ethanol is removed as much as possible from the total lipid and the extracted total lipid ia fractionated to soluble fraction and insoluble fraction using an acetone based solvent or hexane based solvent as eluate and then the solvent is cleaned from the insoluble fraction to give a crude phospholipid.; Then the crude phospholipid is fractionated to phosphatidyl choline and phosphatidyl ethanolamine with an absorption column chromatography using ethanol based solvent, acetone based solvent or hexane based solvent as an eluate. Then

each phospholipid ingredient is isolated therefrom in a high purity of about 90-95% by a fraction collector.

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#### (54) 【発明の名称】 オキアミリン脂質の分取方法

#### (57) 【特許請求の範囲】

【請求項1】生オキアミを真空凍結乾燥法により脱水し、得られたオキアミをエタノールで総脂質を抽出し、得られた総脂質からエタノールを除去し、アセトンに溶解し、可溶区分と不可溶区分に分画し、不可溶区分を更にアセトンで洗浄し粗リン脂質を得、この粗リン脂質をエタノールを溶離液として、シリカゲルを充填剤として、吸着カラムクロマトグラフィーを用いて90~95%のホスファチジルコリンとホスファチジルエタノールアミンを分画するようにしたことを特徴とするオキアミリン脂質の分取方法。

#### 【発明の詳細な説明】

#### 「産業上の利用分野」

本発明は、オキアミからリン脂質を分離抽出する方 法、特に、生体内において重要な生理活性を示すホスフ ァチジルコリン及びホスファチジルエタノールアミンを 高純度に分取する方法であり、こうして分取されたホス ファチジルコリン及びホスファチジルエタノールアミン 等が記憶力改善剤として利用可能なものである点に特徴 を有する技術に関する。

#### 「従来技術」

最近、高齢化社会を迎えて、老人性痴呆症が大きな社会問題になっている。老人性痴呆症は、神経系の障害を原因として起こるアルツハイマー型痴呆症と、脳血管障害を原因として起こる脳血管性痴呆症との二つの型に大別できる。前者のアルツハイマー型痴呆症の場合には、脳内の神経化学的な変化として、神経伝達物質であるアセチルコリンの生産が著しく低下していることが知られており、この病気の予防や治療法として、低下したコリン系の代謝を補給することにより生理機能を回復せんと

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することが行なわれている。例えば、PCT特許出願公表 昭56-500374号「レシチンを投与することにより病気を 治療するための方法および組成物」、特開昭59-167514号「脳機能亢進剤組成物」、特開昭60-214734号「神経障害及び走化の治療組成物および治療方法」等がそれである。

即ち、コリン含有リン物質であるホスファチジルコリンを摂取することにより、脳内にアセチルコリンを供給し、これによりアルツハイマー型痴呆症やその他の神経障害の予防と治療が期待されている。

また、リン脂質の一種であるホスファチジルエタノールアミンはS-アデノシルメチオンニンからのメチル基移転反応によりホスファチジルコリンに変換される。従って、当該ホスファチジルエタノールアミンもアルツハイマー型痴呆症やその他の神経障害の予防と治療剤としての利用が期待されている。

本発明者は、特に、グリセロリン脂質である、これら ホスファチジルコリン及びホスファチジルエタノールア ミンといったリン脂質に注目し、これを記憶力改善剤の 原料として利用が可能な状態で工業的に分取する方法を 研究開発せんとしたものである。

従来、天然物からリン脂質を工業的に精製する場合の原料といえば大豆が一般的であり、大豆リン脂質は主に健康食品等として、商品化されている。従来の大豆リン脂質精製法は、まず原料大豆をクロロホルム・メタノール系の溶媒で総脂質を抽出し、次に当該総脂質をアセトンで分画し、可溶性区分と不溶性区分に分ける。当該アセトン可溶性区分には中性脂質、コレステロール、遊離脂肪酸等が分画されており、またアセトン不溶性区分にはリン脂質が分画されている。そこで、次に、アセトン不溶性区分を90%エタノールで処理して、アルコールに溶けるホスファチジルコリンと不溶性のホスファチジルエタノールアミンとを得る。

また、「Juranl of Chromatography」,Vol,365 (198 6) p. 229-235には、オキアミから脂質を抽出し、ヘキサンを溶離液としてシリカゲルカラムを用いてカラムクロマトグラフィーを行い、流出時間ごとにフラクション分画をおこない、ホスファチジルコリンとホスファチジルエタノールアミンとを単離するオキアミリン脂質の分取方法が記載されている。

「発明が解決しようとする問題点」

しかし、前記大豆を原料としたリン脂質の精製法の場合には、得られるホスファチジルコリン及びホスファチジルエタノールアミンとも純度が70%~80%程度であり、90%以上の高純度の精製物を得ることはなかなか困難であった。また、上記のように、クロロホルムメタノールを使用する方法は、いかに精製分画しても有害成分が残留してる恐れがあるため、食品や薬品には使用しにくいという問題があった。

また後者のオキアミリン脂質の分取方法は、オキアミ

にはホスファチジルコリンが存在していること、シリカ ゲルカラムを用いてカラムクロマトグラフィーを行いフ ラクション分画をして単離するオキアミリン脂質の分取 方法が記載されている。しかし、当該分取方法でも、オ キアミリン脂質を効率的に且つ高純度で取得することが できなかった。

本発明者は、オキアミが豊富な蛋白質資源として注目されているが、腐敗し易く、水分が多過ぎることから保存と運送にコストがかかり過ぎるとして、その有効な利用法が確立していないこと、また、オキアミにはリン脂質が多く含んでいるが、この有効成分であるリン脂質に着目して付加価値が高く経済性のある高額な医薬品等に利用しようとする技術開発が、いまだなされていないことに気が付いた。

そこで本発明者は、未利用の水産資源であるオキアミを原料として、これから有用なリン脂質を高純度で得ることができれば、オキアミの有効利用法として非常に有益であると考え、その精製法の研究開発を進め、完成したのが本発明である。即ち、本発明は、オキアミを原料として、総脂質を分画し、得られた総脂質から高純度のホスファチジルコリン及びホスファチジルエタノールアミンを高純度化して精製単離することを特徴とする分取方法と、そうして得られた生理活性物質を用いて記憶力改善効果を有する記憶力改善剤として利用する技術である。

「問題点を解決する手段」

本発明は、上記問題点を解決するため、次のような手段を採用したものである。

特許を受けようとする第1発明は、生オキアミを真空 凍結乾燥法により脱水し、得られたオキアミをエタノー ルで総脂質を抽出し、得られた総脂質からエタノールを 除去し、アセトンに溶解し、可溶区分と不溶区分に分画 し、不溶区分を更にアセトンで洗浄し粗リン脂質を得、 この粗リン脂質をエタノールを溶離液として、シリカゲ ルを充填剤として、吸着カラムクロマトグラフィーを用 いて90~95%のホスファチジルコリンとホスファチジル エタノールアミンを分画するようにしたことを特徴とす るオキアミリン脂質の分取方法である。

第一工程:船内急速凍結生オキアミのブロック中には、90%以上が水分であるため、脱水方法が問題になる。そこで本発明では、吸着カラムクロマトグラフィーを用いた分取の前処理として、真空凍結乾燥装置を用いて脱水し乾燥オキアミとする。このとき水分含量が6%以下になるように脱水乾燥するのが望ましい。すると、水溶性蛋白質のエタノール抽出物への混入が抑制できるので、分別成分の純度を高めることができる。

第二工程:第一工程により得られた乾燥オキアミをエタ ノールでホモジナイズして総脂質を抽出する。

第三工程:次に総脂質からエタノールを出来るだけ除去 したうえ、アセトンを溶媒となし、可溶区分と不溶区分

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とに分画する。アセトンを溶媒とした場合には、リン脂質の大部分は不溶区分にあるので、これから溶媒を洗浄すれば、容易に粗リン脂質が得られる。

第四工程:この粗リン脂質をエタノール系溶媒を溶離液となし、吸着カラムクロマトグラフィーを用いてホスファチジルコリンやホスファチジルエタノールアミンに分画し、これからフラクションコレクターにより各リン脂質成分を90%~95%前後の高純度にて単離する。

本発明は、以上のようにして90%~95%前後の高純度のホスファチジルコリンもしくはホスファチジルエタノールアミンなどのオキアミリン脂質を分取する方法である。

この場合、オキアミより単離したホスファチジルコリンもしくはホスファチジルエタノールアミン、またはこれらの誘導体のうち少なくとも一種以上を有効成分として含有して記憶力改善剤を構成するようにしてもよい。この記憶力改善剤は、錠剤、カプセル、顆粒、液状等の形態として、薬品化することができるものである。「作用」

アルツハイマー型痴呆症の場合には、脳内の神経化学 的な変化として、神経伝達物質であるアセチルコリンの 生産が著しく低下していることが知られており、この病 気の予防や治療法として、低下したコリン系の代謝を補 給することにより生理機能を回復せんとすることが行な われている。

特に、人の場合、コリンまたはコリンに解離する天然産出化合物レシチンを経口投与した場合、脳アセチルコ

リンの合成および放出を増進するのに十分な容量の血液 コリン量の増加をもたらすともに、脳脊髄液のコリン量 も増加する生理機能のあること解っている。

従って、オキアミからリン脂質であるホスファチジルコリンをいかに効率良く、しかも安全性を保って抽出するか、それを薬剤として摂取することにより、脳内にアセチルコリンを供給し、これによりアルツハイマー型痴呆症やその他の神経障害の予防と治療を期待するが、特に記憶力改善剤効果を期待しようとするのが本発明である。

#### 「実施例」

以下、本発明を実施例に基ずき詳細に説明する。 <実施例1.>

船内急速凍結生オキアミ20Kgを真空乾燥装置を用いて水分含量4%前後になるまで乾燥させて乾燥オキアミ2.2Kgを得た。この原料である乾燥オキアミの脂質組成をイアトロスキャン法で分析した結果は、表1.の通りであった。

次に、こうして得た乾燥オキアミ2kgをエタノール40kgでホモジナイズして総脂質の抽出を行なった。その後、再抽出はエタノール20kgで同様に行なった。

抽出物である総脂質を濃縮して、できるだけエタノールを除去した後、当該総脂質をアセトンに溶解し、可溶区分と不溶区分に分画する。すると大部分のリン脂質は不溶区分に区画される。そこで、当該不溶区分に分画された物質にアセトン洗浄を数回繰り返して、粗リン脂質408gを得た。

表1. 乾燥オキアミの脂質組成

脂質組成	重量 %
ホスファチジルコリン	31.1
ホスファチジルエタノールアミン	7. 5
トリグリセリド	43.2
遊離脂肪酸	6. 5
その他	5.7

次に、前記粗リン脂質400gをエタノールに2000mlに溶 解し、全自動分取型高速液体クロマトグラフィーに装着 した分取カラム(カラム長さ×カラム径:50cm×50mm、 断面積19.6cm³) に粒径10μmの球状シリカゲル(吸着 剤)を充填したものに、1バッチ当たり20mlを自動注入 した。溶離液はエタノール100%を流速30ml/minで流 し、カラム恒温層は40℃で、ピーク検出は紫外部吸収検 出器 (205μm) を用いてモニターしたところ、第1図 に示したクロマトグラムが得られたので、最初のピーク の分画区分をAとなし、2番目の大きなビークの分画区 分をBとしてフラクションコレクターを用いて分取し た。分画区分Bのホスファチジルコリンの純度はイヤト ロスキャン法で分析したところ98%以上であった。1バ ッチのサイクルタイムは30分で、原料溶液を30分毎に自 動充填して100サイクルで約50時間要して、乾燥オキア ミ2kgから高純度ホスファチジルコリンを約239g分取し た。

また、分画区分Aから同様に純度95%以上の高純度のホスファチジルエタノールアミンを約45g分取した。 <実施例2.>

ウエクスラー方式の記憶ないし知能指数試験をしたところ記憶指数123であった記憶喪失にかかっている患者に、オキアミのから第1実施例にて分取した高純度ホスファチジルコリン(純度98%)を6週間に渡って1日3

回食事毎に10gづつ食品に混入して経口投与した。

試験治療前と高純度ホスファチジルコリン摂取終了の6週間後に、患者からコリン測定用血液資料を採取しておき、血漿資料を分離し、凍結し、そしてそのコリン含量について慣用の放射性酵素法により分析した。その結果は、試験治療前採取した血液中の血漿コリン量が13.4±1.2ナノモル/mlであったのに対し、高純度ホスファチジルコリン投与から4時間後に得られた血液中の血漿コリン量が31.3±2.5ナノモル/mlに増加していた(P<0.01)。しかも、高純度ホスファチジルコリン摂取の6週間後には、患者の記憶指数は、142に向上していた。「効果」

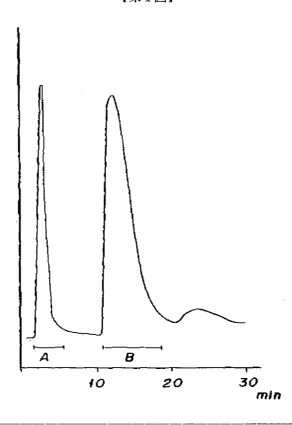
第1請求項に係る保護を受けようとする発明は、未利用の水産資源であるオキアミを原料として、これから有用なホスファチジルコリン及びホスファチジルエタノールアミンを90%以上という高純度で精製単離することができる分取方法である。この分取方法は、精製単離成分が高純度であるというだけでなく、その精製過程において、毒性を持った溶剤などが一切使用されていないので、安全性が高く、薬品などにも安心して利用できる点に特徴がある。

### 【図面の簡単な説明】

第1図は本発明に係るクロマトグラムが得られた組成成分の分画表である。

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フロントページの続き

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(54) Title: LOW VISCOSITY PHOSPHOLIPID COMPOSITIONS

(57) Abstract: The invention relates to processing crustaceans such as krill to oils comprising phospholipids that are Newtonian fluids and/or and have low viscosity, and in particular to the production of oils containing astaxanthin and phospholipids that show Newtonian fluidity and have a low viscosity.

### LOW VISCOSITY PHOSPHOLIPID COMPOSITIONS

#### FIELD OF THE INVENTION

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The invention relates to processing crustaceans such as krill to oils comprising phospholipids that are Newtonian fluids and/or and have low viscosity, and in particular to the production of oils containing astaxanthin and high levels of phospholipids that show Newtonian fluidity and have a low viscosity.

#### 10 BACKGROUND OF THE INVENTION

Krill is a small crustacean which lives in all the major oceans world-wide. For example, it can be found in the Pacific Ocean (Euphausia pacifica), in the Northern Atlantic (Meganyctiphanes norvegica) and in the Southern Ocean off the coast of Antarctica (Euphausia superba). Krill is a key species in the ocean as it is the food source for many animals such as fish, birds, sharks and whales. Krill can be found in large quantities in the ocean and the total biomass of Antarctic krill (E. superba) is estimated to be in the range of 300-500 million metric tons. Antarctic krill feeds on phytoplankton during the short Antarctic summer. During winter, however, its food supply is limited to ice algae, bacteria, marine detritus as well as depleting body protein for energy. Virtue et al., Mar. Biol. 126, 521-527. For this reason, the nutritional values of krill vary during the season and to some extent Phleger et al., Comp. Biochem. Physiol. 131B (2002) 733. In order to annually. accommodate variations in food supply, krill has developed an efficient enzymatic digestive apparatus resulting in a rapid breakdown of the proteins into amino acids. Ellingsen et al., Biochem. J. (1987) 246, 295-305. This autoproteolysis is highly efficient also post mortem, making it a challenge to catch and store the krill in a way that preserves the nutritional quality of the krill. Therefore, in order to prevent the degradation of krill the enzymatic activity is either reduced by storing the krill at low temperatures or the krill is made into a krill meal.

During the krill meal process the krill is cooked so that all the active enzymes are denatured in order to eliminate all enzymatic activity. Krill is rich in phospholipids which act as emulsifiers. Thus it is more difficult to separate water, fat and proteins using mechanical separation methods than it is in a regular fish meal production line. In addition, krill becomes solid, gains weight and loose liquid more easily when mixed with hot water. Eventually this may lead to a gradual build up of coagulated krill proteins in the cooker and a non-continuous

operation due to severe clogging problems. In order to alleviate this, hot steam must be added directly into the cooker. This operation is energy demanding and may also result in a degradation of unstable bioactive components in the krill such as omega-3 fatty acids, phospholipids and astaxanthin. The presence of these compounds, make krill oil an attractive source as a food supplement, a functional food products and a pharmaceutical for the animal and human applications.

Omega-3 fatty acids have recently been shown to have potential effect of preventing cardiovascular disease, cognitive disorders, joint disease and inflammation related diseases such as rheumatoid arthritis. Astaxanthin is a strong antioxidant and may therefore assist in promoting optimal health. Hence, there is a need for a method of processing krill into a krill meal at more gentle conditions which prevents the degradation of these valuable bioactive compounds.

#### SUMMARY OF THE INVENTION

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The invention relates to processing crustaceans such as krill to oils comprising phospholipids that are Newtonian fluids and/or and have low viscosity, and in particular to the production of oils containing astaxanthin and high levels of phospholipids that show Newtonian fluidity and have a low viscosity.

In some embodiments, the present invention provides compositions comprising less than about 150, 100, 10, 5, 2 or 1 mg/kg astaxanthin or from about 0.1 to about 1, 2, 5, 10 or 200 mg/kg astaxanthin, preferably endogenous, naturally occurring astaxanthin, from about 20% to about 50%, 15% to 45%, or 25% to 35% phospholipids on a w/w basis, and about 15% to 60%, about 20% to 50%, or about 25% to 40% protein on a w/w basis, wherein said phospholipids comprise omega-3 fatty acid residues. In some embodiments, the composition comprises a lipid fraction having an omega-3 fatty acid content of from about 5% to about 30%, from 10% to about 30%, or from about 12% to about 18% on a w/w basis. In some embodiments, the phospholipids comprise greater than about 60%, 65%, 80%, 85% or 90% phosphatidylcholine on a w/w basis. In some embodiments, the phospholipids comprise less than about 15%, 10%, 8% or 5% ethanolamine on a w/w basis. In some embodiments, the compositions comprise from about 1% to 10%, preferably 2% to 8%, and most preferably about 2% to 6% alkylacylphosphatidylcholine. In some embodiments, the compositions comprise from about 40% to about 70% triacylglycerol on a w/w basis. In further embodiments, the compositions comprise less than about 1% cholesterol. In some

embodiments, the protein comprises from about 8% to about 14% leucine on a w/w basis and from about 5% to 11% isoleucine on a w/w basis.

In some embodiments, the present invention comprises an aqueous phase and a solid phase, said solid phase comprising from about 20% to about 40% phospholipids on a w/w basis, and about 20% to 50% protein on a w/w basis, wherein said phospholipids comprise from about 10% to about 20% omega-3 fatty acid residues.

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In other embodiments, the present invention provides krill compositions comprising astaxanthin, a protein fraction, and a lipid fraction, wherein said lipid fraction comprises less than about 10%, 5% or 3% phospholipids on a w/w basis. In some embodiments, the phospholipids comprise less than about 15%, 10% or 5% phosphatidylcholine on a w/w basis.

In some embodiments, the present invention provides a krill meal comprising astaxanthin and from about 8 % to about 31 % lipids, preferably from about 8% to about 10 or 18 % lipids, wherein said lipids comprises greater than about 80% neutral lipids on a w/w basis. In some embodiments, the krill meal comprises less than about 15%, 10%, 5%, 3% or 1% phospholipids. In some embodiments, the phospholipids comprise less than about 15%, 10% or 5% phosphatidylcholine on a w/w basis.

In some embodiments, the present invention provides methods of preparing a phospholipid composition from biological material or biomass comprising: mixing said biological material or biomass with water at a suitable temperature to form a solid phase and an aqueous phase comprising phospholipids and proteins; separating said solid phase from said aqueous phase; heating said aqueous phase at a temperature sufficient to form a phospholipid-protein precipitate; and separating said phospholipid-protein precipitate from said aqueous phase. In some embodiments, the present invention provides a phospholipidprotein precipitate obtained by using the foregoing method. In some embodiments, the biological material or biomass is krill. In other embodiments, the biological material or biomass is selected from crabs, shrimp, calanus, plankton, crayfish, eggs or other phospholipid containing biological materials or biomass. In some embodiments, the methods further comprise the step of forming a meal from said solid phase. In some embodiments, the step of forming a meal comprises: heating the solid phase in the presence of water; separating fat and protein in said solid phase; and drying said protein to form a meal. In some embodiments, the processes further comprise the steps of pressing and drying the coagulum to form a coagulum meal. In some embodiments, the drying is by hot air or steam. In some embodiments, the present invention provides a phospholipid-protein precipitate obtained by using the foregoing method. In some embodiments, the present invention provides a

composition comprising a krill solid phase according to the foregoing methods. In some embodiments, the present invention provides a krill meal obtained by the foregoing methods.

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In some embodiments, the present invention provides processes comprising: extracting a first lipid fraction from a krill biomass; extracting a second lipid fraction from a krill biomass; and blending said first lipid fraction and said second lipid fraction to provide a krill lipid composition having a desired composition. In some embodiments, the one or more of the extracting steps are performed in the absence of substantial amounts of organic solvents. In some embodiments, the first lipid fraction is extracted by: mixing krill with water at a suitable temperature to form a solid phase and an aqueous phase comprising phospholipids and protein; separating said solid phase from said aqueous phase; heating said aqueous phase at a temperature sufficient to form a phospholipid-protein precipitate; separating said phospholipid-protein precipitate from said aqueous phase; and separating said phospholipids from said protein. In some embodiments, the second lipid fraction is extracted by: heating the solid phase in the presence of water; and separating fat and protein in said solid phase. In some embodiments, the first lipid fraction comprises a phospholipid fraction comprising greater than about 90% phosphatidylcholine on a w/w basis. In some embodiments, the second lipid fraction comprises greater than about 80% neutral lipids on a w/w basis.

In some embodiments, the present invention provides processes of producing a phospholipid composition from biological material or biomass comprising: mixing said biological material or biomass with water to increase the temperature of said biological material to about 25 to 80 °C, preferably to about 50 to 75 °C, and most preferably to about 60 to 75 °C to form a first solid phase and a first aqueous phase comprising phospholipids and proteins; separating said first solid phase from said first aqueous phase; and separating a protein and phospholipid fraction from said first aqueous phase. In some embodiments, the biomass is heated to the first temperature for at least 3 minutes, preferably from about 3 minutes to 60 minutes, more preferably from about 3 minutes to 20 minutes, and most preferably from about 3 minutes to 10 minutes. The present invention is not limited to the use of any particular biological materials or biomass. In some embodiments, the biological material is a marine biomass. In some preferred embodiments, the biological material or biomass comprises krill crabs, shrimp, calanus, plankton, crayfish, eggs or other phospholipid containing biological materials or biomass. The present invention is not limited to the use of any particular type of krill. In some embodiments, the krill is fresh, while in other embodiments, the krill is frozen. In some embodiments, the krill is of the species Euphausia superba. In some embodiments, the step of separating a protein and phospholipid fraction

from said first aqueous phase comprises heating said first aqueous phase at a temperature sufficient to form a phospholipid-protein coagulate and separating said phospholipid-protein coagulate from said aqueous phase. In some embodiments, the processes utilize a second heating step. In some embodiments, the first aqueous phase is heated to over 80 °C, preferably to about 80 to 120 °C, and most preferably to about 90 to 100 °C. In some embodiments, the krill milk is held at these temperatures for from about 1 minute to about 60 minutes, preferably about 1 minute to about 10 minutes, and most preferably for about 2 minutes to 8 minutes. In some embodiments, the heating is at atmospheric pressure, while in other embodiments, the pressure is greater than atmospheric pressure. In some embodiments, the processes further comprise the step of pressing said phospholipid-protein coagulate to form a coagulate liquid phase and a coagulate press cake. In some embodiments, the processes further comprise drying said coagulate press cake to form a coagulate meal. In some embodiments, the processes further comprise extracting a coagulate oil from said coagulate meal. In some embodiments, the processes further comprise the steps of pressing and drying the coagulum to form a coagulum meal. In some embodiments, the drying is by hot air or steam.

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In some embodiments, the step of separating a protein and phospholipid fraction from said first aqueous phase comprises filtration of said aqueous phase to provide a phospholipidprotein retentate comprising proteins and phospholipids. In some embodiments, filtration is via membrane filtration. In some embodiments, the filtration comprises filtering said aqueous phase through a microfilter with a pore size of from about 50 to 500 nm. In some embodiments, the processes further comprise the step of dewatering said phospholipid-protein retentate to form a retentate liquid phase and a retentate concentrate. In some embodiments, the processes further comprise the step of removing water from said retentate concentrate so that said retentate concentrate is microbially stable. In some embodiments, the processes further comprise the step of extracting a retentate oil from said retentate concentrate. In some embodiments, the processes further comprise the step of heating said first solid phase and then pressing said first solid phase to form a first press cake and a second liquid phase. In some embodiments, the processes further comprise the step of drying said first press cake to provide a first krill meal. In some embodiments, the processes further comprise the steps of heating said second liquid phase and then separating said second liquid phase to provide a first krill oil and stickwater. In some embodiments, the stickwater is evaporated and added to said first press cake, and a meal is formed from said evaporated stickwater and said first press cake to provide a second krill meal. In some embodiments, the second liquid phase is heated to over

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80 °C, preferably to about 80 to 120 °C, and most preferably to about 90 to 100 °C prior to said separation. In some embodiments, the processes further comprise the step of combining the previously described coagulate oil or the retentate oil and the first krill oil to provide a blended oil. In other embodiments, the coagulate oil, retentate oil, or oil pressed from the first solid phase are combined with the coagulate meal or retentate. In further embodiments, the processes of the present invention comprise the further step of supplementing the meals or oils produced as described above with additional proteins, phospholipids, triglycerides, fatty acids, and/or astaxanthin to produce an oil or meal with a desired defined composition. As such, a person of skill in the art will readily recognize that the processes described above serve as a starting point for producing compositions that are further supplemented in subsequent process steps to produce a desired composition, such a composition containing elevated levels of proteins, lipids or astaxanthin. In some embodiments, the present invention provides the lipidprotein composition produced by the foregoing processes. In some embodiments, the present invention provides the coagulate meal produced by the foregoing processes. In some embodiments, the present invention provides the coagulate oil produced by the foregoing processes. In some embodiments, the present invention provides the retentate meal produced by the foregoing processes. In some embodiments, the present invention provides the retentate oil produced by the foregoing processes. In some embodiments, the present invention provides the krill meal produced by the foregoing processes. In some embodiments, the present invention provides a krill oil produced by the foregoing processes. In some embodiments, the present invention provides a blended oil produced by the foregoing processes. In some embodiments, the compositions of the present invention are supplemented with additional proteins, phospholipids, triglycerides, fatty acids, and/or astaxanthin to produce an oil or meal with a desired defined composition. As such, a person of skill in the art will readily recognize that the compositions described above serve as a starting point for producing compositions that are further supplemented in subsequent process steps to produce a desired composition, such a composition containing elevated levels of proteins, lipids or astaxanthin.

In some embodiments, the present invention provides processes comprising: heating a krill biomass to about 25 to 80 °C, preferably to about 50 to 75 °C, and most preferably to about 60 to 75 °C; separating said krill biomass into solid and liquid phases; extracting a first lipid fraction from said solid phase; extracting a second lipid fraction from said liquid phases; and blending said first lipid fraction and said second lipid fraction to provide a krill lipid composition having a desired composition. In some embodiments, the extracting steps are

performed in the absence of substantial amounts of organic solvents. In some embodiments, the first lipid fraction comprises a phospholipid fraction comprising greater than about 90% phosphatidylcholine on a w/w basis. In some embodiments, the second lipid fraction comprises greater than about 80% neutral lipids on a w/w basis.

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In some embodiments, the present invention provides krill compositions comprising from about 0.01 to about 200 mg/kg astaxanthin, from about 45% to about 65% fat w/w, and about 20% to 50% protein w/w, wherein said fat comprises omega-3 fatty acid residues. In some embodiments, the fat has an omega-3 fatty acid content of from about 10% to 30 %, preferably 15% to about 25% on a w/w basis. In some embodiments, the fat comprises from about 20% to about 50% phospholipids w/w, wherein said phospholipids comprise greater than about 65% phosphatidylcholine w/w and from about 1% to about 10% alkylacylphosphatidylcholine. In some embodiments, the phospholipids comprise less than about 10% ethanolamine on a w/w basis. In some embodiments, the fat comprises from about 40% to about 70% triacylglycerol w/w. In some embodiments, the compositions further comprise less than about 1% cholesterol. In some embodiments, the protein comprises from about 8% to about 14% leucine on a w/w basis and from about 5% to 11% isoleucine on a w/w basis.

In some embodiments, the present invention provides krill compositions comprising from about 10% to about 20% protein w/w, about 15% to about 30% fat w/w, and from about 0.01 to about 200 mg/kg astaxanthin. In some embodiments, the fat has an omega-3 fatty acid content of from about 10% to about 30% on a w/w basis. In some embodiments, the fat comprises from about 30% to about 50% phospholipids w/w. In some embodiments, the phospholipids comprise greater than about 65% phosphatidylcholine w/w. In some embodiments, the phospholipids comprise less than about 10% ethanolamine on a w/w basis. In some embodiments, the fat comprises from about 40% to about 70% triacylglycerol w/w. In some embodiments, the compositions comprise less than about 1% cholesterol. In some embodiments, the protein comprises from about 7% to about 13% leucine on a w/w basis and from about 4% to 10% isoleucine on a w/w basis.

In some embodiments, the present invention provides krill meal press cakes comprising from about 65% to about 75% protein w/w (dry matter), from about 10% to about 25% fat w/w (dry matter), and from about 1 to about 200 mg/kg astaxanthin (wet base). In some embodiments, the fat comprises greater than about 30% neutral lipids and greater than about 30% phospholipids on a w/w basis. In some embodiments, the fat comprises from about 50 to about 60% neutral lipids w/w and from about 40% to about 55% polar lipids w/w.

In some embodiments, the protein comprises from about 5% to about 11% leucine w/w and from about 3% to about 7% isoleucine w/w.

In some embodiments, the present invention provides krill meals comprising from about 65% to about 75% protein w/w (dry matter), from about 10% to about 25% fat w/w (dry matter), and from about 1 to about 200 mg/kg astaxanthin (wet base). In some embodiments, the fat comprises greater than about 30% neutral lipids and greater than about 30% phospholipids on a w/w basis. In some embodiments, the fat comprises from about 50 to about 60% neutral lipids w/w and from about 40% to about 55% polar lipids w/w. In some embodiments, the polar lipids comprise greater than about 90% phosphatidyl choline w/w. In some embodiments, the polar lipids comprise less than about 10% phosphatidyl ethanolamine w/w. In some embodiments, the protein comprises from about 5% to about 11% leucine w/w and from about 3% to about 7% isoleucine w/w.

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In some embodiments, the present invention provides krill oil compositions comprising greater than about 1500 mg/kg total esterified astaxanthin, wherein said esterified astaxanthin comprises from about 25 to 35% astaxanthin monoester on a w/w basis and from about 50 to 70% astaxanthin diester on a w/w basis, and greater than about 20 mg/kg free astaxanthin.

In some embodiments, the present invention provides krill compositions comprising from about 3% to about 10% protein w/w, about 8% to about 20% dry matter w/w, and about 4% to about 10% fat w/w. In some embodiments, the fat comprises from about 50% to about 70% triacylglycerol w/w. In some embodiments, the fat comprises from about 30% to about 50% phospholipids w/w. In some embodiments, the phospholipids comprise greater than about 90% phosphatidyl choline w/w. In some embodiments, the fat comprises from about 10% to about 25% n-3 fatty acids. In some embodiments, the fat comprises from about 10% to about 20% EPA and DHA.

In some embodiments, the krill compositions of the present invention are supplemented with additional proteins, phospholipids, triglycerides, fatty acids, and/or astaxanthin to produce an oil or meal with a desired defined composition. As such, a person of skill in the art will readily recognize that the krill compositions described above serve as a starting point for producing compositions that are further supplemented in subsequent process steps to produce a desired composition, such a composition containing elevated levels of proteins, lipids or astaxanthin.

The meal and oil compositions of the present invention described above are characterized in containing low levels, or being substantially free of many volatile compounds

that are commonly found in products derived from marine biomass. In some embodiments, the meals and oils of the present invention are characterized as being substantially free of one or more of the following volatile compounds: acetone, acetic acid, methyl vinyl ketone, 1penten-3-one, n-heptane, 2-ethyl furan, ethyl propionate, 2-methyl-2-pentenal, pyridine, acetamide, toluene, N,N-dimethyl formamide, ethyl butyrate, butyl acetate, 3-methyl-1,4heptadiene, isovaleric acid, methyl pyrazine, ethyl isovalerate, N,N-dimethyl acetamide, 2heptanone, 2-ethyl pyridine, butyrolactone, 2,5-dimethyl pyrazine, ethyl pyrazine, N,Ndimethyl propanamide, benzaldehyde, 2-octanone, β-myrcene, dimethyl trisulfide, trimethyl pyrazine, 1-methyl-2-pyrrolidone. In other embodiments, the meals and oils of the present invention are characterized in containing less than 1000, 100, 10, 1 or 0.1 ppm (alternatively less than 10 mg/100g, preferably less than 1 mg/100 g and most preferably less than 0.1 mg/100 g) of one or more of the following volatile compounds: acetone, acetic acid, methyl vinyl ketone, 1-penten-3-one, n-heptane, 2-ethyl furan, ethyl propionate, 2-methyl-2-pentenal, pyridine, acetamide, toluene, N,N-dimethyl formamide, ethyl butyrate, butyl acetate, 3methyl-1,4-heptadiene, isovaleric acid, methyl pyrazine, ethyl isovalerate, N,N-dimethyl acetamide, 2-heptanone, 2-ethyl pyridine, butyrolactone, 2,5-dimethyl pyrazine, ethyl pyrazine, N,N-dimethyl propanamide, benzaldehyde, 2-octanone, β-myrcene, dimethyl trisulfide, trimethyl pyrazine, 1-methyl-2-pyrrolidone. In further embodiments, the compositions of the present invention are characterized in comprising less than 10 mg/100g, and preferably less than 1mg/100 g (dry weight) of trimethylamine (TMA), trimethylamine oxide (TMAO) and/or lysophosphatidylcholine.

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In some embodiments, the present invention provides systems for processing of marine biomass comprising: a mixer for mixing marine biomass and water to form a mixture having a defined temperature, wherein said mixture has a first solid phase and a first liquid phase. In some embodiments, the water is heated and said defined temperature of said mixture is from about 25 to 80 °C, preferably to about 50 to 75 °C, and most preferably to about 60 to 75 °C. In some embodiments, the systems further comprise a separator in fluid communication with said mixer for separating said first solid phase and said first liquid phase. In some embodiments, the first separator is a filter. In some embodiments, the systems further comprise a first heater unit in fluid communication with said first separator, wherein said first heater unit heats said first liquid phase to a defined temperature. In some embodiments, the defined temperature is about 80°C to about 100°C, preferably 90°C to about 100°C, most preferably 95°C to about 100°C. In some embodiments, the systems further comprise a microfilter in fluid communication with said mixer, wherein said liquid phase is separated into

a retentate phase and a permeate phase by said microfilter. In some embodiments, the systems further comprise a prefilter in line with said microfilter. In some embodiments, the prefilter is a sieve. In some embodiments, the water is heated and said defined temperature of said mixture is from about 25 to 80 °C, preferably to about 50 to 75 °C, and most preferably to about 60 to 75 °C. In some embodiments, the systems further comprise a first separator in fluid communication with said mixer for separating said first solid phase and said first liquid phase. In some embodiments, the first separator is a filter.

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In some embodiments, the present invention provides krill compositions comprising from about 10% to about 20% protein w/w, about 15% to about 30% fat w/w, from about 0.01% to about 200 mg/kg astaxanthin, and less than about 1 mg/100g trimethyl amine, trimethyl amine, volatile nitrogen, or 1g/100g lysophosphatidylcholine or combinations thereof. In some embodiments, the fat has an omega-3 fatty acid content of from about 10% to about 25% on a w/w basis. In some embodiments, the fat comprises from about 35% to about 50% phospholipids w/w. In some embodiments, the phospholipids comprise greater than about 90% phosphatidylcholine w/w. In some embodiments, the phospholipids comprise less than about 10% ethanolamine on a w/w basis. In some embodiments, the fat comprises from about 40% to about 60% triacylglycerol w/w. In some embodiments, the compositions further comprise less than about 1% cholesterol. In some embodiments, the protein comprises from about 7% to about 13% leucine on a w/w basis and from about 4% to 10% isoleucine on a w/w basis.

In some embodiments, the present invention provides processes for processing of marine biomass comprising: providing a marine biomass and a mixer for mixing marine biomass and water to form a mixture having a defined temperature, wherein said mixture comprises a first solid phase and a first liquid phase. In some embodiments, the defined temperature of said mixture is from about 25 to 80 °C, preferably to about 50 to 75 °C, and most preferably to about 60 to 75 °C. In some embodiments, the processes further comprise the steps of separating said liquid phase from said solid phase, and heating said liquid phase to about 80°C to about 100°C, preferably 90°C to about 100°C, most preferably 95°C to about 100°C, to produce a coagulate. In some embodiments, the coagulate comprises proteins and lipids. In some embodiments, the coagulate from residual liquid by filtering.

In some embodiments, the present invention provides systems for processing of marine biomass comprising: a ship; a trawl net towable from said ship, said trawl net configured to catch the marine biomass; and a mixer for mixing said marine biomass and water to form a mixture having a defined temperature, wherein said mixture has a first solid

phase and a first liquid phase. In some embodiments, the marine biomass is krill. In some embodiments, the krill is fresh krill and the trawl and ship are configured to deliver the fresh krill to the mixer. In some embodiments, system comprises a pump to transfer the biomass from the krill to the ship. In some embodiments, the system comprises a microfilter in fluid communication with said mixer, wherein said microfilter separates said first solid phase and said first liquid phase. In some embodiments, the marine biomass is krill. In some embodiments, the krill is fresh krill.

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In some embodiments, the present invention provides an oil extracted from krill comprising from about 40% to about 60% phospholipids by weight of the oil and about 1 to about 1500 mg/l astaxanthin, said oil having Newtonian fluidity at 25°C. In some embodiments, the oil has a viscosity of about 400 to about 1200 microPascals/sec at 25°C. In some embodiments, the oil comprises about 35% to about 55% w/w triglycerides. In some embodiments, the oil further comprises about 10% to about 35% w/w omega-3 fatty acid residues. In some embodiments, the phospholipids comprise about greater than 90% phosphatidyl choline by weight of the phospholipids. In some embodiments, the oil has a viscosity of about 800 to about 1100 microPascals/sec at 25°C. In some embodiments, the krill is *Euphausia superba*. In some embodiments, the present invention provides a capsule containing the previous oil compositions. In some embodiments, the capsule is a gel capsule.

In some embodiments, the present invention provides an oral dosage form comprising an oil extracted from krill comprising from about 40% to about 60% phospholipids by weight of said oil and about 1 to about 1500 mg/l astaxanthin, the oil having a viscosity of about 700 to about 1200 microPascals/sec at 25°C. In some embodiments, the oral dosage form is a gel capsule. In some embodiments, the oral dosage form is a free flowing oil.

In some embodiments, the present invention provides an oil extracted from krill comprising from about 40% to about 60% phospholipids by weight of the oil and about 1 to about 1500 mg/l astaxanthin, said oil having a viscosity of about 400 to about 1200 microPascals/sec at 25°C. In some embodiments, the oil has Newtonian fluidity at 25°C.

In some embodiments, the present invention provides processes for producing a krill oil having Newtonian fluidity comprising: mixing said krill with water to increase the temperature of the krill to about 25 to 80 °C to form a first solid phase and a first aqueous phase comprising said phospholipids and proteins; separating said first solid phase from said first aqueous phase; heating said first aqueous phase to produce a phospholipid and protein concentrate; and extracting an oil from the phospholipid and protein concentrate. In some embodiments, the oil is extracted with ethanol. In some embodiments, the ethanol is removed

by evaporation under reduced pressure. In some embodiments, the present invention provides krill oils produced by the foregoing processes.

In some embodiments, the present invention provides a pharmaceutical composition comprising one or more of the compositions described above in combination with a pharmaceutically acceptable carrier. In some embodiments, the present invention provides a food product comprising one or of the foregoing compositions. In some embodiments, the present invention provides a dietary supplement comprising one or more of the foregoing compositions. In some embodiments, the present invention provides an animal feed comprising one or more of the foregoing compositions.

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#### **DESCRIPTION OF THE FIGURES**

Figure 1 shows an overview of the process of making krill meal with a two stage cooking process.

Figure 2 is a graph of the Permeate flux as function of dry matter of the retentate (%) (°Brix).

Figure 3 is a graph of Average Flux as function of dry matter in retentate.

Figure 4 is a GC of the neutral fraction extracted from krill coagulate.

Figure 5 is a GC analysis of the neutral fraction extracted from krill coagulate.

Figure 6 is a GC of the polar fraction extracted from krill coagulate.

Figure 7 is a GC analysis of the polar fraction extracted from krill coagulate.

Figures 8a, 8b, and 8c provides graphs depicting Newtonian fluidity at 15°C, 25°C, and 35°C, respectively.

#### 25 **DEFINITIONS**

As used herein, "phospholipid" refers to an organic compound having the following general structure:

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wherein R1 is a fatty acid residue, R2 is a fatty acid residue or –OH, and R3 is a –H or nitrogen containing compound choline (HOCH<sub>2</sub>CH<sub>2</sub>N<sup>+</sup>(CH<sub>3</sub>)<sub>3</sub>OH<sup>-</sup>), ethanolamine (HOCH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub>), inositol or serine. R1 and R2 cannot simultaneously be OH. When R3 is an –OH, the compound is a diacylglycerophosphate, while when R3 is a nitrogen-containing compound, the compound is a phosphatide such as lecithin, cephalin, phosphatidyl serine or plasmalogen.

An "ether phospholipid" as used herein refers to a phospholipid having an ether bond at position 1 the glycerol backbone. Examples of ether phospholipids include, but are not limited to, alkylacylphosphatidylcholine (AAPC), lyso-alkylacylphosphatidylcholine (LAAPC), and alkylacylphosphatidylethanolamine (AAPE). A "non-ether phospholipid" is a phospholipid that does not have an ether bond at position 1 of the glycerol backbone.

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, 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).

As used herein, astaxanthin refers to the following chemical structure:

As used herein, astaxanthin esters refer to the fatty acids esterified to OH group in the astaxanthin molecule.

As used herein, the term w/w (weight/weight) refers to the amount of a given substance in a composition on weight basis. For example, a composition comprising 50% w/w phospholipids means that the mass of the phospholipids is 50% of the total mass of the composition (i.e., 50 grams of phospholipids in 100 grams of the composition, such as an oil).

As used herein, the term "fresh krill" refers to krill that is has been harvested less than about 12, 6, 4, 2 or preferably 1 hour prior to processing. "Fresh krill" is characterized in that products made from the fresh krill such as coagulum comprise less than 1 mg/100g TMA, volatile nitrogen or Trimetylamine oxide-N, alone or in combination, and less than 1g/100 g lysophosphatidylcholine.

As used herein, the term "Newtonian fluid" refers to a fluid whose stress versus strain rate curve is linear and passes through the origin. The constant of proportionality is known as the "viscosity." The term "having Newtonian fluidity" is used in reference to a fluid, for example an oil containing phospholipids, that exhibits fluidity properties that are substantially Newtonian fluid-like, i.e., the stress versus strain rate curve is substantially linear and passes approximately through the origin.

#### DETAILED DESCRIPTION OF THE INVENTION

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The invention relates to processing crustaceans such as krill to oils comprising phospholipids that are Newtonian fluids and/or and have low viscosity, and in particular to the production of oils containing astaxanthin and high levels of phospholipids that show Newtonian fluidity and have a low viscosity. In some embodiments, the present invention provides systems and methods for the continuous processing of fresh or frozen krill into useful products, including krill oil, krill meal, and a krill protein/phospholipid coagulum.

Previous processes for treating marine biomasses such as krill have utilized a single high temperature treatment to provide a proteinaceous product. Pat No. SU220741; "Removing fats from the protein paste "Okean". Gulyaev and Bugrova, Konservnaya i Ovoshchesushil'naya Promyshlennost (1976), (4), 37-8; Amino acid composition of protein-coagulate in krill. Nikolaeva, VNIRO (1967), 63 161-4. However, these methods result in a product with a relatively low lipid content. The present invention describes a process in which the marine biomass such as krill is first heated at moderate temperatures to provide an aqueous phase which is subsequently heated at a higher temperature. This process provides a novel protein-lipid composition that has a higher lipid content than previously described

compositions produced from marine biomasses. The compositions of the present invention are further distinguished from other krill oil supplements marketed for human use in that the described compositions are, in some embodiments, provided as solids or powders comprising a combination of krill lipids, including krill phospholipids and krill triglycerides, and krill-derived protein. These solids/powders may preferably be provided in capsules, gel capsules, or as tablets or caplets.

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In some embodiments, the present invention provides solvent-free methods to produce a phospholipid-containing composition from a biomass such as krill, crabs, Calanus, plankton, eggs, crayfish, shrimp and the like without using organic solvents. In some embodiments, the biomass (preferably krill, freshly harvested or frozen) is heated to a temperature in the range of 25 to 80°C, preferably 40 to 75°C, and most preferably 60 to 75°C in order to dissolve/disperse lipids and proteins from the krill into the water phase, which is called krill milk. In some embodiments, the biomass is heated to and held at this first temperature for at least 3 minutes, preferably from about 3 minutes to 60 minutes, more preferably from about 3 minutes to 20 minutes, and most preferably from about 3 minutes to 10 minutes. In some embodiments, the processes then utilize a second heating step. The proteins and phospholipids are precipitated out of the water phase produced from the first heating step by heating the krill milk (after removal of the krill solids) to a temperature of greater than about 80°C, preferably 80 to 120°C, most preferably 95 to 100°C. In some embodiments, the krill milk is held at these temperatures for from about 1 minute to about 60 minutes, preferably about 1 minute to about 10 minutes, and most preferably for about 2 minutes to 8 minutes. The water phase may be heated at atmospheric pressure, or the water phase may be heated in a closed system at an elevated pressure so that the temperature can be increased above 100°C. Accordingly, in some embodiments, the heating is at atmospheric pressure, while in other embodiments, the pressure is greater than atmospheric pressure. The precipitate formed (hereafter called a coagulum) can be isolated and characterized. In some embodiments, the processes further comprise the steps of pressing and drying the coagulum to form a coagulum meal. In some embodiments, the drying is by hot air or steam.

The solid phase (e.g., krill solids) is preferably used to make a krill meal which also has a novel composition. In other embodiments, the krill milk is microfiltrated. The solid phase produced by microfiltration (called the retentate) is similar to that of the coagulum. Data show that the coagulum and retentate are low in cholesterol. In some embodiments, the retentate and coagulum are substantially free of cholesterol. In some embodiments, the retentate and coagulum comprise less than 1% cholesterol, preferably less than 0.1%

cholesterol. This is a novel method to remove at least a portion of the lipids, such as phospholipids, from the krill. Removal of lipids from krill has previously required solvent extraction using liquids such as ethanol or other polar solvents. Solvent extraction is time-consuming and may also result in loss of material and is therefore not wanted. The krill used to separate out the coagulum had been stored frozen for 10 months prior to the experimentation. It is believed that due to the release of proteolytic enzyme activity during a freezing/thawing process, more protein can be expected to be solubilized based on the processing of frozen krill than from fresh krill.

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In some embodiments, the present invention provides systems and processes for processing a marine biomass. In preferred embodiments, the marine biomass is krill, preferably the Antarctic krill Euphausia superba. Other krill species may also be processed using the systems and processes of the present invention. In some embodiments, the krill is processed in a fresh state as defined herein. In some embodiments, the krill is processed on board a ship as described below within 12, 10, 8, 6, 4, or preferably 2 hours of catching the krill. In some embodiments, the krill is processed on board a ship within 1 or preferably 0.5 hours of catching the krill. In some embodiments, the ship tows a trawl that is configured to catch krill. The krill is then transferred from the trawl to the ship and processed. In some embodiments, the trawl comprises a pump system to pump the freshly caught krill from the trawl to the ship so that the krill can be processed in a fresh state. In preferred embodiments, the pump system comprises a tube that extends below the water the trawl and a pumping action is provided by injecting air into the tube below the waterline so that the krill is continuously drawn or pumped from the trawl, through the tube and on board the ship. Preferred trawling systems with pumps are described in PCT Applications WO 07/108702 and WO 05/004593, incorporated herein by reference.

Some embodiments of the systems and processes of the present invention are shown in Figure 1. As shown in Figure 1, fresh or frozen is krill is mixed in mixer with a sufficient amount of hot water from water heater to increase the temperature of the krill mass to approximately 40 to 75°C, preferably 50 to 75°C, more preferably 60 to 75°C, and most preferably about 60 to 70°C. Many different types of water heaters are useful in the present invention. In some embodiments, the water heater is a steam heated kettle, while in other embodiments, the water heater is a scraped surface heat exchanger. The heated mass is then separated into liquid (krill milk) and krill solid fractions in a filter. In some embodiments, the separation is performed by sieving through a metal sieve. After separation, the krill milk is heated to approximately 90°C to 100°C, preferably to about 95°C to 100°C in a heater. Any

type of suitable water or liquid heater may be used. In preferred embodiments, the heater is a scraped surface heat exchanger. This heating step produced a solid fraction (the coagulum described above) and a liquid fraction. In some preferred embodiments, the separator utilizes a filter as previously described. The present invention is not limited to the use of any particular type of filter. In some embodiments, the filter is a woven filter. In some embodiments, the filter comprises polymeric fibers. The coagulum is introduced into a dewaterer. In some embodiments, the dewaterer is a press such as screw press. Pressing produces a liquid fraction and a press cake. The press cake is dried in a drier to produce coagulum meal.

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The solid krill fraction is introduced into a dewaterer for dewatering. In some embodiments, the dewaterer is a press such as screw press. Pressing produces a press cake and a liquid fraction. The press cake is dried in a drier, such as an air drier or steam drier, to provide krill meal. The liquid fraction is centrifuged to produce a neutral krill oil containing high levels of astaxanthin and stickwater. In preferred embodiments, the stick water is added back into the krill press cake to make a full meal, including the various components of the stick water such as soluble proteins, amino acids, etc.

In alternative embodiments, the krill milk can be treated by microfiltration instead of by heating to form a coagulum. The krill milk is introduced into a microfilter. Microfiltration produces a fraction called a retentate and a liquid permeate. The retentate is concentrated by evaporation under vacuum to stability, water activity <0.5 Aw. Membrane filtration of cooking liquid is preferably performed at about 70 °C with a filter having a pore size of about 10 nm to about 1000nm, more preferably about 50 to about 500 nm, and most preferably about 100 nm. An exemplary filter is the P19-40 100 nm ZrO<sub>2</sub> membrane. In some embodiments, the liquid fraction is prefiltered prior to microfiltration. In preferred embodiments, the prefilter is a roto-fluid sieve (air opening 100 μm).

In yet another embodiment of the invention is a novel and more efficient method of preparing krill meal. By removing the coagulum, the krill meal process is less susceptible to clogging problems and the use of hot steam in the cooker can be avoided. The data disclosed show the coagulum contains a high percentage of phospholipids, hence the separation of the fat in the new krill meal process can be obtained using mechanical methods as in standard fish meal processes. In fact, the separation of fat from the meal is important. Ideally, the krill meal should have a low fat value in order to have satisfactory technical properties. Mechanically separating the fat from the meal will result in a neutral oil rich in astaxanthin. If the neutral oil rich in astaxanthin stays in the meal, the astaxanthin may be degraded during the drying.

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In some embodiments, the present invention provides a krill coagulate and retentate compositions. The compositions are characterized in containing a combination of protein and lipids, especially phospholipids. In preferred embodiments, the compositions are solids or powders (also referred to as a meal). In some embodiments, the compositions comprise from about 20% to about 50% protein w/w, preferably about 30% to 40% protein w/w, and about 40% to 70% lipids w/w, preferably about 50% to 65% lipids w/w, so that the total amount of proteins and lipids in the compositions of from 90 to 100%. In some embodiments, the lipid fraction contains from about 10 g to 30 g omega-3 fatty acid residues per 100 g of lipid, preferably about 15 g to 25 g omega-3 fatty acids residues per 100 g lipids (i.e., from 10 to 30% or preferably from 15 to 25% omega-3 residues expressed w/w as a percentage of total lipids in the composition). In some embodiments, the lipid fraction of the composition comprises from about 25 to 50 g polar lipids per 100 g lipids (25 to 50% w/w expressed as percentage of total lipids), preferably about 30 to 45 g polar lipids per 100 g total lipids (30 to 45% w/w expressed as percentage of total lipids), and about 50 to 70 g nonpolar lipids per 100 g lipids (50 to 70% w/w expressed as percentage of total lipids), so that the total amount of polar and nonpolar lipids is 90 to 100% of the lipid fraction. In some embodiments, the phospholipids comprise greater than about 60% phosphatidylcholine on a w/w basis. In some embodiments, the phospholipids comprise less than about 10% ethanolamine on a w/w basis. In some embodiments, the compositions comprise from about 20% to about 50% triacylglycerol on a w/w basis. In some embodiments, the compositions comprise less than about 1% cholesterol. In some embodiments, the protein fraction comprises from about 8% to about 14% leucine on a w/w basis and from about 5% to 11% isoleucine on a w/w basis. In some embodiments, the compositions comprise less than about 200, 10, 5 or 1 mg/kg naturally occurring or endogenous astaxanthin. In some embodiments, the compositions comprise from about 0.01 to about 200 mg/kg naturally-occurring astaxanthin. It will be recognized that the astaxanthin content of the composition can be increased by adding in astaxanthin from other (exogenous) sources, both natural and non-natural. Likewise, the compositions can be supplemented with exogenous proteins, triglycerides, phospholipids and fatty acids such as omega-3 fatty acids to produce a desired composition.

In yet another embodiment of the invention is a pre-heated krill composition. Non-limiting examples of the pre-heated krill composition is a krill composition comprising lipids with less than 10% or 5% phospholipids, and in particular phosphatidylcholine.

In yet another embodiment of the invention is a novel krill meal product produced from the solid phase left after the first heating step (i.e., the heating step at below 80 C). The

krill meal has good nutritional and technical qualities such as a high protein content, low fat content and has a high flow number. Unexpectedly, the ratios of polar lipids to neutral lipids and EPA to DHA is substantially enhanced as compared to normal krill meal. In some embodiments, the krill meals comprise from about 60% to about 80% protein on a w/w basis, preferably from about 70% to 80% protein on a w/w basis, from about 5% to about 20% fat on a w/w basis, and from about 1 to about 200 mg/kg astaxanthin, preferably from about 50 to about 200 mg/kg astaxanthin. In some embodiments, the fat comprises from about 20 to 40% total neutral lipids and from about 50 to 70% total polar lipids on a w/w basis (total lipids). In some embodiments, the ratio of polar to neutral lipids in the meal is from about 1.5:1 to 3:1, preferably about 1.8:1 to 2.5:1, and most preferably from about 1.8:1 to 2.2:1. In some embodiments, the fat comprises from about 20% to 40% omega-3 fatty acids, preferably about 20% to 30% omega-3 fatty acids. In some embodiments, the ratio of EPA:DHA is from about 1.8:1 to 1:0.9, preferably from about 1.4:1 to 1:1.

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In still other embodiments, the present invention provides oil produced by the processes described above. In some embodiments, the oils comprise greater than about 1800 mg/kg total esterified astaxanthin, wherein said esterified astaxanthin comprises from about 25 to 35% astaxanthin monoester on a w/w basis and from about 50 to 70% astaxanthin diester on a w/w basis, and less than about 40 mg/kg free astaxanthin.

In still further embodiments, the present invention provides oils extracted from the coagulum powder (meal) described above. In some embodiments, the coagulum powder is extracted with ethanol. For example, the coagulum powder may be extracted with a suitable quantity of 96% ethanol for for about one hour at about 15 to 30°C. The mixture of ethanol and coagulum powder is then filtered and the ethanol is removed evaporation, preferably at reduced pressure. In other embodiments, the coagulum powder is extracted by super critical fluid extraction. In some embodiments, the oils comprise from about 40% to about 60% phospholipids by weight of said oil, about 1 to about 1500 mg/l astaxanthin, and have a viscosity of about 700 to about 1200 microPascals/sec (μP/sec) at 25°C. In some embodiments, the oils have oil have a viscosity of about 800 to about 1100 μP/sec at 25°C. In some embodiments, the oils comprise about 35% to about 55% w/w triglycerides. In some embodiments, the oils comprise about 10% to about 35% w/w omega-3 fatty acid residues. In some embodiments, the phospholipids comprise about greater than 90% phosphatidyl choline by weight of the phospholipids. In some embodiments, the krill is *Euphausia superba*. In some embodiments, the oil is provided in a capsule, preferably a gel capsule.

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The compositions of the present invention are highly palatable humans and other animals. In particular the oil and meal compositions of the present invention are characterized as containing low levels of undesirable volatile compounds or being substantially free of many volatile compounds that are commonly found in products derived from marine biomass. In some embodiments, the meals and oils of the present invention are characterized as being substantially free of one or more of the following volatile compounds: acetone, acetic acid, methyl vinyl ketone, 1-penten-3-one, n-heptane, 2-ethyl furan, ethyl propionate, 2-methyl-2pentenal, pyridine, acetamide, toluene, N,N-dimethyl formamide, ethyl butyrate, butyl acetate, 3-methyl-1,4-heptadiene, isovaleric acid, methyl pyrazine, ethyl isovalerate, N,N-dimethyl acetamide, 2-heptanone, 2-ethyl pyridine, butyrolactone, 2,5-dimethyl pyrazine, ethyl pyrazine, N,N-dimethyl propanamide, benzaldehyde, 2-octanone, β-myrcene, dimethyl trisulfide, trimethyl pyrazine, 1-methyl-2-pyrrolidone. In other embodiments, the meals and oils of the present invention are characterized in containing less than 1000, 100, 10, 1 or 0.1 ppm (alternatively less than 10 mg/100g, preferably less than 1 mg/100 g and most preferably less than 0.1 mg/100 g) of one or more of the following volatile compounds: acetone, acetic acid, methyl vinyl ketone, 1-penten-3-one, n-heptane, 2-ethyl furan, ethyl propionate, 2methyl-2-pentenal, pyridine, acetamide, toluene, N,N-dimethyl formamide, ethyl butyrate, butyl acetate, 3-methyl-1,4-heptadiene, isovaleric acid, methyl pyrazine, ethyl isovalerate, N,N-dimethyl acetamide, 2-heptanone, 2-ethyl pyridine, butyrolactone, 2,5-dimethyl pyrazine, ethyl pyrazine, N,N-dimethyl propanamide, benzaldehyde, 2-octanone, β-myrcene, dimethyl trisulfide, trimethyl pyrazine, 1-methyl-2-pyrrolidone. In further embodiments, the compositions of the present invention are characterized in comprising less than 10 mg/100g, and preferably less than 1mg/100 g (dry weight) of trimethylamine (TMA), trimethylamine oxide (TMAO) and/or lysophosphatidylcholine.

In some embodiments, the present invention provides an oral dosage form comprising an oil extracted from krill comprising from about 40% to about 60% phospholipids by weight of said oil and about 1 to about 1500 mg/l astaxanthin, wherein the oil has Newtonian fluidity and/or a viscosity of about 700 to about 1200 microPascals/sec at 25°C. In some embodiments, the compositions of this invention (such as those described in the preceding sections) are contained in acceptable excipients and/or carriers for oral consumption. In some embodiments, the present invention provides a pharmaceutical compositions one or more of the foregoing compositions in combination with a pharmaceutically acceptable carrier. The actual form of the carrier, and thus, the composition itself, is not critical. The carrier may be a liquid, gel, gelcap, capsule, powder, solid tablet (coated caplet or non-coated), tea, or the like.

The composition is preferably in the form of a tablet or capsule and most preferably in the form of a soft gel capsule. Suitable excipient and/or carriers include maltodextrin, calcium carbonate, dicalcium phosphate, tricalcium phosphate, microcrystalline cellulose, dextrose, rice flour, magnesium stearate, stearic acid, croscarmellose sodium, sodium starch glycolate, crospovidone, sucrose, vegetable gums, lactose, methylcellulose, povidone, carboxymethylcellulose, corn starch, and the like (including mixtures thereof). Preferred carriers include calcium carbonate, magnesium stearate, maltodextrin, and mixtures thereof. The various ingredients and the excipient and/or carrier are mixed and formed into the desired form using conventional techniques. The tablet or capsule of the present invention may be coated with an enteric coating that dissolves at a pH of about 6.0 to 7.0. A suitable enteric coating that dissolves in the small intestine but not in the stomach is cellulose acetate phthalate. Further details on techniques for formulation for and administration may be found in the latest edition of Remington's Pharmaceutical Sciences (Maack Publishing Co., Easton, PA).

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The dietary supplement may comprise one or more inert ingredients, especially if it is desirable to limit the number of calories added to the diet by the dietary supplement. For example, the dietary supplement of the present invention may also contain optional ingredients including, for example, herbs, vitamins, minerals, enhancers, colorants, sweeteners, flavorants, inert ingredients, and the like. For example, the dietary supplement of the present invention may contain one or more of the following: ascorbates (ascorbic acid, mineral ascorbate salts, rose hips, acerola, and the like), dehydroepiandosterone (DHEA), Fo-Ti or Ho Shu Wu (herb common to traditional Asian treatments), Cat's Claw (ancient herbal ingredient), green tea (polyphenols), inositol, kelp, dulse, bioflavinoids, maltodextrin, nettles, niacin, niacinamide, rosemary, selenium, silica (silicon dioxide, silica gel, horsetail, shavegrass, and the like), spirulina, zinc, and the like. Such optional ingredients may be either naturally occurring or concentrated forms.

In some embodiments, the dietary supplements further comprise vitamins and minerals including, but not limited to, calcium phosphate or acetate, tribasic; potassium phosphate, dibasic; magnesium sulfate or oxide; salt (sodium chloride); potassium chloride or acetate; ascorbic acid; ferric orthophosphate; niacinamide; zinc sulfate or oxide; calcium pantothenate; copper gluconate; riboflavin; beta-carotene; pyridoxine hydrochloride; thiamin mononitrate; folic acid; biotin; chromium chloride or picolonate; potassium iodide; sodium selenate; sodium molybdate; phylloquinone; vitamin D3; cyanocobalamin; sodium selenite; copper

sulfate; vitamin A; vitamin C; inositol; potassium iodide. Suitable dosages for vitamins and minerals may be obtained, for example, by consulting the U.S. RDA guidelines.

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In further embodiments, the compositions comprise at least one food flavoring such as acetaldehyde (ethanal), acetoin (acetyl methylcarbinol), anethole (parapropenyl anisole), benzaldehyde (benzoic aldehyde), N butyric acid (butanoic acid), d or 1 carvone (carvol), cinnamaldehyde (cinnamic aldehyde), citral (2,6 dimethyloctadien 2,6 al 8, gera nial, neral), decanal (N decylaldehyde, capraldehyde, capric aldehyde, caprinaldehyde, aldehyde C 10), ethyl acetate, ethyl butyrate, 3 methyl 3 phenyl glycidic acid ethyl ester (ethyl methyl phenyl glycidate, strawberry aldehyde, C 16 aldehyde), ethyl vanillin, geraniol (3,7 dimethyl 2,6 and 3,6 octadien 1 ol), geranyl acetate (geraniol acetate), limonene (d, 1, and dl), linalool (linalol, 3,7 dimethyl 1,6 octadien 3 ol), linalyl acetate (bergamol), methyl anthranilate (methyl 2 aminobenzoate), piperonal (3,4 methylenedioxy benzaldehyde, heliotropin), vanillin, alfalfa (Medicago sativa L.), allspice (Pimenta officinalis), ambrette seed (Hibiscus abelmoschus), angelic (Angelica archangelica), Angostura (Galipea officinalis), anise (Pimpinella anisum), star anise (Illicium verum), balm (Melissa officinalis), basil (Ocimum basilicum), bay (Laurus nobilis), calendula (Calendula officinalis), (Anthemis nobilis), capsicum (Capsicum frutescens), caraway (Carum carvi), cardamom (Elettaria cardamomum), cassia, (Cinnamomum cassia), cayenne pepper (Capsicum frutescens), Celery seed (Apium graveolens), chervil (Anthriscus cerefolium), chives (Allium schoenoprasum), coriander (Coriandrum sativum), cumin (Cuminum cyminum), elder flowers (Sambucus canadensis), fennel (Foeniculum vulgare), fenugreek (Trigonella foenum graecum), ginger (Zingiber officinale), horehound (Marrubium vulgare), horseradish (Armoracia lapathifolia), hyssop (Hyssopus officinalis), lavender (Lavandula officinalis), mace (Myristica fragrans), marjoram (Majorana hortensis), mustard (Brassica nigra, Brassica juncea, Brassica hirta), nutmeg (Myristica fragrans), paprika (Capsicum annuum), black pepper (Piper nigrum), peppermint (Mentha piperita), poppy seed (Papayer somniferum), rosemary (Rosmarinus officinalis), saffron (Crocus sativus), sage (Salvia officinalis), savory (Satureia hortensis, Satureia montana), sesame (Sesamum indicum), spearmint (Mentha spicata), tarragon (Artemisia dracunculus), thyme (Thymus vulgaris, Thymus serpyllum), turmeric (Curcuma longa), vanilla (Vanilla planifolia), zedoary (Curcuma zedoaria), sucrose, glucose, saccharin, sorbitol, mannitol, aspartame. Other suitable flavoring are disclosed in such references as Remington's Pharmaceutical Sciences, 18th Edition, Mack Publishing, p. 1288-1300 (1990), and Furia and Pellanca, Fenaroli's Handbook of Flavor Ingredients, The Chemical Rubber Company, Cleveland, Ohio, (1971), known to those skilled in the art.

In other embodiments, the compositions comprise at least one synthetic or natural food coloring (e.g., annatto extract, astaxanthin, beet powder, ultramarine blue, canthaxanthin, caramel, carotenal, beta carotene, carmine, toasted cottonseed flour, ferrous gluconate, ferrous lactate, grape color extract, grape skin extract, iron oxide, fruit juice, vegetable juice, dried algae meal, tagetes meal, carrot oil, corn endosperm oil, paprika, paprika oleoresin, riboflavin, saffron, tumeric, tumeric and oleoresin).

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In still further embodiments, the compositions comprise at least one phytonutrient (e.g., soy isoflavonoids, oligomeric proanthcyanidins, indol 3 carbinol, sulforaphone, fibrous ligands, plant phytosterols, ferulic acid, anthocyanocides, triterpenes, omega 3/6 fatty acids, conjugated fatty acids such as conjugated linoleic acid and conjugated linolenic acid, polyacetylene, quinones, terpenes, cathechins, gallates, and quercitin). Sources of plant phytonutrients include, but are not limited to, soy lecithin, soy isoflavones, brown rice germ, royal jelly, bee propolis, acerola berry juice powder, Japanese green tea, grape seed extract, grape skin extract, carrot juice, bilberry, flaxseed meal, bee pollen, ginkgo biloba, primrose (evening primrose oil), red clover, burdock root, dandelion, parsley, rose hips, milk thistle, ginger, Siberian ginseng, rosemary, curcumin, garlic, lycopene, grapefruit seed extract, spinach, and broccoli.

In still other embodiments, the compositions comprise at least one vitamin (e.g., vitamin A, thiamin (B1), riboflavin (B2), pyridoxine (B6), cyanocobalamin (B12), biotin, ascorbic acid (vitamin C), retinoic acid (vitamin D), vitamin E, folic acid and other folates, vitamin K, niacin, and pantothenic acid). In some embodiments, the particles comprise at least one mineral (e.g., sodium, potassium, magnesium, calcium, phosphorus, chlorine, iron, zinc, manganese, flourine, copper, molybdenum, chromium, selenium, and iodine). In some particularly preferred embodiments, a dosage of a plurality of particles includes vitamins or minerals in the range of the recommended daily allowance (RDA) as specified by the United States Department of Agriculture. In still other embodiments, the particles comprise an amino acid supplement formula in which at least one amino acid is included (e.g., 1-carnitine or tryptophan).

In further embodiments, the present invention provide animal feeds comprising one or more the compositions described in detail above. The animal feeds preferably form a ration for the desired animal and is balanced to meet the animals nutritional needs. The compositions may be used in the formulation of feed or as feed for animals such as fish, including fish fry, poultry, cattle, pigs, sheep, shrimp and the like.

#### **EXAMPLE 1**

Four portions of krill were analysed for dry matter, fat, and protein. Most of the variation in the composition can be expected to be due to variation in the sampling. To include the effect of variation in storage time after thawing, raw material samples were also taken at different times during the working day. The observed variation in raw material input is inherent in all calculations of fat, dry matter and protein distributions based on the reported examples.

Table 1. Composition of krill (g/100 g)

		Fat free					
	Dry matter	Fat	dry matter	Protein			
Krill 1	21,40	7,80	13,60	11,80			
Krill 2	22,13	7,47	14,66	12,96			
Krill 3	23,78	7,44	16,34	14,60			
Krill 4	23,07	7,55	15,52	13,83			
Mean	22,60	7,57	15,03	13,30			
SD	1,04	0,16	1,17	1,20			
RSD	4,6 %	2,2 %	7,8 %	9,0 %			

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#### **EXAMPLE 2**

In this example a novel method for preparing krill meal was investigated. 800 g of preheated water (95-100 °C) and 200g of frozen krill (0 °C) were mixed in a cooker (cooker 1) at a temperature of 75 °C for 6 minutes. Next, the heated krill and the hot water were separated by filtration. The preheated krill was further cooked (cooker 2) by mixing with 300 g hot water (95 °C) in a kitchen pan and kept at 90 °C for 2 minutes before separation over a sieve  $(1,0 \times 1,5 \text{ mm opening})$ . The heated krill was separated from the liquid and transferred to a food mixer and cut for 10 seconds. The disintegrated hot krill was added back to the hot water and centrifuged at  $8600 \times g$  (RCF average) for 10 minutes. The supernatant corresponding to a decanter liquid (Dl) was decanted off. The liquid from cooking step 1 was heated to 95-100 °C to coagulate the extracted protein. The coagulum was separated over a

sieve  $(1.0 \times 1.5 \text{ mm opening})$  and a weight of 40 g was found. Figure 1 shows an overview of the process of making krill meal with a two stage cooking process.

#### **EXAMPLE 3**

The total volatile nitrogen (TVN), trimethylamine (TMA) and trimethylamine oxide (TMAO) content were determined in the four products from the cooking test in example 2 (Table 2). The krill was fresh when frozen, so no TMA was detected in the products. The results show that TMAO is evenly distributed in the water phase during cooking of krill.

Table 2. Distribution of total volatile nitrogen (TVN), trimethylamine (TMA) and trimethylamine oxide (TMAO) in the products from the cooking procedure.

Products from test no.	10		Coagulum	Coagulated	Decanter	Decanter	
	- 0	Krill	from cooker	cooker liquid	solids	liquid	SUM
Weight (wb)	g	200	97,6	711,1	90,3	294,7	
Dry matter	g/100 g	21,4	14,2	1,0	22,2	0,9	
Analytical values							
Total volatile nitrogen	mg N/100 g	8	1,3	1,2	2,3	1	
Trimetylamine-N	mg N/100 g	<1	<1	<1	<1	<1	
Trimetylamine oxid-N	mg N/100 g	107	19,2	13,5	10,4	13,1	
Quantities							
Total volatile nitrogen	mg N	15,0	1,3	8,5	2,1	2,9	14,8
Trimetylamine-N	mg N	-	-	-	-	-	-
Trimetylamine oxid-N	mg N	214	18,7	96,0	9,4	38,6	163
Distribution							
Total volatile nitrogen	% of input	100 %	8 %	57 %	14 %	20 %	99 %
Trimetylamine-N	% of input						
Trimetylamine oxid-N	% of input	100 %	9 %	45 %	4 %	18 %	76 %

In addition, fat, dry matter and astaxanthin were determined in the products (Table 3). It was observed that the major part of the astaxanthin in the krill was found in the press cake (Table 3). Only a minor part is found in the coagulum which contains more than 60 % of the lipid in the krill raw material. The cooking procedure with leaching of a protein-lipid emulsion increases the concentration of astaxanthin in the remaining fat. The results also show that the water free coagulum contains approximately 40% dry matter and 60% fat. The dry matter consist of mostly protein.

Table 3. Distribution of astaxanthin in the products from the cooking procedure.

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Products from test no.	10		Coagulu m	Coagulate d	Decante r	Decante r	
		Krill	from cooker	cooker liquid	solids	liquid	SUM
Weight (wb)	g	200	97,6	711,1	90,3	294,7	
Fat	g/100 g	7,8	10,3	0,1	5,3	0,2	
Fat free dry matter	g/100 g	13,6	3,9	0,9	16,9	0,8	
Analytical values							
Fri Astaxanthin	mg/kg	3	<1	<1	4,5	<1	
Astaxanthin esters	mg/kg	33	1,2	<0,02	59	0,18	
Conc. in lipid							
Fri Astaxanthin	mg/kg lipid	38	-	-	85	-	
Astaxanthin esters	mg/kg lipid	423	12	-	1111	113	
Quantities							
Free Astaxanthin	mg	0,6	-	-	0,4	-	0,4
Astaxanthin esters	mg	6,6	0,1	-	5,3	0,1	6,2
Distribution							
Free Astaxanthin	% of input	100 %	-	-	68 %	-	68 %
Astaxanthin esters	% of input	100 %	2 %	-	81 %	1 %	83 %

The coagulum from the cooking experiment in Example 2 were analysed for lipid classes. The coagulum lipid was dominated by triacylglycerol and phosphatidyl choline with a small quantity of phosphatidyl ethanolamine (Table 4).

Table 4. Distribution of lipid classes in the coagulum from cooking experiments.

		Krill	Coagulum	Coagulum
Experiment			F5	F6
Fat (Bligh & Dyer)	g/100 g sample	7,8	11,8	9,9
Triacylglycerol	g/100 g fat	47	40	50
Diacylglycerol	g/100 g fat	<0,5	1	0,7
Monocylglycerol	g/100 g fat	<1	<1	<1
Free fatty acids	g/100 g fat	12	0,2	0,4
Cholesterol	g/100 g fat	0,3	<0,3	<0,3
Cholesterol esters	g/100 g fat	0,8	< 0,3	<0,3
Phosphatidyl				
ethanolamine	g/100 g fat	5,3	2,3	2,2
Phosphatidyl inositol	g/100 g fat	<1	<1	<1
Phosphatidyl serine	g/100 g fat	<1	<1	<1
Phosphatidyl choline	g/100 g fat	33	43,1	42,3
Lyso-Phosphatidyl				
choline	g/100 g fat	2,4	<1	<1
Total polar lipids	g/100 g fat	41,3	45,5	44,5
Total neutral lipids	g/100 g fat	61,0	41,3	51,2
Sum lipids	g/100 g fat	102,3	86,8	95,7

The proportion of phosphatidyl choline increased from 33 % in krill to 42 - 46 % in the coagulum. The other phospholipids quantified, phosphatidyl ethanolamine and lysophosphatidyl choline, had lower concentrations in the coagulum than in krill. The free fatty acids were almost absent in the coagulum.

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The cooking time in test F5 was 6.75 min, in test F6 it was 4.00 min. The results in Table 4 show no dependence of the distribution of the lipid classes with the cooking time.

The amino acid composition of the coagulum is not much different the amino acid composition in krill. There seems to be a slight increase in the apolar amino acids in the coagulum compared to krill (Table 5). For a protein to have good emulsion properties it is the distribution of amino acids within the protein that is of importance more than the amino acid composition.

Table 5. Amino acids in coagulum from cooking Example 2.

		Coagulum	Coagulum	
		F 10-2	70-100°C	Krill
		mar/apr 2007	24.06.2006	24.06.2006
	- /1.00 -	2007	24.06.2006	24.00.2006
Aspartic acid	g/100 g protein	8,8	10,8	7,8
Tispartie dela	g/100 g	0,0	10,0	,,0
Glutamic acid	protein	10,1	11,6	10,7
	g/100 g			,
Hydroxiproline	protein	< 0,10	< 0,10	< 0,10
	g/100 g			
Serine	protein	4,3	4,6	3,0
	g/100 g			
Glycine	protein	3,7	3,4	4,1
TT*-4* 1*	g/100 g	1.7	1.6	1.6
Histidine	protein	1,7	1,6	1,6
Arginine	g/100 g protein	4,4	4,4	5,7
Arginine	g/100 g	7,7	7,7	5,7
Threonine	protein	5,2	5,6	3,4
	g/100 g	- ,-	-,-	-,.
Alanine	protein	4,7	4,6	4,7
	g/100 g			
Proline	protein	4,2	4,3	3,9
	g/100 g			
Tyrosine	protein	4,3	4,7	2,7
T 7 11	g/100 g			
Valine	protein	6,4	6,6	4,2
Methionine	g/100 g	2.1	2.1	2.4
Memonine	protein	2,1	2,1	2,4
Isoleucine	g/100 g protein	8,0	8,5	4,5
1501cacine	g/100 g	0,0	0,0	•,~
Leucine	protein	10,8	11,6	6,7
	g/100 g			,
Phenylalanine	protein	4,3	4,3	3,6
	g/100 g			
Lysine	protein	7,5	8,2	6,2
	g/100 g	0.75		
Cysteine/Cystine	protein	0,75		
Truntanhan	g/100 g	0.62		
Tryptophan	protein	0,63		

Sum amino acids	91,9	96,9	75,2
Polar amino	47 %	48 %	51 %
Apolar amino			
acids	53 %	52 %	49 %

The fatty acid profile of the coagulum is presented in Table 6. The content of EPA (20:5) is about 12.4 g/100 g extracted fat and the content of DHA (22:6) is about 5.0 g/100 g extracted fat.

# 5 Table 6. Fatty acid content of coagulum

Fatty acid	Unit	Amount
14:0	g/100 extracted fat	11,5
16:0	g/100 extracted fat	19,4
18:0	g/100 extracted fat	1,1
20:0	g/100 extracted fat	<0,1
22:0	g/100 extracted fat	<0,1
16:1 n-7	g/100 extracted fat	7,0
18:1 (n-9) + (n-7) + (n-5)	g/100 extracted fat	18,4
20:1 (n-9) + (n-7)	g/100 extracted fat	1,3
22:1 (n-11) + (n-9) + (n-7)	g/100 extracted fat	0,8
24:1 n-9	g/100 extracted fat	0,1
16:2 n-4	g/100 extracted fat	0,6
16:3 n-4	g/100 extracted fat	0,2
16:4 n-4	g/100 extracted fat	<0,1
18:2 n-6	g/100 extracted fat	1,2
18:3 n-6	g/100 extracted fat	0,1
20:2 n-6	g/100 extracted fat	<0,1
20:3 n-6	g/100 extracted fat	<0,1
20:4 n-6	g/100 extracted fat	0,2
22:4 n-6	g/100 extracted fat	<0,1
18:3 n-3	g/100 extracted fat	0,8
18:4 n-3	g/100 extracted fat	2,5
20:3 n-3	g/100 extracted fat	<0,1
20:4 n-3	g/100 extracted fat	0,4
20:5 n-3	g/100 extracted fat	12,4
21:5 n-3	g/100 extracted fat	0,4
22:5 n-3	g/100 extracted fat	0,3
22:6 n-3	g/100 extracted fat	5,0

# **EXAMPLE 4**

To evaluate the two stage cooking process described above, a laboratory scale test was performed. The tests are described below.

#### Materials and methods

**Raw material.** Frozen krill were obtained by Aker Biomarine and 10 tons were stored at Norway Pelagic, Bergen, and retrieved as required. The krill was packed in plastic bags in cardboard boxes with  $2\times12.5$  kg krill. The boxes with krill were placed in a single layer on the floor of the process plant the day before processing. By the time of processing the krill varied from +3 °C to -3 °C.

### 10 Analytical methods.

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**Protein, Kjeldahl's method:** Nitrogen in the sample is transformed to ammonium by dissolution in concentrated sulfuric acid with cupper as catalyst. The ammonia is liberated in a basic distillation and determined by titration, (ISO 5983:1997(E), Method A 01). Uncertainty: 1 %.

**Protein, Combustion:** Liberation of nitrogen by burning the sample at high temperature in pure oxygen. Detection by thermal conductivity. Percent protein in the sample is calculated by a multiplication of analysed percent nitrogen and a given protein factor, (AOAC Official Method 990.03, 16th ed. 1996, Method A 25).

**Moisture:** Determination of the loss in mass on drying at 103 °C during four hours (ISO 6496 (1999). Method A 04). Uncertainty: 4 %.

**Ash:** Combustion of organic matter at 550 °C. The residue remaining after combustion is defined as the ash content of the sample. (ISO 5984:2002. Method A 02). Uncertainty: 3 %.

Fat, Ethyl acetate extraction: Absorption of moisture in wet sample by sodium sulphate, followed by extraction of fat by ethyl acetate (NS 9402, 1994 (modified calculation). Method A 29).

**Fat, Soxhlet:** Extraction of fat by petroleum ether. Mainly the content of triglycerides is determined, (AOCS Official Method Ba 3-38 Reapproved 1993. Method A 03).

Fat, Bligh and Dyer: Extraction of fat by a mixture of chloroform, methanol, and water in the proportion 1:2:0.8 which build a single phase system. Addition of chloroform and water gives a chloroform phase with the lipids and a water/methanol phase. The lipids are determined in an aliquot of the chloroform phase after evaporation and weighing. The extraction includes both triglycerides and phospholipids. (E.G. Bligh & W.J. Dyer: A rapid method of total lipid extraction and purification. Can.J.Biochem.Physiol. Vol 37 (1959). Metode A 56).

Astaxanthin: Extraction with ethanol and di-chloromethane. Polar products are removed by open column chromatography on silica gel. Isomers are separated on normal phase HPLC on Si 60 column and detection at 470 nm. (Schierle J. & Härdi W. 1994. Determination of stabilized astaxanthin in Carophyll® Pink, premixes and fish feeds. Edition 3. Revised Supplement to: Hoffman P, Keller HE, Schierle J., Schuep W. Analytical methods for vitamins and carotenoids in feed. Basel: Department of Vitamin Research and Development, Roche. Method A 23)

**Moisture in oil:** Determination of actual water content of fats and oils by titration with Karl Fischer reagent, which reacts quantitatively with water, (AOCS Official Method CA 2e-84. Reapproved 1993. Method A 13).

Dry matter in stick water during processing is correlated to refract meter which gives ° Brix. Amino acids were determined as urea derivatives by reversed phase HPLC with fluorescence detection. (Cohen S. A. and Michaud D. P., Synthesis of a Fluorescent Derivatizing Reagent, 6-Aminoquinolyl-N-Hydroxysuccinimidyl Carbamate, and Its Application for the Analysis of Hydrolysate Amino Acids via High-Performance Liquid Chromatography. Analytical Biochemistry 211, 279-287, 1993. Method A42). TVB-N, TMA-N and TMAO-N were determined in a 6% trichloro-acetic acid extract by micro diffusion and titration. (Conway, E. I., and A. Byrne. An absorption apparatus for the micro determination of certain volatile substances. Biochem. J. 27:419-429, 1933, and Larsen, T, SSF rapport nr. A-152, 1991). Fatty acids were determined by esterifying the fatty acids to methyl esters, separate the esters by GLC, and quantify by use of C23:0 fatty acid methyl ester as internal standard.( AOCS Official Method Ce 1b-89, Method A 68). Lipids were separated by HPLC and detected with a Charged Aerosol Detector. Vitamins A, D and E were analysed at AnalyCen, Kambo.

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#### Results and discussion

Raw material of krill. Table 7 gives the results of analysis of the raw material of the krill that was used in the pilot trials. Besides the first trial, the same shipment of krill was used for all trials. The dry matter was about 21-22 %, fat 6 %, protein 13-14 %, salt 1 % pH, total volatile nitrogen (TVN) 18 mgN/100g, trimethylamine (TMA) 4 mg N/100g and trimethylamineoxide (TMAO) 135 mg N/100g. Compared to fish pH, TMAO and salt (C1-) is high for krill.

Table 7. Analysis of raw krill on wet base (wb)

Sample:	Raw material	of krill								
Analysis:	Dry matter	Fat, B&D	Protein	Ash	Salt	pН	TVN	TMA	TMAO	
Date:	g/100 g	g/100 g	g/100 g	g/100 g	g/100 g		mg N/100 g	mg N/100 g	mg N/100 g	Marks
07.08.2007	22,8	7,1	13,5	2,5						Saga Sea 04.07.06 Lot. L1
18.09.2007	21,3	6,0								
04.10.2007	21,6	6,3	13,5							Krillråstoff CO5S
04.10.2007	20,5	5,9	12,8							Krillråstoff AO6S
25.10.2007	22,1	6,0	13,9	2,9	1,1	7,4	20,8	5,8	128,3	Krillråstoff CO5S
25.10.2007	21,3	6,0	13,2	2,7	1,1	7,4	15,0	2,3	140,6	Krillråstoff AO6S
22.11.2007	21,9	5,9				7,8	17,9	3,5	123,7	
Average	21,6	6,2	13,5	2,7	1,1	7,4	17,9	4,0	134,5	

Table 8 gives the analysis of raw krill on dry base. If these figures are multiplied with 0.93 it will give the figures on meal base with 7 % water.

## 5 Table 8 Analysis of raw krill on dry base (db)

Sample:	Raw material	of krill						
Analysis:	Dry matter	Fat, B&D	Protein	Ash	Salt	TVN	TMA	TMAO
Date:	g/100 g	g/100 g	g/100 g	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g
07.08.2007	100	31,1	59,2	11,0				
18.09.2007	100	28,2		0,0				
04.10.2007	100	29,2	62,5	0,0				
04.10.2007	100	28,8	62,4	0,0				
25.10.2007	100	27,1	62,9	13,1	5,0	94,1	26,1	580,5
25.10.2007	100	28,2	62,0	12,7	5,2	70,6	10,9	660,2
22.11.2007	100	26,9				81,7	16,0	564,8
Average	100	28,5	62,5	12,3	5,1	82,4	18,5	620,4

Separation of coagulum and pressing for krill oil. 99 kg krill was processed by adding batches of 20 kg krill to 80 l of water at 95 °C in a steam heated kettle (200 l). The steam on the kettle was closed, and the krill and water were gently mixed manually for 3 minutes, and the mixed temperature became 75 °C (heating step no. 1). The heated krill was separated from the water by sieving. Sieved preheated krill (75°C) was added 20 kg hot water and heated to 85 °C within a minute, (heating step 2). The krill was sieved again and feed into the press. The liquid from step1 (krill milk) was coagulated at 95 °C. All the krill was cooked and the press liquid was separated for oil. From 99 kg krill about 0.5 kg of unpolished krill oil was separated from the press liquid. Tables 9 and 10 provide an analysis of cooked krill after first cooking step on wet base and dry base.

Table 9 Analysis of cooked krill on wet base (wb)

Sample:	Cooked krill							
Analysis:	Dry matter	Fat, B&D	Protein	Ash	pН	TVN	TMA	TMAO
Date:	g/100 g	g/100 g	g/100 g	g/100 g		mg N/100 g	mg N/100 g	mg N/100 g
07.08.2007	20,2	4,7	13,5	2,2				
18.09.2007	19,8	4,6						
25.10.2007	15,2	3,2	10,3	2,0	8,2	10,5	3,5	75,4

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Table 10 Analysis of cooked krill on dry base (db)

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Sample:	Cooked krill						
Analysis:	Dry matter	Fat, B&D	Protein	Ash	TVN	TMA	TMAO
Date:	g/100 g	g/100 g	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g
07.08.2007	100,0	23,3	66,8	10,9			
18.09.2007	100	23,2					
25.10.2007	100	21,1	67,8	13,2	69,3	23,1	496,3

Compared to raw krill (Table 8) there is a reduction in dry matter for cooked krill. The fat content in dry matter is reduced because of the fat in the krill milk which is separated from the cooked krill. The content of protein is increased on dry base, but the ash seems to be at the same level. TMAO in the krill is reduced and is found in the cooking liquid.

Micro filtration. The krill milk (70 °C) from step 1 was coagulated at > 95 °C and separated from the liquid through microfiltration (Soby Miljøfilter). Coagulum was then pressed in a press and dried. Tables 11 and 12 gives analyses of coagulum on wet base and dry base. The dry matter of the coagulum was between 12.8 and 16.7 %. On dry base the fat content about 60 % and TMAO 340 mg N/100 g. The dry matter of the coagulum increased to 34-38 % by pressing. The fat content also increased on dry base (Table 13), but the TMAO was reduced to 145 mg N/100 g. After washing the press cake with 1 part water to 1 part press cake of coagulum and then press again, the TMAO was reduced to 45 mg N/100g on dry base (Table 18).

Table 11 Analysis of coagulum on wet base (wb)

Sample:	Coagulum						
Analysis:	Dry matter	Fat, B&D	Protein	Ash	TVN	TMA	TMAO
Date:	g/100 g	g/100 g	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g
10.10.2007	12,8	7,9					
25.10.2007	14,3	8,3	5,4	1,0	5,9	2,3	48,6
31.10.2007	16,7	9,3	6,2				
Average	14,6	8,5	5,8				

Table 12 Analysis of coagulum on dry base (db)

Sample:	Coagulum						
Analysis:	Dry matter	Fat, B&D	Protein	Ash	TVN	TMA	TMAO
Date:	g/100 g	g/100 g	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g
10.10.2007	100	61,7					
25.10.2007	100	58,0	37,8	7,0	41,0	16,4	340,1
31.10.2007	100	55,7	37,1				
Average	100	58,5	37,4				

Table 13 Analysis of press cake from coagulum on wet base

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Sample:	Press cake of coagulum					Raw krill	Coagulum	Coagulum PK
Analysis:	Dry matter	Fat, B&D	TVN	TMA	TMAO	worked up	perss cake	per kg raw krill
Date:	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g	kg	kg	kg/kg
22.11.2007	38,8	23,6	7,9	4,5	56,1	1000	54,2	0,0542
11.12.2007	33,8	22,5	3,4	0	45,3	500	21,92	0,0438
11.12.2007*	33,6	21,3	0	0	15,3	500	15	0,0300
*) After 1 wash (Pre	ess cake : water = 1:1)							

**Membrane filtration.** Another way to collect the lipids from the krill milk is to separate by membrane filtration. For this to be possible the milk must not coagulate, but be brought to the membrane filter from the sieve (heating step no. 1).

Before the krill milk could enter the membrane filter the milk is pre-filtrated, which was done by the sieve (100  $\mu$ m). The opening of the micro-filter was 100 nm. 80 kg krill was processed by starting by 80 kg water (95 °C) and 20 kg krill into the kettle as described. For the first 2 batches of krill clean water was used (160 kg), but for the last 2 batches permeate from the membrane filter was used instead of water. The membrane filtration was followed with a refract meter calibrated for sugar solution (°Brix). The Brix-value is near the dry matter concentration in the process liquids. The flux value for the filter at about 60 °C was 350 l/m2/h for retentate with 7.8 °Brix (refract meter) and reduced to 290 l/m2/h when the Brix value increased to 9.9 °. The Brix value for the permeate was only 1 ° due to high dilution when the amount to be filtered is small. See Figures 2 and 3. The permeate was golden and transparent.

All permeate was evaporated in a kettle to > 65 ° Brix. Retentate, 2 liter, was evaporated in a laboratory evaporator at 70 °C and 12 mm Hg. At 27.5 °Brix the retentate was still flowing well. As the concentration continued the retentate became more and more viscous, first as a paste and finely to a dry mass. The concentrated retentate (27 °Brix), permeate (> 65 °Brix) and dry retentate were analyzed and the results are given in Table 14 on sample base (% wb) and Table 15 on dry matter base (% db) (sample no 1, 2 and 3). A sample of coagulum was dried as for the retentate (sample no 4).

Table 14 Analysis of concentrate from retentate, permeate and coagulum on wet base (wb)

	Dry matter	Fat (polar+apolar)	Crude Protein	Ash	TVN	TMA	TMAO	Water activity
		Bligh & Dyer						25 °C
Sample	% wb	% wb	% wb	% wb	mg N/100g wb	mg N/100g wb	mg N/100g wb	aw
No. 1 Concentrate of retentat	26,0	16,3	9,5	1,6	5,7	<1	99	0,978
No. 2 Consentrate of permeat	72,7	1,0	51,1	24,7	138	110	1 157	0,385
No. 3 Vakuum dried retentate	64,9	39,3	24	4,1	12,8	29,4	196	0,875
No. 4 Vakuum died coagulum	60,3	37,1	20,9	4,4	52,9	28,1	216	0,912

Table 15 Analysis of concentrate from retentate, permeate and coagulum on dry matter base (db)

	Dry matter	Fat (polar+apolar)	Crude Protein	Ash	TVN	TMA	TMAO
		Bligh & Dyer					
Sample	% db	% db	% db	% db	mg N/100g db	mg N/100g db	mg N/100g db
No. 1 Concentrate of retentat	100,0	62,7	36,5	6,2	21,9	<1	382
No. 2 Consentrate of permeat	100,0	1,4	70,3	34,0	190	152	1 592
No. 3 Vakuum dried retentate	100,0	60,6	37,0	6,3	19,7	45,3	302
No. 4 Vakuum died coagulum	100,0	61,5	34,7	7,3	87,7	46,6	358

These results indicate that micro filtration of krill milk was promising and is an alternative to coagulate the krill milk. The protein portion was high in taurine. The content of fat, protein, ash and TMAO were almost similar between retentate and coagulum. Permeate can be concentrated to 70 % dry matter and will have a water activity below 0.4 at 25 °C which means that it can be stored at ambient temperature.

Press cake and press liquid. Tables 16 and 17 provide an analysis of press cake on wet and dry base from the different trials. The average amount of press cake per kg raw krill was found to be 0.23 kg. The dry matter of the press cake was between 44 and 48 %. The fat content in dry matter was reduced from 21 % before to 15-20 % after pressing. This will give a press cake meal from 14 to 18.5 % fat, about 67 % protein and 7 % moisture. TMAO was reduced from about 500 mg N/100g dry matter in cooked krill to 95mg N/100g dry matter in the press cake.

Table 16 Analysis on wet base (wb) of press cake and calculations

Sample:	Press cake						Raw krill	Press cake	Kg press cake
Analysis:	Dry matter	Fat, B&D	Protein	TVN	TMA	TMAO	worked up		per kg raw krill
Date:	g/100 g	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g	kg	kg	kg/kg
18.09.2007	48,1	8,0					327	90	0,28
04.10.2007	47,9	7,0	34,8						
10.10.2007	44,8	9,3					250	55	0,22
31.10.2007	47,4	7,2	33,8				709	143	0,20
22.11.2007	44,4	8,1		8,4	2,1	42,2	1000	226	0,23
11.12.2007	43,8	7,3		5,6	2,2	46,7	500	117	0,23
Average:	46,1	7,8	34,3	7	2,2	44,5			0,23

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Table 17 Analysis on dry base (db) of press cake

Press cake					
Dry matter	Fat, B&D	Protein	TVN	TMA	TMAO
g/100 g	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g
100	16,6				
100	14,6	72,7			
100	20,8				
100	15,2	71,3			
100	18,2		18,9	4,7	95,0
100	16,7		12,8	5,0	106,6
100	17,0	72,0	15,9	4,9	100,8

Oil was produced from the krill solids by centrifugation. Table 18. The oil was almost free for water and the content of astaxanthin was quite high (1.8 g/kg).

Table 18 Analysis of krill oil

		Date:	Date:
Tricanter oil (krill oil)		31.10.2007	22.11.2007
Astaxanthin, Free	mg/kg	22	29
Trans	mg/kg	12	14
9-cis	mg/kg	2,3	3,2
13-cis	mg/kg	5,4	7,8
Astaxanthin, Esters	mg/kg	1802	1785
Diester	mg/kg	1142	1116
Monoester	mg/kg	660	669
Astaxanthin - total	mg/kg	1824	1814
Water, Karl F.	g/100 g	0,17	0,04
FFA	g/100 g		0,9
Vitamin A	IE/kg		602730
Vitamin D3	IE/kg		<1000
Vitamin E (alfa-tokoferol)	mg/kg		630

10 Table 19 Analysis of press cake from coagulum on dry base

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Sample:	Press cake of coagulum				
Analysis:	Dry matter	Fat, B&D	TVN	TMA	TMAO
Date:	g/100 g	g/100 g	mg N/100 g	mg N/100 g	mg N/100 g
22.11.2007	100	60,8	20,4	11,6	144,6
11.12.2007	100	66,6	10,1	0,0	134,0
11.12.2007*	100	63,4	0,0	0,0	45,5
*) After 1 wash (Pre	ess cake : water = 1:1)				

The yield of coagulum press cake was about 5 % of raw krill. The compositions of coagulum and retentate from micro filtration is compared in Table 20. There was hardly any difference between the products from the two process alternatives. Press cake of coagulum was dried, and Table 21 gives the analysis of the coagulum and final coagulum meal. The proximate composition based on dry matter did not change during drying, and the amino acid composition and fatty acid composition is near identical. There was some loss of

phospholipids during drying. This is most probable caused by oxidation of fatty acids, but other chemical modification of the phospholipids may also be of consequence.

Table 20 Analysis of Retentate from micro filtration and Coagulum

		Retentat 25.10.07	Coagulum 25.10.07
Protein	g/100 g	5,8	5,4
Dry matter	g/100 g	13,5	14,3
Ash	g/100 g	1,1	1,0
Fat (B&D)	g/100 g	7,3	8,3
Н		8,5	
ΓFN	mg N/100 g	5,9	5,9
ГМА	mg N/100 g	2,3	2,3
ГМАО	mg N/100 g	61,0	48,6
ipd classes:			
Friacylglycerol	g/100 g extracted fat	59,0	51
Diacylglycerol	g/100 g extracted fat	1,3	1
Monocylglycerol	g/100 g extracted fat	<1	<1
Free fatty acids	g/100 g extracted fat	3,8	3,2
Cholesterol	g/100 g extracted fat	<0,5	<0,5
Cholesterol esters	g/100 g extracted fat	1,0	0,8
Phosphatidyl ethanolamine	g/100 g extracted fat	1,8	3
Phosphatidyl inositol	g/100 g extracted fat	<1	<1
Phosphatidyl serine	g/100 g extracted fat	<1	<1
Phosphatidyl choline	g/100 g extracted fat	35,0	40
yso-Phosphatidyl choline	g/100 g extracted fat	0,8	1,2
Total polar lipids	g/100 g extracted fat	37,6	44,2
Total neutral lipids	g/100 g extracted fat	67,1	56,0
Sum lipids	g/100 g extracted fat	103,4	100,2
Fatty acid composition:		1	,
14:0	g/100 g extracted fat	10,6	10,4
16:0	g/100 g extracted fat	16,4	16,2
18:0	g/100 g extracted fat	1,1	1,2
20:0	g/100 g extracted fat	0,1	0,1
22:0	g/100 g extracted fat	<0,1	<0,1
16:1 n-7	g/100 g extracted fat	6,3	6,4
18:1 (n-9)+(n-7)+(n-5)	g/100 g extracted fat	15,5	15,4
20:1 (n-9)+(n-7)	g/100 g extracted fat	1,1	1,1
22:1 (n-11)+(n-9)+(n-7)	g/100 g extracted fat	0,6	0,5
24:1 n-9	g/100 g extracted fat	0,1	0,1
16:2 n-4	g/100 g extracted fat	0,5	0,5
16:3 n-4	g/100 g extracted fat	0,2	0,2
18:2 n-6	g/100 g extracted fat	1,4	1,4
18:3 n-6	g/100 g extracted fat	0,2	0,2
20:2 n-6	g/100 g extracted fat	0,1	0,1
20:3 n-6	g/100 g extracted fat	0,1	0,1
20:4 n-6	g/100 g extracted fat	0,3	0,3
22:4 n-6	g/100 g extracted fat	<0,1	<0,1
18:3 n-3	g/100 g extracted fat	0,7	0,7
18:4 n-3	g/100 g extracted fat	1,7	1,7
20:3 n-3	g/100 g extracted fat	<0,1	<0,1
20:4 n-3	g/100 g extracted fat	0,3	0,3
20:5 n-3 (EPA)	g/100 g extracted fat	10,5	10,3
21:5 n-3	g/100 g extracted fat	0,3	0,3
22:5 n-3	g/100 g extracted fat	0,5	0,4
22:6 n-3 (DHA)	g/100 g extracted fat	5,1	5,0
Sum saturated fat acides	g/100 g extracted fat	28,2	27,9
Sum monoene fat acides	g/100 g extracted fat	23,6	23,4
Sum PUFA (n-6) fat acides	g/100 g extracted fat	2,1	25,4
Sum PUFA (n-3) feat acides	g/100 g extracted fat	19,1	18,7
Sum PUFA (II-3) leat acides  Sum PUFA fat acides total	g/100 g extracted fat	21,9	21,4
	g/100 g extracted fat	73,7	72,7
Sum fat acides total	Id/1001 d avtracted for		

Table 21 Analysis of Coagulum press cake and meal dried in a Rotadisc dryer on wet and dry base

		Coagulum	Coagulum	Coagulum	Coagulum
		press cake	meal	press cake	meal
		22.11.2007	22.11.2007	22.11.2007	22.11.2007
Analysis:	-/100 -	wb	wb	db	db
Protein	g/100 g	14,6	35,3	37,6	37,4
Moisture	g/100 g	61,2	5,7	0,0	0,0
Fat B&D	g/100 g	23,6	55,1	60,8	58,4
Ash	g/100 g		5,9	44.0	6,3
TMA	mg N/100 g	4,5	7	11,6	7
TMAO	mg N/100 g	56,1	140	144,6	148
Fatty acid composition:					
14:0	g/100 g extracted fat	10,4	10,4		
16:0	g/100 g extracted fat	17	17		
18:0	g/100 g extracted fat	1,2	1,2		
20:0	g/100 g extracted fat	0,1	0,1		
22:0	g/100 g extracted fat	0,1	0,1		
16:1 n-7	g/100 g extracted fat	6,4	6,4		
18:1 (n-9)+(n-7)+(n-5)	g/100 g extracted fat	15,2	15,3		
20:1 (n-9)+(n-7)	g/100 g extracted fat	1,1	1,1		
22:1 (n-11)+(n-9)+(n-7)	g/100 g extracted fat	0,5	0,6		
24:1 n-9	g/100 g extracted fat	0,1	0,1		
16:2 n-4	g/100 g extracted fat	0,5	0,5		
16:3 n-4	g/100 g extracted fat	0,2	0,2		
18:2 n-6	g/100 g extracted fat	1,5	1,4		
18:3 n-6	g/100 g extracted fat	0,2	0,2		l
20:2 n-6	g/100 g extracted fat	0,1	0,1		
20:3 n-6	g/100 g extracted fat	<0,1	<0,1	<b> </b>	<del>                                     </del>
20:4 n-6			0.3		<b>-</b>
20:4 n-6 22:4 n-6	g/100 g extracted fat	0,3 <0,1	<0,1		<b>.</b>
	g/100 g extracted fat			<b></b>	<u> </u>
18:3 n-3	g/100 g extracted fat	0,7	0,7		ļ
18:4 n-3	g/100 g extracted fat	1,7	1,7		
20:3 n-3	g/100 g extracted fat	<0,1	<0,1		
20:4 n-3	g/100 g extracted fat	0,4	0,4		
20:5 n-3 (EPA)	g/100 g extracted fat	10,9	10,5		
21:5 n-3	g/100 g extracted fat	0,3	0,3		
22:5 n-3	g/100 g extracted fat	0,3	0,3		
22:6 n-3 (DHA)	g/100 g extracted fat	5,3	5,1		
Sum saturated fat acides	g/100 g extracted fat	28,7	28,7		
Sum monoene fat acides	g/100 g extracted fat	23,3	23,3		
Sum PUFA (n-6) fat acides	g/100 g extracted fat	2	2		
Sum PUFA (n-3) feat acides	g/100 g extracted fat	19.7	19		
Sum PUFA fat acides total	g/100 g extracted fat	22,4	21,7		
Sum fat acides total	g/100 g extracted fat	74.4	73,8		
Amino acids:	gried g extracted fat	(-7,-7	70,0		
Aspartic acid	g/100 g protein	10,5	10,5		
		11,2	11,6		<b></b>
Glutamic acid	g/100 g protein	<0,10			<b>.</b>
Hydroxiproline	g/100 g protein	4.3	<0,10		
Serine	g/100 g protein		4,2		
Glycine	g/100 g protein	2	4		
Histidine	g/100 g protein		1,9		
Arginine	g/100 g protein	4,8	4,7		
Threonine	g/100 g protein	4,9	4,9		
Alanine	g/100 g protein	4,8	4,9		
Proline	g/100 g protein	4,2	4,1	ļ	<u> </u>
Tyrosine	g/100 g protein	3,7	3,5		
Valine	g/100 g protein	6	5,9	ļ	
Methionine	g/100 g protein	2,4	2,4	ļ	<u> </u>
Isoleucine	g/100 g protein	6,9	6,7	L	
Leucine	g/100 g protein	9,6	9,4		
Phenylalanine	g/100 g protein	4,5	4,4	l	
Lysine	g/100 g protein	7,7	7,6		
Sum AA	g/100 g protein	91,5	90,7		
Lipid classes:					
Triacylglycerol	g/100 g extracted fat	48	63		
Diacylglycerol	g/100 g extracted fat	1,2	1,3		
Monocylglycerol	g/100 g extracted fat	<1	<1		ŀ
Free fatty acids	g/100 g extracted fat	3,2	3,1	İ	l
Cholesterol	g/100 g extracted fat	1,2	<0,5	i	
Cholesterol esters	g/100 g extracted fat	0,5	0,9		
Phosphatidyl ethanolamine	g/100 g extracted fat	3,1	1,1		
Phosphatidyl inositol	g/100 g extracted fat	<1	<1		
Phosphatidyl serine	g/100 g extracted fat	<1	<1	<del> </del>	<b>-</b>
Phosphatidyl choline	g/100 g extracted fat	38	34	<del>                                     </del>	<b>-</b>
Lyso-Phosphatidyl choline	g/100 g extracted fat	1,2	<1	l	<b>-</b>
Total polar lipids	g/100 g extracted fat	42	34,8	<b> </b>	l
Total polar lipids Total neutral lipids	g/100 g extracted fat			<b> </b>	l
Sum lipids	g/100 g extracted fat	54,6 96,7	67,9 103,6	-	-
Julii lipius	grioo g extracted rat	90,1	100,0	I	

**Krill meal.** Final krill meal was produced. Press cake and press cake with stick water concentrate were dried in a hot air dryer or steam drier. Table 22.

Table 22 Analysis of krill meal from

		Forberg	Forberg	Rota disc.
		Air dried	Air dried	Steam dried
		Press cake	Krill meal	Krill meal
Date: 22.11.2007		meal of krill	with stickwater	with stickwater
Wet base:				
Protein	g/100 g	66,4	63,6	66,3
Moisture	g/100 g	5,9	7,1	3,7
Fat Soxhlet	g/100 g	8,7	10,4	
Fat B&D	g/100 g	15,9	15,6	15,2
Ash	g/100 g	9,8	13,0	13,4
Salt	g/100 g	1,3	4,3	4,4
Water sol. protein	g/100 g prot.	11,1	28,0	27,1
pН		8,6	8,3	
TVN	mg N/100 g	18,8	39,9	38,6
TMA	mg N/100 g	11,1	22,2	29,8
TMAO	mg N/100 g	109,7	442,1	399,5
Dry matter base:				
Protein	g/100 g db	70,6	68,5	
Fat Soxhlet	g/100 g db	9,2	11,2	
Fat B&D	g/100 g db	16,9	16,8	15,8
Ash	g/100 g db	10,4	14,0	
Salt	g/100 g db	1,4	4,6	
TVN	mg N/100 g db	20,0	42,9	40,1
TMA	mg N/100 g db	11,8	23,9	30,9
TMAO	mg N/100 g db	116,6	475,9	414,9
Astaxanthin on wet base:			_	
Astaxanthin, Free	mg/kg	4,6	3,6	<1
Trans	mg/kg	2,5	1,9	<1
9-cis	mg/kg	0,4	0,4	<1
13-cis	mg/kg	1,3	0,9	<1
Astaxanthin, Esters	mg/kg	112,0	100	58,0
Diester	mg/kg	80,0	72,0	50,0
Monoester	mg/kg	32,0	27,0	8,1
Astaxanthin - total	mg/kg	116,6	103,6	58,0
staxanthin on fat base:				
Astaxanthin, Fritt	mg/kg fat	28,9	23,1	<7
Trans	mg/kg fat	15,7	12,2	<7
9-cis	mg/kg fat	2,5	2,6	<7
13-cis	mg/kg fat	8,2	5,8	<7
Astaxanthin, Estere	mg/kg fat	704,4	641,0	381,6
Diester	mg/kg fat	503,1	461,5	328,9
Monoester	mg/kg fat	201,3	173,1	53,3
Astaxanthin - totalt	mg/kg fat	733,3	664,1	381,6
Amino acids:				
Aspartic acid	g/100 g protein	10,6	9,2	9,2
Glutamic acid	g/100 g protein	14,1	12,4	12,3
Hydroxiproline	g/100 g protein	<0,5	<0,5	0,1
Serine	g/100 g protein	4,2	3,7	3,8
Glycine	g/100 g protein	4,4	4,4	4,5
Histidine	g/100 g protein	2,3	1,9	1,9
Arginine	g/100 g protein	6,6	6,0	6,1
Threonine	g/100 g protein	4,3	3,7	4,1
Alanine	g/100 g protein	5,4	4,9	5,3
Proline	g/100 g protein	3,7	4,1	4
Tyrosine	g/100 g protein	4,4	3,1	4,7
Valine	g/100 g protein	5,1	4,4	4,5
Methionine	g/100 g protein	3,2	2,7	2,7
Isoleucine	g/100 g protein	5,3	4,5	4,5
Leucine	g/100 g protein	8,0	6,9	6,9
Phenylalanine	g/100 g protein	4,6	3,9	4
Lysine	g/100 g protein	8,2	7,0	6,6
Sum AA	g/100 g protein	94,4	82,8	85,2
ipide classes:	460			-
Triacylglycerol	g/100 g extracted fat		41,0	63
Diacylglycerol	g/100 g extracted fat		1,7	1,3
Monocylglycerol	g/100 g extracted fat		<1	<1
Free fatty acids	g/100 g extracted fat		8,8	3,1
Cholesterol	g/100 g extracted fat		2,4	<0,5
Cholesterol esters	g/100 g extracted fat		<0,5	0,9
Phosphatidyl ethanolamine	g/100 g extracted fat		3,6	1,1
DESCRIPTION OF A	g/100 g extracted fat		<1	<1
Phosphatidyl inositol			<1	<1
Phosphatidyl serine	g/100 g extracted fat		4	
Phosphatidyl serine Phosphatidyl choline	g/100 g extracted fat		43,0	34
Phosphatidyl serine Phosphatidyl choline Lyso-Phosphatidyl choline	g/100 g extracted fat g/100 g extracted fat		1,1	<1
Phosphatidyl serine Phosphatidyl choline	g/100 g extracted fat			

#### **EXAMPLE 5**

Coagulum meal produced as described in Example 4 was extracted using lab scale SFE. 4,885g of coagulum (freeze dried over night) via a two step extraction: 1) SFE: CO<sub>2</sub>, 500 Bar, 60°C, 70min at a medium flow rate of 1,8ml/min of CO<sub>2</sub>; 2) SFE: CO<sub>2</sub>+15%EtOH, 500 Bar, 60°C, 70min at a medium flow rate of 2,5ml/min of CO<sub>2</sub>+EtOH. The first step extracted 1,576g of extracted neutral fraction (NF). As shown in Figures 4 and 5, the analysis at HPLC show lower than the detectable limit content on PL in the NF. It was extracted about 32.25% of the total material. Table 29 provides the peak areas of the components of the neutral fraction as determined by GC.

Table 29.

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Rel.Area	Peakname	Ret.Time	Area	Height	Rel.Area
%		min	mV*min	mV	%
0,29	n.a.	17,455	0,2864	2,271	0,29
19,49	C14:0	24,073	19,0301	105,696	19,49
21,16	C16:0	32,992	20,6601	88,859	21,16
11,99	C16:1	36,197	11,7032	48,125	11,99
3,5	n.a.	37,28	3,4166	14,344	3,5
1,57	n.a.	43,331	1,5375	6,141	1,57
15,6	n.a.	46,425	15,2285	58,605	15,6
8,81	n.a.	46,873	8,5983	30,65	8,81
0,93	n.a.	50,499	0,9055	3,164	0,93
1,56	n.a.	51,292	1,5216	5,746	1,56
1,67	n.a.	57,312	1,6281	4,78	1,67
2,03	n.a.	60,985	1,98	6,963	2,03
0,02	n.a.	67,761	0,0189	0,116	0,02
0,11	n.a.	68,833	0,1066	0,423	0,11
0,11	n.a.	71,705	0,1028	0,497	0,11
0,08	n.a.	74,053	0,0806	0,398	0,08
	C20:5				
3,92	EPA	74,489	3,826	12,07	3,92
0,11	n.a.	80,519	0,1095	0,48	0,11
	C22:5				
0,08	DPA	85,369	0,0785	0,41	0,08
	C22:6	0= =0=	4.0=40		
1,3	DHA	87,787	1,2719	4,253	1,3

The second step extracted a polar fraction of 1,023g corresponding to 20,95% of the total material. The polar fraction consisted mostly of PL and just less than 1% TG. See Figures 6 and 7. Table 30 provides the peak areas of the components of the polar fraction as determined by GC.

Table 30.

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Rel.Area	Peakname	Ret.Time	Area	Height	Rel.Area
%		min	mV*min	mV	%
2,87	C14:0	24,025	4,8099	28,243	2,87
28,5	C16:0	33,084	47,7079	182,756	28,5
1,82	C16:1	36,155	3,0402	13,166	1,82
1,13	n.a.	43,304	1,8848	8,208	1,13
3,89	n.a.	46,336	6,5129	27,429	3,89
5,46	n.a.	46,852	9,1467	35,825	5,46
2,15	n.a.	51,265	3,6015	14,095	2,15
1,6	n.a.	57,121	2,6735	7,213	1,6
1,72	n.a.	60,944	2,8832	10,686	1,72
2,03	n.a.	68,259	3,3913	8,025	2,03
	C20:5				
30,09	EPA	74,599	50,3768	163,312	30,09
	C22:6				
12,11	DHA	87,832	20,2774	68,714	12,11

The coagulate was dried over night with a weight loss of about 5,53% w/w. The total extracted was about 53,2% of the starting weight of the dried material.

#### **EXAMPLE 6**

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Freshly harvested krill were processed into coagulum on board the ship either 10 minutes or six hours post harvest. The coagulum produced from both the 10 minute post harvest krill and the 6 hour post harvest krill contained less than 1mg/100g volatile nitrogen, less than 1 mg/100 g trimethylamine (TMA), and less than 1g/100g lysophosphatidylcholine. This can be compared to the coagulum produced from frozen krill in Example 4 above, which contained higher levels of volatile nitrogen, and lysophosphatidylcholine. The methods of the invention which utilize freshly harvested krill provide krill products that are characterized in being essentially free of TMA, volatile nitrogen, and lysophosphatidylcholine.

#### **EXAMPLE 7**

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Coagulum meal, 250 g, and krill oil were mixed in a kitchen mixer. The aim was to add 300 – 500 mg astaxanthin/kg coagulum meal. If the oil contains 1500 mg astaxanthin/kg krill oil, at least 200 g oil should be added to one kg of coagulum meal. The flow of the meal was markedly reduced by addition of 10 % oil, and the oil came off on the packaging when the addition of oil was increased to 14 and 20 %. 3.5 kg coagulum from was thawed and milled on a Retsch ZM1 with a 2 mm sieve. The quantity of milled powder was 2.96 kg. The 2.96 kg dried coagulum was added 300 g krill oil in three portions. The knives in the mixer (Stephan UM12) were to far from the bottom to give a good mixing, so the mixture was mixed by hand and mixer intermittently. The astaxanthin content in the final mixture was 40 % lower than calculated. New analyses of astaxanthin were performed on the oil and on the fortified meal. The krill oil had been stored in a cold room at 3 °C for 4 months, and the astaxanthin content in the oil did not change during this storage. A new sample were drawn from the fortified meal after 4 weeks frozen storage, and the astaxanthin content was the same in both samples (Table 31).

Table 31. Composition of steam dried coagulum fortified with 10 % krill oil.

		Analysed	Calculated	New analysis	New analysis
		Meal with oil	Meal with oil	Krill oil	Meal with oil
Dry matter	g/100 g	98.0	99.2		
Protein	g/100 g		33.6		
Fat (B&D)	g/100 g	58.9	60.7		
Ash	g/100 g		5.9		
Water soluble protein	g/100 g protein		15.8		
TFN	mg N/100 g		10		
TMA	mg N/100 g		10		
TMAO	mg N/100 g		113		
Astonomikin Franc		2.5	4.0	27	2.0
Astaxanthin, Free	mg/kg	2.5	4.9	27	2.8
Trans	mg/kg	1.4	2.5	14	1.5
9-cis	mg/kg	0.35	0.6	3.1	0.4
13-cis	mg/kg	0.57	1.2	6.2	0.7
Astaxanthin, Esters	mg/kg	193	338	1805	197
Diester	mg/kg	126	216	1128	127
Monoester	mg/kg	67	122	677	70
Astaxanthin - total	mg/kg	196	343	1832	200
Astaxanthin, Free	mg/kg lipid	4.2	8.1		
Trans	mg/kg lipid	2.4	4.2		
9-cis	mg/kg lipid	0.6	1.0		

13-cis	mg/kg lipid	1.0	2.0
Astaxanthin, Esters	mg/kg lipid	328	556
Diester	mg/kg lipid	214	356
Monoester	mg/kg lipid	114	200
Astaxanthin - total	mg/kg lipid	332	564
Ffa	g/100 g extracted fat		4.4
T-1-11 P-1-1-			20.7
Total polar lipids	g/100 g extracted fat		39.7
Total neutral lipids	g/100 g extracted fat		60.1

The astaxanthin content in fortified coagulum meal is 58 % of the amount in the ingredients. This reduction in astaxanthin takes place during mixing of dried coagulum and krill oil, and indicate that dried coagulum is easily oxidized.

### 5 Example 8

The dried coagulum meal was extracted by supercritical fluid extraction. The extracted oil was analyzed as presented in Tables 32-34.

10 Table 32. Lipid composition

Table 32: Elpia composition	
Phosphatidylcholine	34 g/100 g lipid
Phosphatidylethanolamine	1,3 g/100 g lipid
Triglycerides	48 g/100 g lipid
Cholesterol	n.d.
Free fatty acids	1,0 g/100 g lipid

Table 33. Fatty acid profile

Total saturated fatty acids	26,3 g/100 g lipid
Total omega-3 fatty acids	18,1 g/100 g lipid
Total fatty acids	67,3 g/100 g lipid

Table 34. Miscellaneous properties

Astaxanthin	130 mg/kg
TMAO	87 mg N/100 g

TMA	<1 mg N/100 g
Viscosity at 25°C	61 mPa s

### Example 9

Coagulum meal prepared as described above was administered to two human subjects and absorption of the product was determined by measuring omega-3 fatty acids in total lipids and in phospholipids in plasma. Subject 1 consumed 8g of coagulum in combination with yoghurt, whereas subject 2 consumed 8g of krill oil without yoghurt. The data is presented in Tables 35 (Subject 1) and 36 (Subject 2).

Table 35

Time (h)	C20:5 W3 (EPA)	C22:5 W3 (DPA)	C22:6 W3(DHA)
0	0.117	0.062	0.267
0.5	0.118	0.063	0.270
1	0.113	0.061	0.260
1.5	0.117	0.064	0.272
2	0.116	0.063	0.271
2.5	0.119	0.063	0.271
3	0.123	0.065	0.281
3.5	0.122	0.063	0.275
4	0.123	0.063	0.275
5	0.141	0.065	0.294
6	0.153	0.064	0.286
7	0.154	0.062	0.277
8	0.165	0.063	0.292
10	0.167	0.063	0.291
12	0.163	0.061	0.275
16	0.169	0.062	0.301
24	0.173	0.074	0.323

Table 36

Time (h)	C20:5 W3 (EPA)	C22:5 W3 (DPA)	C22:6 W3(DHA)
0	0.146	0.052	0.260
0.5	0.142	0.052	0.260
1	0.146	0.054	0.268
1.5	0.142	0.053	0.263
2	0.145	0.054	0.267
2.5	0.140	0.053	0.258

3	0.143	0.054	0.264
3.5	0.155	0.056	0.278
4	0.155	0.055	0.277
5	0.179	0.057	0.295
6	0.217	0.057	0.316
7	0.204	0.057	0.304
8	0.211	0.060	0.320
10	0.187	0.057	0.293
12	0.171	0.054	0.272
16	0.166	0.052	0.272
24	0.169	0.061	0.290

These data show that absorption patterns of the coagulum and krill oil are different for the two subjects. The EPA pattern in subject 1 (coagulum) shows that a high EPA level is maintained over a long time despite the fact that coagulum contains less lipid than the krill oil. The coagulum has also enriched the circulating PL pool which could be an indication of absorption/incorporation of krill oil fatty acids in PL form. We have previously observed that krill oil is more efficient in enriching tissue lipid fatty acid profiles than fish oil. These data indicate that coagulum is even more bioeffective than krill oil.

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### Example 10.

The phospholipid content of the retentate was further analyzed by NMR. Table 37 provides the results.

Table 37.

Phospholipid	% (w/w)
Phosphatidylcholine	16,5
Alkylacylphosphatidylcholine	1,7
Lyso-alkylacylphosphatidylcholine	0,28
2-lysophosphatidylcholine	0,52
Phosphatidylethanolamine	0,59
N-acylphosphatidylethanolamine	3,6
Total phospholipid	23,23

### Example 11

This example provides an analysis of the volatile compounds in oil extracted from krill meal and oil extracted from coagulum meal. Table 38. Briefly, oil was extracted by SFE from regular krill meal or meal prepared from coagulum as described above. The oil prepared from coagulum meal had substantially reduced amounts of volatile compounds as compared to the oil prepared from regular krill meal. In particular, 1-penten-3-one was detected in oil prepared from regular krill meal and was absent in oil prepared from coagulum meal. 1-pentene-3-one have previously been identified has a key marker of fishy and metallic off-flavor in fish oil and fish oil enriched food products (Jacobsen et al., J. Agric Food Chem, 2004, 52, 1635-1641).

Table 38.

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	TIC peak area		TIC peak area	
Compound	(Krill oil extracted from krill meal using SFE)	Description	(Krill oil extracted from coagulum using SFE)	Description
dimethyl amine	180403283		22848535	
trimethyl amine	255213688	old fish, strong bad	49040416	old fish
ethanol	394615326	fresh	1426886614	vodka, ethanol
acetone	875959		0	
acetic acid	36136270	weak smell	0	
methyl vinyl ketone	515892		0	
2-butanone	2807131	sweet	23124362	
ethyl acetate	6231705		404501	
1- [dimethylamino]- 2-propanone	23316404		15380603	
1-penten-3-one	5627101	rubbery	0	weak dishcloth

n-heptane	291386		0	
2-ethyl furan	1640866	weak sweet	0	
ethyl propionate	909959		0	
2-methyl-2- pentenal	6996219		0	
pyridine	2085743		0	
acetamide	6169014	pleasant	0	
toluene	4359806		0	
N,N-dimethyl formamide	177968590	garden hose, mint	0	garden hose
ethyl butyrate	1122805		0	
2-ethyl-5-methyl furan	1550476	good, flower	427805	
butyl acetate	306001		856292	
3-methyl-1,4- heptadiene	1617339		0	weak smell, rubber
isovaleric acid	1528541	foot sweat, weak	0	
methyl pyrazine	1335979	peculiar	0	
ethyl isovalerate	1043918	fruity	0	fruity
N,N-dimethyl acetamide	9895351		0	smell, solvent
2-heptanone	7397187	blue cheese	0	
2-ethyl pyridine	317424		0	
butyrolactone	652076	butter, pleasant	0	
2,5-dimethyl pyrazine	2414087		0	
ethyl pyrazine	1909284	metallic	0	soft
N,N-dimethyl propanamide	1160830	unpleasant	0	
benzaldehyde	3134653		0	
2-octanone	2068169	disgusting	0	
β-myrcene	2618870		0	

dimethyl trisulfide	3279406	sewer	0	
n-decane	1851488		331629	
trimethyl pyrazine	4186679	unpleasant	0	
1-methyl-2- pyrrolidone	9577873		0	
eucalyptol	0	peppermint	868411	
asetofenoni	1146348	smell, pleasant	350688	

# Example 12

Krill meal produced by the traditional process (Tables 39-42) was compared with krill meal produced from the solid fraction remaining after removal of krill milk (Tables 43-46).

Table 39

14:0	g/100g total fat	8,3
16:0	g/100g total fat	15,4
18:0	g/100g total fat	1,0
20:0	g/100g total fat	<0,1
22:0	g/100g total fat	<0,1
16:1 n-7	g/100g total fat	4,7
18:1 (n-9)+(n-7)+(n-5)	g/100g total fat	13,5
20:1 (n-9)+(n-7)	g/100g total fat	0,9
22:1 (n-11 )+(n-9)+(n-7)	g/100g total fat	0,6
24:1 n-9	g/100g total fat	0,1
16:2 n-4	g/100g total fat	0,6
16:3 n-4	g/100g total fat	0,3
18:2 n-6	g/100g total fat	1,1
18:3 n-6	g/100g total fat	0,1
20:2 n-6	g/100g total fat	<0,1
20:3 n-6	g/100g total fat	<0,1
20:4 n-6	g/100g total fat	0,3
22:4 n-6	g/100g total fat	<0,1
18:3 n-3	g/100g total fat	0,8
18:4 n-3	g/100g total fat	1,8
20:3 n-3	g/100g total fat	<0,1
20:4 n-3	g/100g total fat	0,4
20:5 n-3	g/100g total fat	11,3
21:5 n-3	g/100g total fat	0,4
22:5 n-3	g/100g total fat	0,3
22:6 n-3	g/100g total fat	6,5

Table 40

* Fat Bligh & Dyer	%	22,8
Sum saturated fatty acids	g/100g total fat	24,7
Sum monounsaturated	g/100g total fat	19,8
fatty acids		
Sum PUFA (n-6)	g/100g total fat	1,6
Sum PUFA (n-3)	g/100g total fat	21,5
Sum PUFA	g/100g total fat	24,0
Sum fatty acids total	g/100g total fat	68,5

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# Table 41

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Triacylglycerol	g/100g total fat
Diacylgyycerol	g/100g total fat
Monoacylglycerol	g/100g total fat
Free fatty acids	g/100g total fat
Cholesterol	g/100g total fat
Cholesterol ester	g/100g total fat
Phosphatidylethanolamine	g/100g total fat
Phosphatidylinositol	g/100g total fat
Phosphatidylserine	g/100g total fat
Phosphatidylcholine	g/100g total fat
Lyso-Phosphatidylcholine	g/100g total fat
Total polar lipids	g/100g total fat
Totale neutral lipids	g/100g total fat
Total sum lipids	g/100g total fat

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# Table 42

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Protein Kjeldahl (N*6,25)	%	60,9
Total	%	92,7
Salt (NaCI)	%	2,9
Trimetylamine-N	Mg N/100 gram	4
Trimethylaminoxide-N	Mg N/100 gram	149
Free Astaxanthin	Mg/kg	<1
Astaxanthin ester	Mg/kg	122

46 1,0 <1

1,6

0,8

37 2,0 36,2 54,0

Table 43

14:0	g/100g total fat	5,0
16:0	g/100g total fat	13,9
18:0	g/100g total fat	0,8
20:0	g/100g total fat	<0,1
22:0	g/100g total fat	<0,1
16:1 n-7	g/100g total fat	3,0
18:1 (n-9)+(n-7)+(n-5)	g/100g total fat	11,4
20:1 (n-9)+(n-7)	g/100g total fat	0,5
22:1 (n-11 )+(n-9)+(n-7)	g/100g total fat	0,4
24:1 n-9	g/100g total fat	0,1
16:2 n-4	g/100g total fat	0,4
16:3 n-4	g/100g total fat	0,2
18:2 n-6	g/100g total fat	1,2
18:3 n-6	g/100g total fat	0,1
20:2 n-6	g/100g total fat	0,1
20:3 n-6	g/100g total fat	0,1
20:4 n-6	g/100g total fat	0,4
22:4 n-6	g/100g total fat	<0,1
18:3 n-3	g/100g total fat	0,7
18:4 n-3	g/100g total fat	1,2
20:3 n-3	g/100g total fat	0,1
20:4 n-3	g/100g total fat	0,3
20:5 n-3	g/100g total fat	13,1
21:5 n-3	g/100g total fat	0,3
22:5 n-3	g/100g total fat	0,3
22:6 n-3	g/100g total fat	10,0

Table 44

* Fat Bligh & Dyer	%	10,2
Sum saturated fatty acids	g/100g total fat	19,7
Sum monounsaturated	g/100g total fat	15 <b>5</b> 3
fatty acids		
Sum PUFA (n-6)	g/100g total fat	1,8
Sum PUFA (n-3)	g/100g total fat	26,1
Sum PUFA	g/100g total fat	28,5
Sum fatty acids	g/100g total fat	63,5
		10

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Table 45

Triacylglycerol	g/100g total fat	155
Diacylgyycerol	g/100g total fat	0,7
Monoacylglycerol	g/100g total fat	<1
Free fatty acids	g/100g total fat	0,9
Cholesterol	g/100g total fat	3,1
Cholesterol ester	g/100g total fat	<0,5
Phosphatidylethanolamine	g/100g total fat	12,8
Phosphatidylinositol	g/100g total fat	<1
Phosphatidylserine	g/100g total fat	<1
Phosphatidylcholine	g/100g total fat	269
Lyso-Phosphatidylcholine	g/100g total fat	1,3
Total polar lipid	g/100g total fat	63,2
Total neutral lipid	g/100g total fat	29,7
Total sum lipid	g/100g total fat	92,9

Table 46

Protein Kjeldahl (N*6,25)	%	73,9
Total	%	96,2
Salt (NaCI)	%	1,9
Trimetylamine-N	Mg N/100 gram	7
Trimethylaminoxide-N	Mg N/100 gram	224
Free Astaxanthin	Mg/kg	2,8
Astaxanthin ester	Mg/kg	89

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### Example 13

Dried coagulum meal or powder produced as described above was extracted with ethanol. The results of the extraction are described in Tables 47-50. The total phospholipids content was 48.6 g/100g. The viscosity was analyzed at 15, 25, and 35° C expressed in micro Pascal per second (µP/sec) and the results were 1850, 960 and 535 µP/sec respectively. This can be compared with oil extracted from conventional krill meal with 46.0 g/100g total phospholipids analyzed at 15, 25, and 35° C, which have viscosities of 9670, 6150 and 2520 μP/sec, respectively.

Table 47

14:0	g/100g oil	8.8
16:0	g/100g oil	16.0
18:0	g/100g oil	1.0
20:0	g/100g oil	<.01
22:0	g/100g oil	<0,1
16:1 n-7	g/100g oil	4.3
18:1 (n-9)+(n-7)+(n-5)	g/100g oil	13.4
20:1 (n-9)+(n-7)	g/100g oil	0.9
22:1 (n-11 )+(n-9)+(n-7)	g/100g oil	0.5
24:1 n-9	g/100g oil	0.1
16:2 n-4	g/100g oil	0.6
16:3 n-4	g/100g oil	0.4
18:2 n-6	g/100g oil	1.2
18:3 n-6	g/100g oil	0.2
20:2 n-6	g/100g oil	0.1
20:3 n-6	g/100g oil	<0.1
20:4 n-6	g/100g oil	0.2
22:4 n-6	g/100g oil	< 0.1
18:3 n-3	g/100g oil	0.9
18:4 n-3	g/100g oil	2.5
20:3 n-3	g/100g oil	0.1
20:4 n-3	g/100g oil	0.4
20:5 n-3	g/100g oil	11.8
21:5 n-3	g/100g oil	0.4
22:5 n-3	g/100g oil	0.3
22:6 n-3	g/100g oil	5.7

Table 48

Sum saturated fatty acids	g/100g oil	25.9
Sum monounsaturated	g/100g oil	19.2
fatty acids		5
Sum PUFA (n-6)	g/100g oil	1.6
Sum PUFA (n-3)	g/100g oil	22.1
Sum PUFA	g/100g oil	24.7
Sum fatty acids	g/100g oil	69.7

Table 49

Triacylglycerol	g/100g oil	1.434
Diacylgyycerol	g/100g oil	1.3
Monoacylglycerol	g/100g oil	<1
Free fatty acids	g/100g oil	2.2
Cholesterol	g/100g oil	0.6
Cholesterol ester	g/100g oil	0.7
Phosphatidylethanolamine	g/100g oil	1.4
Phosphatidylinositol	g/100g oil	<1
Phosphatidylserine	g/100g oil	<1
Phosphatidylcholine	g/100g oil	205 2.1
Lyso-Phosphatidylcholine	g/100g oil	2.1
Total polar lipid	g/100g oil	48.6
Total neutral lipid	g/100g oil	48.3
Total sum lipid	g/100g oil	96.9

Table 50

Trimetylamine-N	Mg N/100 gram	251
Trimethylaminoxide-N	Mg N/100 gram	<del>2</del> 87
Free Astaxanthin	Mg/kg	1.4
Astaxanthin ester	Mg/kg	120
Water	%	0.62
Volatile comp.	%	1.7
Ash	%	4.0
Acid insoluble ash	%	206
Viscosity at 15C	μP/sec	1850
Viscosity at 25C	μP/sec	960
Viscosity at 35C	μP/sec	535

### Example 14

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Dried coagulum meal or powder produced as described above was extracted with ethanol. Samples were heated to 15°C, 25°C, and 35°C in a Grafnt waterbath. The viscosity was determined in a rotational viscosimeter (Rheomat 30, Contraves AG, Zürich). The viscosity in the sample is measured with cup C. The measuring cup with sample and spindle is placed in a water bath and when the temperature of the sample has reached the required temperature the measurement is started. The rotational velocity of the spindle increases from 0 to 340 min<sup>-1</sup> during 1 minute, and then the speed decreases to 0 min<sup>-1</sup> during 1 minute. For calculating viscosity the output signal from the sample is compared to the signal from the viscosity standard. The shear rates at 340 min<sup>-1</sup> are for cup A: 647 s<sup>-1</sup>, cup B: 133 s<sup>-1</sup>, cup C: 74 s<sup>-1</sup>, and cup D: 36 s<sup>-1</sup>. For Newtonian liquids the viscosity for the liquid is reported, for non-Newtonian liquids the viscosity at the highest shear rate is reported together with the shear rate. A series of measurements starts with a viscosity standard. For the samples tested, the C cup and 5000 mPa·s standard were used (Brookfield viscosity standards). When the curve is linear, as for this sample, the viscosity is independent of the shear rate, it has a Newtonian flow. The results are shown in Figures 8 a, b, and c.

All publications and patents mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the described method and system of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are obvious to those skilled in the relevant fields are intended to be within the scope of the following claims.

#### **CLAIMS**

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5 1. An oil extracted from krill comprising from about 40% to about 60% phospholipids by weight of said oil and about 1 to about 1500 mg/l astaxanthin, said oil having Newtonian fluidity at 25°C.

- 2. The oil of Claim 1, said oil having a viscosity of about 400 to about 1200 microPascals/sec at 25°C.
  - 3. The oil of claim 1 or claim 2, further comprising about 35% to about 55% w/w triglycerides.
- 15 4. The oil of any of claims 1 to 3 further comprising about 10% to about 35% w/w omega-3 fatty acid residues.
  - 5. The oil of any of any of claims 1 to 4, wherein said phospholipids comprise about greater than 90% phosphatidyl choline by weight of said phospholipids.

6. The oil of any of any of claims 1 to 5, said oil having a viscosity of about 800 to about 1100 microPascals/sec at 25°C.

- 7. The oil of any of any of claims 1 to 6, wherein said krill is *Euphausia superba*.
- 8. An oil extracted from krill comprising from about 40% to about 60% phospholipids by weight of said oil and about 1 to about 1500 mg/l astaxanthin, said oil having a viscosity of about 400 to about 1200 microPascals/sec at 25°C.
- 30 9. The oil of Claim 8, wherein said oil has Newtonian fluidity at 25°C.
  - 10. A capsule containing the oil of any of claims 1 to 9.
  - 11. The capsule of Claim 10, wherein said capsule is a gel capsule.

- 12. An oral dosage form comprising an oil as claimed in claims 1 to 9.
- 13. The oral dosage form of Claim 12, wherein said oral dosage form is a gel capsule.
- 14. The oral dosage form of Claim 12, wherein said oral dosage form is a free flowing oil.
- 16. A process for producing a krill oil of claims 1 to 9 comprising:
- mixing krill with water to increase the temperature of said krill to about 25 to 80 °C to form a first solid phase and a first aqueous phase comprising said phospholipids and proteins;

separating said first solid phase from said first aqueous phase;

heating said first aqueous phase to produce a phospholipid and protein concentrate;

and

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- extracting an oil from said phospholipid and protein concentrate.
- 17. A krill oil produced by the process of Claim 16.

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FIGURE 1

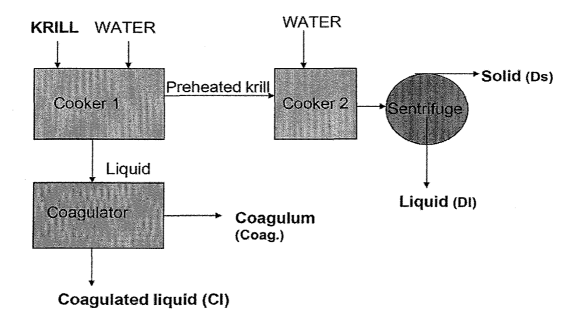


FIGURE 2

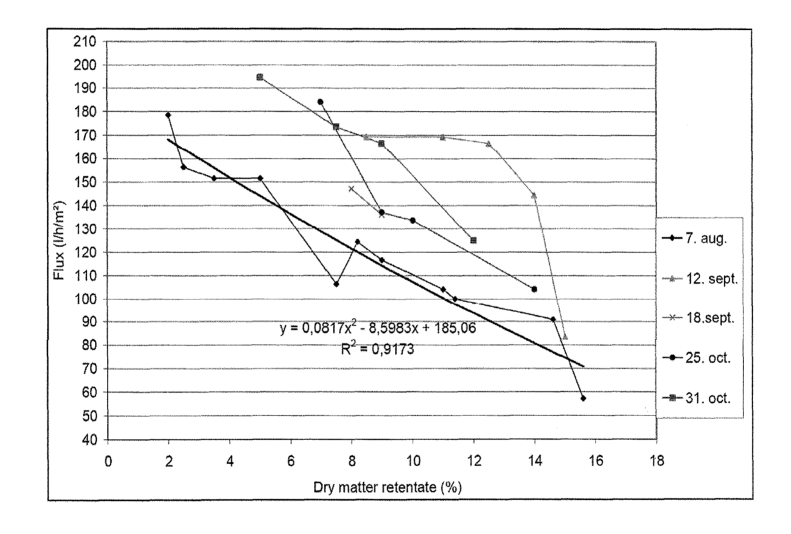


FIGURE 3

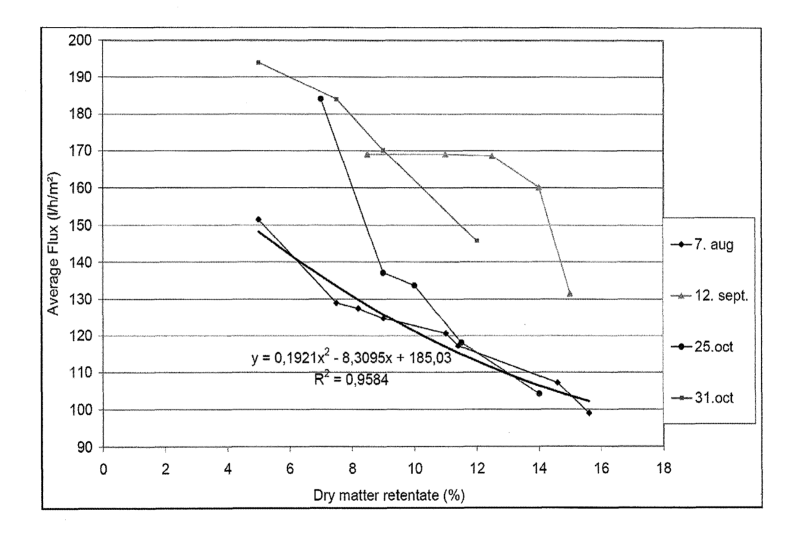
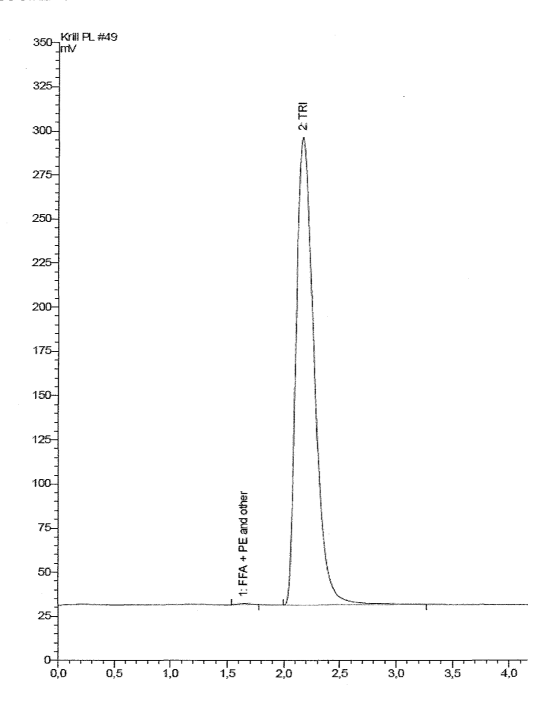
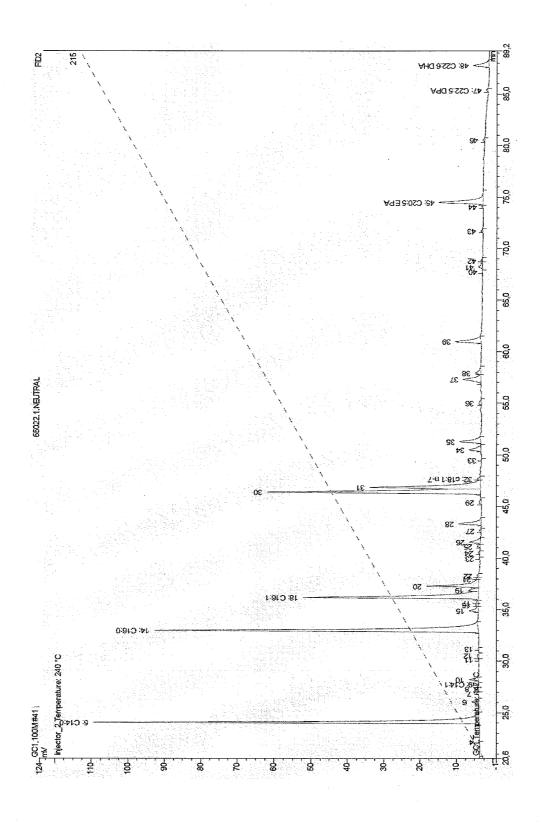


FIGURE 4

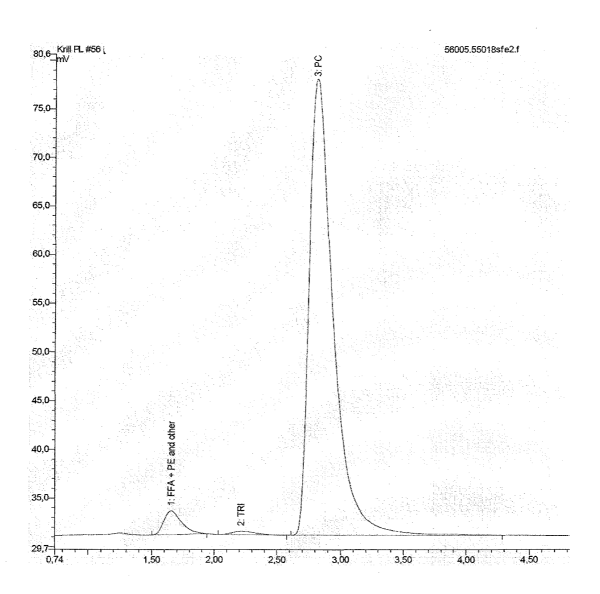




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FIGURE 5

FIGURE 6



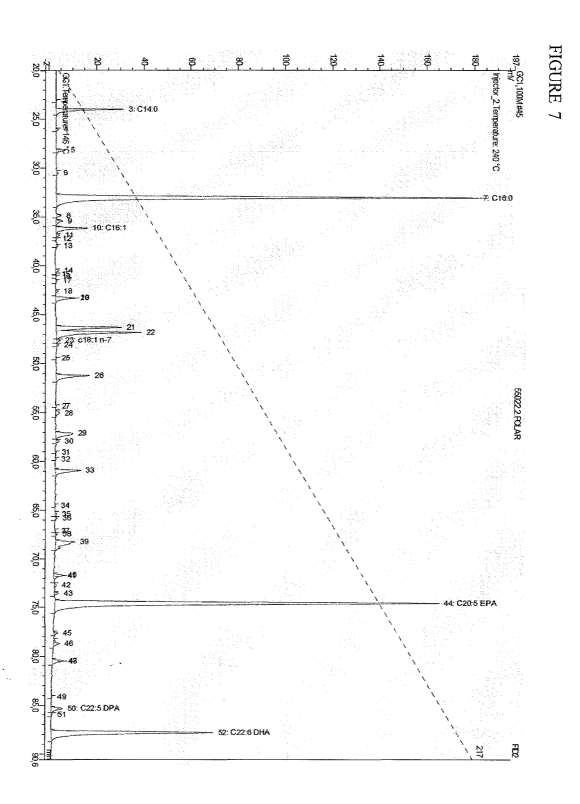
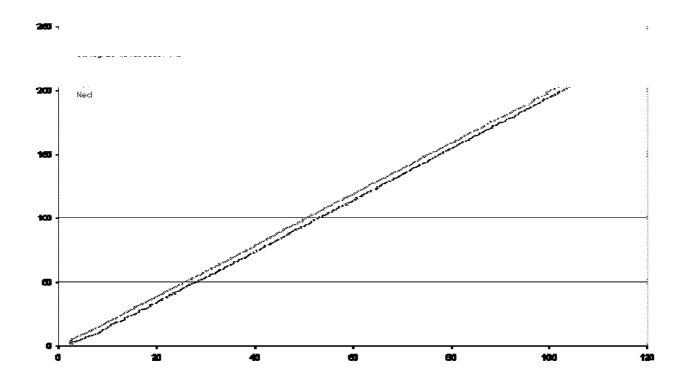
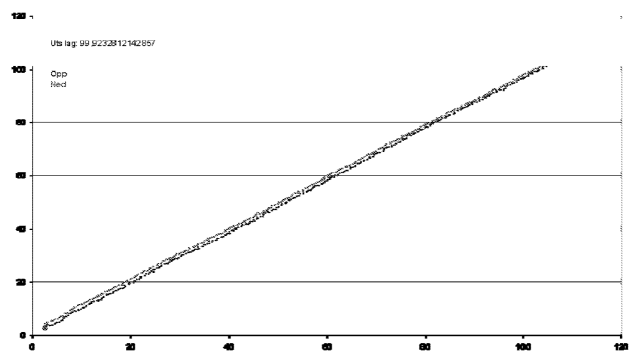


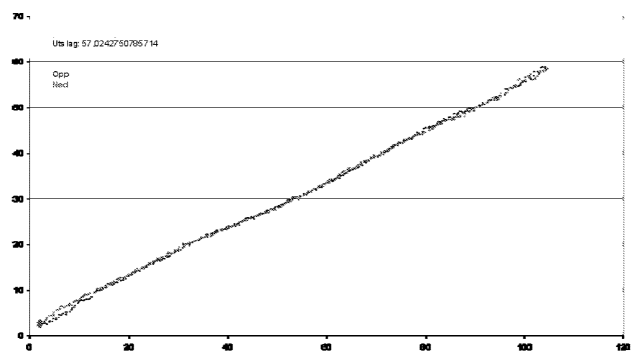
Figure 8a











#### INTERNATIONAL SEARCH REPORT

International application No

PCT/IB2010/000512 A. CLASSIFICATION OF SUBJECT MATTER INV. A23L1/325 A23D9 A23D9/013 A23L1/275 A61K35/56 ADD. According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) A23L A23D A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, FSTA, BIOSIS, EMBASE C. DOCUMENTS CONSIDERED TO BE RELEVANT Category\* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X WO 2008/117062 A1 (AKER BIOMARINE ASA 1-14,17[NO]; GOLDING LOUISE [GB]; BRUHEIM INGE [NO]; GRIIN) 2 October 2008 (2008-10-02) the whole document X WO 2007/080515 A1 (AKER BIOMARINE ASA 16 [NO]; LARSEN PETER MOSE [DK]; FEY STEPHEN JOHN [DK]) 19 July 2007 (2007-07-19) page 3, paragraph 4 - page 5, paragraph 1: claims 1-28; example 1 X,P WO 2009/027692 A2 (AKER BIOMARINE ASA 1 - 17[NO]; GOLDING LOUISE [GB]; OEISTEIN HOESTMARK [NO];) 5 March 2009 (2009-03-05) the whole document -/--Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international "X" document of particular relevance: the claimed invention filing date cannot be considered novel or cannot be considered to "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the "O" document referring to an oral disclosure, use, exhibition or document is combined with one or more other such doc ments, such combination being obvious to a person skilled other means document published prior to the international filing date but "&" document member of the same patent family later than the priority date claimed Date of the actual completion of the international search Date of mailing of the international search report 13/07/2010 24 June 2010 Authorized officer Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Korb, Margit

Fax: (+31-70) 340-3016

# INTERNATIONAL SEARCH REPORT

International application No
PCT/IB2010/000512

ategory*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
A	YAMAGUCHI K ET AL: "SUPERCRITICAL CARBON DIOXIDE EXTRACTION OF OILS FROM ANTARCTIC KRILL"  JOURNAL OF AGRICULTURAL AND FOOD CHEMISTRY, AMERICAN CHEMICAL SOCIETY, US LNKD- DOI:10.1021/JF00071A034, vol. 34, no. 5, 1 January 1986 (1986-01-01), pages 904-907, XP001183110 ISSN: 0021-8561	1-17	
	TAKAICHI S ET AL: "Fatty acids of astaxanthin esteres in krill determined by mild mass spectrometry"  COMPARATIVE BIOCHEMISTRY AND PHYSIOLOGY. PART B, BIOCHEMISTRYAND MOLECULAR BIOLOGY, ELSEVIER, OXFORD, GB LNKD-DOI:10.1016/S1096-4959(03)00209-4, vol. 136, 1 January 2003 (2003-01-01), pages 317-322, XP008110880 ISSN: 1096-4959 the whole document	1-17	

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Information on patent family members

International application No
PCT/IB2010/000512

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[Continued on next page]

(54) Title: METHOD FOR PROCESSING CRUSTACEANS TO PRODUCE LOW FLUORIDE/LOW TRIMETHYL AMINE PRODUCTS THEREOF

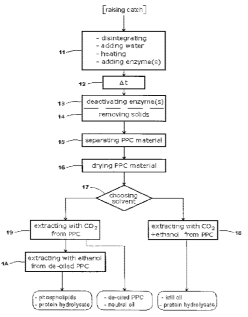


Fig. 1

(57) Abstract: The present invention contemplates the creation of a low fluoride crustacean oil processed from a phospholipid-protein complex (PPC) formed immediately upon a crustacean (i.e., for example, krill) catch. Further, the crustacean oil may also have reduced trimethyl amine and/or trimethyl amino oxide content. The process comprises disintegrating the crustaceans into smaller particles, adding water, heating the result, adding enzyme(s) to hydrolyze the disintegrated material, deactivating the enzyme(s), removing solids from the enzymatically processed material to reduce fluoride content of the material, separating and drying the PPC material. Then, using extraction with supercritical CO<sub>2</sub> or supercritical dimethyl ether, and/or ethanol as solvents, krill oil, inter alia, is separated from the PPC. In the extraction the krill oil can be separated almost wholly from the feed material.

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## Method For Processing Crustaceans To Produce Low Fluoride/Low Trimethyl Amine Products Thereof

#### FIELD OF THE INVENTION

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The invention relates to a method for processing crustaceans (i.e., for example, krill) rich in lipids to produce compositions low in fluoride, trimethyl amine and trimethyl amine oxide comprising phospholipids, proteinaceous nutrients and oil (i.e., for example, neutral lipids and/or triglycerides).

## 10 BACKGROUND OF THE INVENTION

The crustaceans, especially krill, represent a vast resource as biological material. The amount of Antarctic krill ( $Euphausia\ superba$ ), depending on the calculation method and investigation, is roughly 1 to  $2x10^9$  tons and the possible weight of the annual catch is estimated at 5 to  $7x10^6$  tons. These small crustaceans, which live in the cold waters around the Antarctic, are interesting as a source for proteins, lipids such as phospholipids, polyunsaturated fatty acids etc., chitin/chitosan, astaxanthin and other carotenoids, enzymes and other materials.

Several methods for isolating above-mentioned materials have been developed. One problem is that the products may contain unwanted trace material included in the exoskeleton (also called integument or cuticle) of the crustaceans. For example, krill accumulates fluoride in their exoskeleton, thereby increasing the fluoride amount of any produced material either through the inclusion of parts of the exoskeleton or through extraction processes not taking into account the transfer of fluoride to the final material. In this case free fluoride or loosely bound fluoride may diffuse from the exoskeletal material and into the further processed material, making the end product high in fluoride ions and/or fluorinated compounds.

Fluoride is a compound that in high concentrations is detrimental for the health of land-dwelling animals as well as all kind of fish and crustaceans and especially fresh-water fish species, since fluoride atoms have the tendency of entering into the bone structure of such organisms and creating fluorosis, or weakening of the bone structure similar in its effect to osteoporosis, but different since it is the bone structure itself, and not the porosity of the bone that is affected. Skeletal fluorosis is a condition characterized by skeletal abnormalities and joint pain. It is caused by pathological bone formation due to the mitogenic action of fluoride on osteoblasts. In its more severe forms, skeletal fluorosis causes kyphosis, crippling

and invalidism. Secondary neurological complications in the form of myelopathy, with or without radiculopathy, may also occur. High fluoride intake has also been shown to be toxic to the male reproductive system in rat experiments, and in humans high fluoride intake and symptoms of skeletal fluorosis have been associated with decreased serum testosterone levels. Consequently, if krill material is used as a starting material for food or feed products, precautions have to be taken for removing fluoride through the processing steps. However, the diffusion of fluoride and the presence of miniscule particles of the exoskeleton represent a problem that is difficult to overcome when processing krill material in an industrial scale.

Polar lipids such as phospholipids are essential for cell membranes and are also called membrane lipids. For most known animal species the content of polar lipids is nearly constant. However, this does not hold for the Antarctic krill. The phospholipids content varies from 2% up to 10% depending on the season. The high content, e.g. more than 5%, of the phospholipids is in principle good, but means also a problem, because it may result in strong emulsions in industrial processes. The emulsions complicate the separation of the lipid and proteinaceous fractions in the processes, such as hydrolysis.

The krill oil is one the valuable products made from krills. It contains *inter alia* phospholipids, triglycerides and carotenoid astaxanthin while being essentially free of protein, carbohydrates and minerals. Different portions of the krill material are separated from each other by, *inter alia*: i) crushing krill mechanically; ii) pressing them, iii) hydrolysis with heat and enzymes; iv) centrifugal force in rotating devices; and v) solvent extraction.

What is needed in the art are significant improvements to these rather conventional approaches and are described within many embodiments of the present invention (*infra*). For example, a disintegrated raw crustacean material may be separated and/or extracted into various enriched low-fluoride, low trimethyl amine and/or low trimethyl amine oxide crustacean meal and/or oil compositions.

## **SUMMARY**

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The invention relates to a method for processing crustaceans (i.e., for example, krill) rich in lipids to produce compositions low in fluoride, trimethyl amine and trimethyl amine oxide comprising phospholipids, proteinaceous nutrients and oil (i.e., for example, neutral lipids and/or triglycerides).

In one embodiment, the present invention contemplates a crustacean oil composition comprising phospholipids and less than approximately 0.5 ppm fluoride. In one embodiment, the crustacean oil composition further comprises less than approximately 0.001% (w/w)

trimethyl amine. In one embodiment, the crustacean oil composition further comprises less than approximately 0.02% (w/w) trimethyl amine oxide. In one embodiment, the phospholipids are between approximately 39-52 wt%, wherein said phospholipids comprise at least approximately 65% phosphatidylcholine and at least approximately 2.4 wt% lysophasphatidylcholine. In one embodiment, the crustacean oil further comprises triglycerides, neutral lipids, approximately 20 - 26 wt% Omega-3 (e.g., n-3) fatty acids, and at least approximately 0.8 wt% free fatty acids. In one embodiment, the crustacean oil composition is krill oil.

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In one embodiment, the present invention contemplates a crustacean phospholipid-peptide complex (PPC) composition comprising a matrix of hydrolyzed protein, phospholipids and between approximately 200-500 ppm fluoride. In one embodiment, the phospholipids are at least 40 wt%. In one embodiment, the crustacean PPC composition further comprises approximately 0.044% (w/w) trimethyl amine and approximately 0.354% (w/w) trimethyl amine oxide. In one embodiment, the crustacean PPC composition further comprises at least 40% (w/w) triglycerides.

In one embodiment, the present invention contemplates a crustacean de-oiled phospholipid-peptide complex (PPC) composition comprising a matrix of hydrolyzed protein, between approximately 200-500 ppm fluoride, approximately 35% total fat, approximately 16.6% eicosapentaenoic acid, approximately 10.0% docosahexaenoic acid and at least 0.1 wt% free fatty acids. In one embodiment, wherein the total fat comprises less than 20% triglycerides, and approximately 69% other lipid components. In one embodiment, total fat comprises approximately 35.2% fatty acids, wherein approximately 30 wt% of said fatty acids are n-3 fatty acids. In one embodiment, the total lipids further comprise at least 68% phospholipids. In one embodiment, the de-oiled PPC further comprises approximately 2.2% lysophosphatidyl choline. In one embodiment, the de-oiled PPC further comprises approximately 115 mg/kg astaxanthin.

In one embodiment, the present invention contemplates a method for creating low fluoride crustacean compositions, comprising: a) disintegrating a crustacean catch into a material having a particle size ranging between approximately 1 – 25 millimeters; and b) separating said disintegrated crustacean material into a phospholipid-peptide complex (PPC) composition subfraction, wherein said subfraction comprises a fluoride content of less than 500 ppm. In one embodiment, the method further comprises extracting said PPC composition subfraction with a fluid comprising a solvent wherein a low fluoride oil is created, said oil having a fluoride content of less than 0.5 ppm. In one embodiment, the

extracting further creates a low trimethyl amine/trimethyl amine oxide oil, wherein said trimethyl amine is less than approximately 0.001% (w/w) and said trimethyl amine oxide is less than approximately 0.02% (w/w). In one embodiment, the separating is performed without emulsification. In one embodiment, the solvent comprises a non-polar solvent. In one embodiment, the solvent comprises at least one polar solvent. In one embodiment, the solvent comprises said non-polar solvent and said at least one polar solvent. In one embodiment, the non-polar solvent includes, but is not limited to, supercritical carbon dioxide and supercritical dimethyl ether. In one embodiment, the polar solvent includes, but is not limited to, ethanol and acetone. In one embodiment, the method further comprises hydrolyzing said crustacean material before said separating. In one embodiment, the extracting further creates a de-oiled PPC composition. In one embodiment, the polar solvent separates a phospholipid composition and a protein hydrolysate composition from said deoiled PPC composition. In one embodiment, the extracting comprises less than ten hours. In one embodiment, the extracting comprises less than five hours. In one embodiment, the extracting comprises less than two hours. In one embodiment, the crustacean material is krill material. In one embodiment, the separating comprises a centrifugal force of between approximately 1,000 - 1,800 g. In one embodiment, the separating comprises a centrifugal force of between approximately 5,000 - 10,000 g.

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In one embodiment, the present invention contemplates a composition comprising a mixture of a low fluoride crustacean PPC and a low fluoride de-oiled PPC, wherein said fluoride level ranges between approximately 200 – 500 ppm. In one embodiment, the crustacean PPC is krill PPC. In one embodiment, the crustacean de-oiled PPC is krill de-oiled PPC. In one embodiment, the crustacean PPC and crustacean de-oiled PPC are in a 1:1 ratio. In one embodiment, the mixture comprises a milled fine powder. In one embodiment, the powder comprises a particle size of approximately 250 μm. In one embodiment, the composition comprises a peroxide level of less than 0.1 %;(mEq/kg). In one embodiment, the composition further comprises microencapsulated polyunsaturated Omega-3 fatty acids. In one embodiment, the composition further comprises zinc oxide. In one embodiment, the composition further comprises marine peptides. In one embodiment, the composition further comprises at least one supplemental amino acid.

In one embodiment, the present invention contemplates a method, comprising formulating a composition comprising a low fluoride crustacean PPC and a low fluoride crustacean de-oiled PPC, wherein said fluoride level ranges between approximately 200 –

500 ppm. In one embodiment, the method further comprises milling said composition into a powder. In one embodiment, the method further comprises tabletting said composition into a tablet. In one embodiment, the method further comprises encapsulating said composition into a capsule. In one embodiment, the method further comprises mixing said powder with a food product. In one embodiment, the formulating further comprises microencapsulated polyunsaturated Omega-3 fatty acids. In one embodiment, the formulating further comprises zinc oxide. In one embodiment, the formulating further comprises marine peptides. In one embodiment, the formulating further comprises at least one supplemental amino acid.

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In one embodiment, the present invention contemplates a composition comprising a mixture of a low fluoride crustacean PPC and a crustacean protein hydrolysate, wherein said fluoride level ranges between approximately 200 – 500 ppm. In one embodiment, the crustacean PPC is krill PPC. In one embodiment, the crustacean protein hydrolysate is a krill protein hydrolysate. In one embodiment, the crustacean PPC and crustacean protein hydrolysate are in a 1:1 ratio. In one embodiment, the mixture comprises a milled fine powder. In one embodiment, the powder comprises a particle size of approximately 250 μm. In one embodiment, the composition comprises a peroxide level of less than 0.1 %;(mEq/kg). In one embodiment, the composition further comprises microencapsulated polyunsaturated Omega-3 fatty acids. In one embodiment, the composition further comprises marine peptides. In one embodiment, the composition further comprises marine peptides. In one embodiment, the composition further comprises at least one supplemental amino acid.

In one embodiment, the present invention contemplates a method, comprising formulating a composition comprising a low fluoride crustacean PPC and a crustacean protein hydrolysate, wherein said fluoride level ranges between approximately 200 – 500 ppm. In one embodiment, the method further comprises milling said composition into a powder. In one embodiment, the method further comprises tabletting said composition into a tablet. In one embodiment, the method further comprises encapsulating said composition into a capsule. In one embodiment, the method further comprises mixing said powder with a food product. In one embodiment, the formulating further comprises microencapsulated polyunsaturated Omega-3 fatty acids. In one embodiment, the formulating further comprises marine peptides. In one embodiment, the formulating further comprises marine peptides. In one embodiment, the formulating further comprises marine peptides. In one embodiment, the formulating further comprises at least one supplemental amino acid.

In one embodiment, the present invention contemplates a phospholipid-peptide complex (PPC) composition comprising a range between approximately 40 - 50% lipids and less than 0.5 mg/kg fluoride. In one embodiment, the lipids comprise phospholipids. In one embodiment, the present invention contemplates an oil composition comprising approximately 400-500 grams/kg phospholipids, approximately 200-260 grams/kg Omega-3 fatty acids, less than 0.5 mg/kg fluoride, approximately 15 grams/kg lysophosphatidic acid, and less than approximately 8 grams/kg free fatty acids. In one embodiment, the present invention contemplates a de-oiled phospholipid-peptide complex (PPC) composition comprising approximately 300-400 grams/kg lipids, wherein approximately 0.1-1.0 % are free fatty acids and a range between approximately 22-27 % (w/w) that are Omega-3 fatty acids. In one embodiment, the lipids comprise phospholipids. In one embodiment, the present invention contemplates a crustacean lipid composition comprising at least 75% phospholipids. In one embodiment, the lipid composition comprises between approximately 75% - 90% phospholipids. In one embodiment, the lipid composition comprises between approximately 75% - 80% phospholipids. In one embodiment, the present invention contemplates a dried protein hydrolysate composition comprising approximately 70 - 80% protein, approximately 1.5 - 3.0% lipids, and approximately 5 - 7% ash.

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In one embodiment, the present invention contemplates a method, comprising: a) providing; i) a hydrolyzed and disintegrated crustacean material; ii) at least one horizontal centrifuge capable of separating said hydrolyzed crustacean material; and iii) a fluid comprising a solvent; and b) separating said hydrolyzed crustacean material into a high fluoride solid fraction and a low fluoride hydrolyzed material fraction with a first horizontal centrifuge; c) separating said low fluoride hydrolyzed material fraction into a phospholipidpeptide complex (PPC) composition subfraction and a concentrated hydrolysate subfraction with a second horizontal centrifuge; and d) contacting said PPC composition subfraction with said fluid, wherein a low fluoride oil is extracted. In one embodiment, the disintegrated crustacean material has particle sizes between approximately 1 - 25 millimeters. In one embodiment, the first horizontal centrifuge separates said hydrolyzed crustacean material without emulisification. In one embodiment, the solvent comprises a non-polar solvent. In one embodiment, the non-polar solvent comprises supercritical CO<sub>2</sub>. In one embodiment, the solvent comprises a polar solvent. In one embodiment, the polar solvent comprises ethanol. In one embodiment, the second horizontal centrifuge comprises an extended separation pathway. In one embodiment, the contacting is performed at a pressure of less than 300 bar. In one embodiment, the non-polar solvent further extracts a de-oiled PPC composition from

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said PPC composition subfraction. In one embodiment, the ethanol separates a phospholipid composition and a protein hydrolysate composition from said de-oiled PPC composition. In one embodiment, the de-oiled PPC is separated from the PPC in less than ten hours. In one embodiment, the de-oiled PPC is separated from the PPC in less than five hours. In one embodiment, the de-oiled PPC is separated from the PPC in less than two hours. In one embodiment, the hydrolyzed crustacean material comprises hydrolyzed krill material. In one embodiment, the separating said hydrolyzed crustacean material is performed at a centrifugal force of between approximately 1,000 - 1,800 g. In one embodiment, the separating said low fluoride hydrolyzed material fraction is performed at a centrifugal force of between approximately 5,000 - 10,000 g. In one embodiment, the method produces a phospholipidpeptide complex (PPC) composition comprising a range between approximately 40%-50% lipid and less than 0.5 mg/kg fluoride. In one embodiment, the method produces an oil composition comprising approximately 400-500 grams/kg phospholipids, approximately 200-260 grams/kg Omega-3 fatty acids, less than 0.5 mg/kg fluoride, approximately 15 grams/kg lysophosphatidic acid, and less than approximately 8 grams/kg free fatty acids. In one embodiment, the method produces a de-oiled phospholipid-peptide complex (PPC) composition comprising approximately 300-400 grams/kg lipids, wherein approximately 0.1-1.0 % are free fatty acids and a range between approximately 20-28 % (w/w) are Omega-3 fatty acids. In one embodiment, the method produces a crustacean lipid composition comprising at least 75% phospholipids. In one embodiment, the lipid composition comprises a range between approximately 75% - 90% phospholipids. In one embodiment, the lipid composition comprises a range between approximately 75% - 80% phospholipids. In one embodiment, the method produces a dried protein hydrolysate composition comprising approximately 70 - 80% protein, approximately 1.5 - 3.0% lipids, and approximately 5 - 7 % ash.

In one embodiment, the present invention contemplates a system comprising: a) a solvent unit comprising at least one non-polar solvent inlet; b) an extraction tank unit in fluidic communication with the solvent unit, wherein the tank comprises an inlet configured to receive a phospholipid-protein complex composition; c) a separator unit comprising an outlet configured to release a low fluoride oil composition and residual co-solvent, wherein the separator is in fluidic communication with the tank; d) an absorbent unit in fluidic communication with the separator unit, wherein the absorbent unit is capable of recycling the non-polar solvent. In one embodiment, the non-polar solvent is a supercritical fluid. In one embodiment, the supercritical fluid comprises carbon dioxide. In one embodiment, the

supercritical fluid comprises dimethyl ether. In one embodiment, the solvent unit further comprises a co-solvent inlet. In one embodiment, the co-solvent is a polar solvent. In one embodiment, the polar solvent is ethanol or acetone. In one embodiment, the at least one non-polar solvent inlet comprises an unused non-polar solvent inlet. In one embodiment, the at least one non-polar solvent inlet comprises a recycled non-polar solvent inlet. In one embodiment, the solvent unit further comprises a fluid pump. In one embodiment, the tank unit is pressurized by the fluid pump. In one embodiment, the solvent unit further comprises a heater. In one embodiment, the phospholipid-protein complex composition in the tank unit is heated by the heater. In one embodiment, the separator outlet is in fluid communication with an evaporator. In one embodiment, the separator further comprises a horizontal centrifuge. In one embodiment, the horizontal centrifuge is a decanter centrifuge having an extended separation pathway. In one embodiment, the phospholipid-protein complex composition. In one embodiment, the low fluoride crustacean phospholipid-protein complex composition is a low fluoride krill phospholipid-protein complex composition.

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In one embodiment, the present invention contemplates a method for processing crustaceans, especially krills, in which method the crustaceans are disintegrated into smaller particles, fresh water is added to the disintegrated material, the water with the disintegrated material is heated and enzyme(s) are added for hydrolyzing the disintegrated material and said enzyme(s) is/are deactivated, the method further comprising steps: a) removing solids from the hydrolyzed material to reduce fluoride content of the material; b) separating phospholipid-peptide complex material and concentrated hydrolysate fraction from each other; c) drying said phospholipid-peptide complex material; and d) dividing the drying result, or PPC, to components by extraction(s) using at least a supercritical CO<sub>2</sub> as solvent, wherein the processing of crustaceans is started as soon as a crustacean catch has been decked on a ship or boat. In one embodiment, the fluoride content solids are removed from the hydrolyzed material by a decanter. In one embodiment, the phospholipid-peptide complex material and concentrated hydrolysate fraction are separated from each other by a sedicanter with high centrifugal forces and long clarification/separation zones to avoid an emulsification. In one embodiment, the method further comprises using in the extraction ethanol as a co-solvent in addition to the supercritical CO<sub>2</sub> to separate: i) a krill oil consisting of phospholipids and triglycerides, or neutral oil, and ii) a protein hydrolysate from the PPC. In one embodiment, the pressure of the solvent being at most 300 bar. In one embodiment, the extraction includes two steps: i) first using only the supercritical CO<sub>2</sub> as solvent to

separate de-oiled PPC from the PPC; and ii) second using only ethanol as solvent to separate phospholipids and protein hydrolysate from the de-oiled PPC. In one embodiment, the duration of the step when said de-oiled PPC is extracted from the PPC is at most three hours. In one embodiment, the method produces a phospholipid-peptide complex (PPC) composition comprising approximately 40% - 50% lipid and approximately 0.5 mg/kg fluoride. In one embodiment, the lipid comprises phospholipids. In one embodiment, the method produces an oil composition comprising approximately 400-500 grams/kg phospholipids, approximately 200-260 grams/kg Omega-3 fatty acids, approximately 0.5 mg/kg fluoride, approximately 15 grams/kg lysophosphatidic acid, and less than approximately 8 grams/kg free fatty acids. In one embodiment, the method produces a deoiled phospholipid-peptide complex (PPC) composition comprising approximately 300-400 grams/kg lipids, wherein approximately 0.1-1.0 % are free fatty acids and approximately 22-27 % (w/w) are Omega-3 fatty acids. In one embodiment, the method produces a crustacean phospholipid composition comprising approximately 75% polar lipids. In one embodiment, the method produces a dried protein hydrolysate composition comprising approximately 70 -80% protein, approximately 1.5 - 3.0% lipids, and approximately 5 - 7% ash.

#### **DEFINITIONS**

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The term "disintegrated material" as used herein refers to any biological material that has been subjected to a mechanical destruction and/or disruption that results in a composition having particle sizes of between approximately 1 - 25 millimeters, preferably between approximately 3 - 15 millimeters, more preferably between approximately 5 - 10 millimeters and most preferably approximately 8 millimeters.

The term "hydrolyzed material" as used herein refers to any biological material that has been subjected to high heat and/or enzymatic treatment. Such hydrolyzed materials would be expected to have phospholipid/peptide components that are physically separated from the components of the chitinous exoskeleton.

The term "crustacean" as used herein refers to any marine organism have a hard outside shell (e.g., a chitinous exoskeleton combined with a carbonate) encompassing a fleshy interior that is a living organism. More specifically, the crustaceans are usually considered a large class of mostly aquatic arthropods that have a chitinous or calcareous and chitinous exoskeleton, a pair of often much modified appendages on each segment, and two pairs of antennae. For example, a crustacean may include but not limited to, krill, lobsters, shrimps, crabs, wood lice, water fleas, and/or barnacles.

The term "horizontal centrifuge" refers to any device that is capable of rotating a mixture in the Z-plane (as opposed to the X-plane and/or Y-plane as with conventional centrifuges). This rotation is generated by a screw-type conveyor element aligned horizontally within a tube shaped enclosure. The induced centrifugal force then layers the heavier particles to the outside edges of the enclosure, while the lighter particles form layers closer to the center of the enclosure. Some horizontal centrifuges are modified to comprise an extended separation pathway and induce high gravitational forces (e.g., a sedicanter).

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The term "polar solvent" as used herein refers to any compound, or compound mixture, that is miscible with water. Such polar solvent compounds include, but are not limited to, ethanol, propanol and/or ethyl acetate.

The term "non-polar solvent" as used herein refers to any compound, or compound mixture, that is not miscible with water. Such non-polar solvent compounds include, but are not limited to, hexane, pentane, dimethyl ether and/or CO<sub>2</sub>. Either dimethyl ether or CO<sub>2</sub> may be used in a supercritical phase.

The term "supercritical" refers to any mixture comprising a chemical (e.g., for example, carbon dioxide (CO<sub>2</sub>) or dimethyl ether) in a fluid state while held at, or above, its critical temperature and critical pressure where its characteristics expand to fill a container like a gas but with a density like that of a liquid. For example, carbon dioxide becomes a supercritical fluid above 31.1 °C and 72.9 atm/7.39 MPa. Carbon dioxide usually behaves as a gas in air at standard temperature and pressure (STP), or as a solid called dry ice when frozen. If the temperature and pressure are both increased from STP to be at or above the critical point for carbon dioxide, it can adopt properties midway between a gas and a liquid. As contemplated herein, supercritical CO<sub>2</sub> can be used as a commercial and industrial solvent during chemical extractions, in addition to its low toxicity and minimal environmental impact. The relatively low temperature of the process and the stability of CO<sub>2</sub> also allows most compounds (i.e., for example, biological compounds) to be extracted with little damage or denaturing. In addition, because the solubility of many extracted compounds in CO<sub>2</sub> may vary with pressure, supercritical CO<sub>2</sub> is useful in performing selective extractions.

The term "fluoride" as used herein interchangeably and refer to any compound containing an organofluoride and/or an inorganic fluoride.

The term "high fluoride solid fraction" as used herein refers to a composition containing the vast majority of a crustacean's exoskeleton following a low g-force (e.g., between approximately 1,000 - 1,800 g) horizontal centrifugation separation of a hydrolyzed and disintegrated crustacean material. This fraction contains small particles of exoskeleton of

the crustacean that retains the vast majority of fluoride (i.e., for example, between 50 - 95%) in these organisms.

The term "low fluoride" as used herein may refer to the product of any method and/or process that reduced the fluoride from the original material by approximately 10-fold (i.e., for example, from 5 ppm to 0.5 ppm). For example, 'a low fluoride crustacean phospholipid-protein complex' comprises ten-fold less fluoride than 'a low fluoride hydrolyzed and disintegrated crustacean material'.

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The term "low fluoride hydrolyzed material fraction" as used herein refers to a composition containing the vast majority of a crustacean's fleshy internal material following a low g-force (e.g., between approximately 1,000 - 1,800 g) horizontal centrifugation separation of a hydrolyzed and disintegrated crustacean material. This fraction contains small particles of phospholipids, neutral lipids, proteins and/or peptides that is largely devoid of any fluoride (i.e., for example, between 5% - 50% of the raw hydrolyzed and disintegrated material).

The term "a low fluoride phospholipid-peptide complex composition subfraction" as used herein refers to a low fluoride composition containing the vast majority of lipid material following a high g-force (e.g., between approximately 5,000 - 10,000 g) horizontal centrifugation separation of a low fluoride hydrolyzed material fraction.

The term "concentrated hydrolysate composition subfraction" as used herein refers to a low fluoride composition containing the vast majority of water soluble lean material following a high g-force (e.g., between approximately 5,000 - 10,000 g) horizontal centrifuge separation of a low fluoride hydrolyzed material fraction.

The term "low fluoride oil" as used herein refers to a lipid-rich composition created by the extraction of a phospholipid-peptide complex composition subfraction using a selective extraction process, such as with a supercritical carbon dioxide fluid. Such a process removes approximately ten-fold of the fluoride from the raw hydrolyzed and disintegrated crustacean material.

The term "de-oiled phospholipid-peptide complex" as used herein refers to a low fluoride composition containing the vast majority of dry matter composition created by the extraction of a phospholipid-peptide complex composition subfraction using selective extraction process, such as a supercritical carbon dioxide fluid. A de-oiled PPC generally comprises a reduced triglyceride content in comparison to PPC.

The term "phospholipid composition" as used herein refers to a low fluoride composition comprising a high percentage of polar lipids (e.g., approximately 75%) created

by the extraction of a de-oiled phospholipid-peptide complex using a co-solvent, such as ethanol.

The term "protein hydrolysate" as used herein refers to a low fluoride composition comprising a high percentage of protein (e.g., approximately 70 - 80%) created by the extraction of a de-oiled phospholipid-peptide complex using a co-solvent, such as ethanol.

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The term "immediately" as used herein refers to a minimum practical period between decking a crustacean catch in a trawl bag and/or net coupled with a direct transfer to a suitable disintegraor. For example, this minimum practical period should preferably not exceed 60 minutes, more preferred to not exceed 30 minutes, even more preferred to not exceed 15 minutes.

The term "hydrolysis" as used herein refers to any break and/or disruption made in a protein structure of a disintegrated crustacean material, wherein in the naturally occurring protein sequences become shorter (i.e.., for example, by breaking peptide bonds of the amino acid sequence primary structure) and/or denatured (i.e., for example, an unfolding of the amino acid sequence secondary, tertiary and/or quaternary structure). This process may be controlled by hydrolytic enzyme(s). For example, one or more exogenous proteolytic enzymes (e.g. alkalase, neutrase, and enzymes derived from microorganisms or plant species) may be used in the process. Co-factors such as specific ions can be added depending on the used enzymes. The selected enzyme(s) can also be chosen for reducing emulsions caused by high content of phospholipids in the raw material. Besides the temperature, the hydrolysis takes place within optimal or near-optimal pH and sufficient time. For example, the exogenous enzyme alkalase the optimum pH is about 8, optimum temperature about 60°C and the hydrolysis time 40-120 minutes.

The term "solvent unit" refers to any enclosed volume configure to heat and pressurize a mixture of supercritical carbon dioxide fluid and/or a co-solvent (e.g., ethanol). Such an enclosed volume may be constructed out of any suitable material including but not limited to metals (e.g., steel, aluminum, iron etc.), plastics (e.g., polycarbonate, polyethylene etc.), fiberglass (etc.).

The term "extraction tank" refers to any enclosed volume configured to withstand heat and pressure sufficient to perform lipid and protein extraction from a raw biomass using a supercritical carbon dioxide fluid. As designed, the extraction tank contemplated herein is configured such that the solvents containing the extracted lipids and proteins rise to the tank top for transfer to a separator unit. Such an enclosed volume may be constructed out of any

suitable material including but not limited to metals (e.g., steel, aluminum, iron etc.), plastics (e.g., polycarbonate, polyethylene etc.), fiberglass (etc.).

The term "separator unit" refers to any enclosed volume configured with a centrifuge capable of separating the components of the extracted lipids and proteins received from an extraction tank. The respective extraction components exit the separator unit via outlet ports such that the remaining solvents (i.e., supercritical CO<sub>2</sub>) are transferred to an absorbent unit for recycling. Such an enclosed volume may be constructed out of any suitable material including but not limited to metals (e.g., steel, aluminum, iron etc.), plastics (e.g., polycarbonate, polyethylene etc.), fiberglass (etc.).

The term "absorbent unit" refers to any enclosed volume configured with materials that will remove contaminants from a supercritical CO<sub>2</sub> fluid. Such materials may include, but are not limited to charchol, coal, purifying gases, plastic polymer resins and/or filtration cartridges comprising single or dual-flat extruded nets (Tenax UK LTD, Wrexham, North Wales LL13 9JT, UK). Such an enclosed volume may be constructed out of any suitable material including but not limited to metals (e.g., steel, aluminum, iron etc.), plastics (e.g., polycarbonate, polyethylene etc.), fiberglass (etc.).

The term "in fluidic communication" refers to any means by which a fluid can be transported from one location to another location. Such means may include, but are not limited to pipes, buckets and/or troughs. Such means may be constructed out of any suitable material including but not limited to metals (e.g., steel, aluminum, iron etc.), plastics (e.g., polycarbonate, polyethylene etc.), fiberglass (etc.).

## BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 presents a flow diagram of one embodiment of a method to produce a low fluoride crustacean material.

Figure 2 presents a longitudinal centrifuge with an extended separation path. This specific example is a FLOTTWEG SEDICANTER horizontal decanter centrifuge.

Figure 3 depicts one example of an extraction plant suitable for use in the presently disclosed method. For example, the plant comprises a solvent unit (21), an extraction tank (22), separators (23) and adsorbents (24).

Figure 4 present exemplary data showing the extraction efficiencies of two different runs in accordance with one embodiment of the present invention.

#### DETAILED DESCRIPTION OF THE INVENTION

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The invention relates to a method for processing crustaceans (i.e., for example, krill) rich in lipids to produce compositions low in fluoride, trimethyl amine and trimethyl amine oxide comprising phospholipids, proteinaceous nutrients and oil (i.e., for example, neutral lipids and/or triglycerides).

Krill oil comprises lipids extracted with solvents from krill biomass. Krill biomass can be either fresh, whole krill (WO2008/060163A1), frozen whole krill (Neptune Technologies & Bioresources Inc., Canada), lyophilized whole krill (JP2215351) or krill meal (US20080274203). Solvents used in extracting lipids from krill biomass have been reported as acetone + ethanol (WO2000/23546; WO2002/102394), ethanol + hexane (Enzymotec Ltd), ethanol alone (JP2215351; Aker BioMarine ASA, Norway) or supercritical CO<sub>2</sub> + ethanol co-solvent (US2008/0274203; WO2008/060163). Solvent-free technology for obtaining krill oil has also been developed (US20110224450A1). Krill oil comprises a lipid fraction of raw krill biomass that is essentially free of protein, carbohydrates and/or minerals. Krill oil also comprises neutral lipids (e.g., mostly triglycerides), polar lipids (e.g., mostly phospholipids) and carotenoid astaxanthin. Although it is not necessary to understand the mechanism of an invention, it is believed that the lipid and/or fatty acid compositions of krill oil vary depending of the season.

In some embodiments, the present invention contemplates methods of processing crustacean biomass having unexpected findings including, but not limited to: i) removal of most of the exoskeleton from the crustacean biomass that results in low level of fluorides in a PPC composition and very low levels of fluoride in krill oil extracted from the PPC composition by a non-polar solvent (e.g., supercritical CO<sub>2</sub>) and, optionally, a polar cosolvent (e.g., ethanol); ii) a level of fluorides in the crustacean oil that is less than 0.5 ppm in contrast to conventional krill oil with fluoride content of approximately 5 - 100 ppm; iii) crustacean oil extracted from PPC by supercritical CO<sub>2</sub> and ethanol co-solvent has a minimal brown color suggesting that minimal degradation of astaxanthin or formation of tertiary oxidation products has occurred; iv) a reduced dark/brown color as measured on a Hunter L\* scale; v) a reduced pyrrole content as measured by absorption at 570 nm; v) minimal contents of free fatty acids (i.e., for example, 0.8 g/100 g of oil (~ 0.8% w/w)) and lysophosphatidylcholine (i.e., for example, 1.5 g/100 g of oil (~ 1.5% w/w)). These findings suggest that the lipids of crustacean biomass have undergone minimal hydrolysis during the initial processing steps producing PPC

## I. Historical Overview of Crustacean Processing Methods

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Publication GB 2240786 discloses a method for processing krill including removing a part of the fluoride content of krill. The removing is based on passing electric current through pulverized krill. However, fluoride-containing solid particles remain in the material.

Publication US 2011/0224450 (Sclabos Katevas et al., herein incorporated by reference) discloses a method for obtaining krill oil from whole raw krills using *inter alia* cooking, separating by decanter, and pressing. No solvents and extraction are used.

Publication WO 2008/060163 (Pronova Biopharma AS) discloses a method for obtaining krill oil using supercritical CO<sub>2</sub> and either ethanol, methanol, propanol or isopropanol as co-solvent. Fresh or pre-heated (about 90 °C) whole krills are used as the extraction feed material.

Publication WO 02/102394 (Neptune Technologies & Bioresources) discloses a method for obtaining krill oil using in different phases acetone and ethanol or e.g. ethyl acetate as solvents. Frozen whole krill is used as feed material.

Publication JP 2215351 (Taiyo Fishery) discloses a method for obtaining krill oil using ethanol as solvent. Lyophilized whole krills are used as feed material.

Publication US 2008/0274203 (Aker Biomarine ASA, Bruheim et al.)(herein incorporated by reference) discloses a method for obtaining krill oil from krill meal using supercritical fluid extraction in a two-stage process. Stage 1 removes the neutral lipid by extracting with neat supercritical CO<sub>2</sub> or CO<sub>2</sub> plus approximately 5% of a co-solvent. Stage 2 extracts the actual krill oils using supercritical CO<sub>2</sub> in combination with approximately 20% ethanol.

There are a number of problems associated with these conventionally known technologies of extracting krill lipids, including but not limited to: i) whole crustacean biomass contains high fluoride exoskeleton particles that results in the production of fluoride-contaminated crustacean oil; ii) crustacean oil having a brownish hue color may arise from exposing astaxanthin to excessive heat during crustacean biomass processing. Specifically, the brown color can arise from degradation of astaxanthin and/or from accumulation of the end products of non-enzymatic browning (e.g., Strecker degradation products or polymerized pyrroles). Although it is not necessary to understand the mechanism of an invention, it is believed that a brown color resulting from this non-enzymatic process results from oxidative degradation due to a reaction of secondary lipid oxidation products with amino groups from amino acids or proteins creating so-called tertiary oxidation products; iii) freezing the crustacean biomass for transportation to an extraction plant can result in relative stability, but

some changes in the product are known to occur over time, for example, one characteristic change in frozen krill is a partial hydrolysis of the lipids resulting in the accumulation of free fatty acids (FFA) arising from degradation of triglycerides, phospholipids and/or lysophospholipids, specifically lysophophatidylcholine (LPC), arising from hydrolysis of phosphatidylcholine; and iv) the use of heat and frozen storage can induce oxidation of lipids and proteins in crustacean biomass, where primary oxidation leads into formation of secondary oxidation products that are volatile and can be detected in krill oil as off-flavors or undesirable odor; and v) the separation of the krill oil from the feed material is quite inefficient, wherein only about a half of the oil can be extracted.

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#### II. Production Of Low Fluoride Crustacean Materials

In one embodiment, the present invention contemplates a method comprising forming a phospholipid-peptide complex (PPC) composition from a crustacean (i.e., for example, krill) immediately after the catch has been brought upon on board (e.g., decked) a boat and/or ship (i.e., for example, a fishing vessel). The process of creating the PPC composition comprises disintegrating the crustaceans into a disintegrated material comprising smaller particles (i.e., for example, between approximately 1 - 25 millimeters), adding water, heating the disintegrated material, adding enzyme(s) to hydrolyze the disintegrated material, deactivating the enzyme(s), removing solids (i.e., for example, exoskeleton, shell, and/or carapace) from the enzymatically processed material to reduce the fluoride content of the material, separating and drying the PPC composition. Preferably, the PPC composition is transferred to an on-shore facility (i.e., a fish oil extraction plant) where a low-fluoride crustacean oil is separated from the PPC composition using solvents including, but not limited to, supercritical CO<sub>2</sub> and/or ethanol. Using alternative extractions, de-oiled PPC compositions, phospolipids and/or protein hydrolysate compositions are also separated from the PPC composition.

- An advantage of some embodiments of the invention is that these crustacean products, like krill oil, have a low fluoride content. This is due to the fact that the solid crusteacean exoskeletal particles (i.e., for example, shell and/or carapace) are effectively removed from mass to be processed.
- Another advantage of the invention is that crustacean oil can be separated effectively, almost completely, from the disintegrated crustacean material (e.g., feed material) during the extraction. This is due to the fact that, in the extraction process with, for example, a supercritical CO<sub>2</sub> solvent, the feed material comprises a PPC composition. Although it is

not necessary to understand the mechanism of an invention, it is believed that the phospholipids of the feed material are embedded in a matrix of hydrolyzed protein which means that the close association between the phospholipids and hydrophobic/phosphorylated proteins is broken thus facilitating the extraction of the lipids.

5 - An advantage of the invention is that relatively low pressure and temperature can be used in the extraction, which means lower production costs.

- A further advantage of the invention is that disposal of residual solvents, common when using other more conventional lipid solvents, is avoided when using supercritical CO<sub>2</sub> as a solvent.
- A further advantage of the invention is that phosphatidylserine (PS), free fatty acids (FFA) and lysophosphocholine (LPC) contents are very low in the end products.
  - A further advantage of the invention is that a low fluoride crustacean oil product (i.e., for example, a low fluoride krill oil) has very little brown color. It is believed in the art that appearance of a brown color in crustacean oil indicates that unfavorable processes are occuring during the the manufacture of the feed material (e.g., a disintegrated crustacean material).

### A. Processing Of Crustaceans

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The present invention provides an industrial method for processing catches of crustaceans comprising a number of steps beginning with a very early and substantially complete removal of the crustacean's exoskeleton (i.e., for example, the crust, carapace and/or shell). Although it is not necessary to understand the mechanism of an invention, it is believed that the crustacean exoskeleton comprises a vast majority of fluoride in the organism. Consequently, this step thereby results in a substantial removal of fluoride from the crustacean material. The method also uses longitudinal centrifugation techniques that prevents separation problems caused by emulsions when processing a raw material with high content of phospholipids.

The method according to the present invention is initiated immediately after decking a catch of crustacean. It is of importance that the method according to the present invention is initiated as soon as possible after the crustacean catch has been decked since fluoride starts to leak/diffuse immediately from the exoskeleton into the crustacean's flesh and juices.

When using the term "immediately" in connection with starting the process according to the present invention this relates to the period from decking the crustacean catch and to the initial disintegration of the crustacean. This period of time should be kept to a minimum, and should preferably not exceed 60 minutes, more preferred not exceed 30 minutes, even more

preferred not exceed 15 minutes, and should include a direct transfer of the crustacean catch from the trawl bag and/or net to a suitable disintegrator. A disintegrator of the crustacean material may be a conventional pulping, milling, grinding or shredding machine.

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The crustacean catch is initially loaded into a disintegration appratus where the crustacean catch is subjected to pulping, milling, grinding and/or shredding to create a disintegrated crustacean material. The temperature of the disintegration process is around the ambient temperature of the water (i.e., for example, between approximately -2 and +1° C, but more preferably between approximately +0° C to +6° C) and may be performed by any convenient disintegration method. This disintegration process is also conventionally done by the previous known processing methods, and represents one of the obstacles according to the prior art because it produces large amounts of exoskeletal particles from the crustacean mixing in the milled material and producing a disintegrated paste with a high fluoride content. However, this high fluoride content is one of the reasons why the prior art processed crustacean material has limited applications and is less suitable for food, feed or corresponding food or feed additives compared to other marine raw materials e.g. pelagic fish.

According to the present invention the crustacean material is separated into a particle size suitable for a further separation step that does not interfer with the subsequent extraction steps. The disintegrating process is performed continuously and produces particle sizes up to 25 mm, a preferred particle size range is between approximately 0.5 - 10 mm and a more preferred size range is between approximately 1.0 - 8 mm.

Although it is not necessary to understand the mechanism of an invention, it is believed that this small particle size distribution represents one of advantages of the present invention because the fluoride has a tendency to leak out of the milled material and mingle with the rest of the raw material. However, this leaking process takes time and is not rapid enough to negatively impact a subsequent enzymatic hydrolysis step, provided the hydrolysis step is performed within specific parameters with respect to time and optimal, or near-optimal conditions, such as pH and temperature and optionally with the addition of co-factors such as specific ions depending on the used enzymes.

The temperature of the disintegrated material may, according to the present invention, be elevated to a temperature suitable for the subsequent enzymatic hydrolysis. Preferably, the temperature may be increased within seconds (e.g., 1-300 seconds, more preferred 1-100 seconds, even more preferred 1-60 seconds, most preferred 1-10 seconds) subsequent to the

disintegrating step for reducing the processing time and thereby preventing diffusion of fluoride and for preparing the material for the enzymatic hydrolysis.

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According to the present invention enzymes may be added directly to the disintegrated material or through the added water or both, before, during or after the disintegration process.

According to the present invention, exogenous proteolytic enzymes (e.g., alkalase, neutrase, enzymes derived from microorganisms including, but not limited to, *Bacillus subtilis* and/or *Aspergillus niger*, and/or or enzymes derived from plant species) may be added before, during or after the disintegration, and before, during or after the heating of the disintegrated material. The added enzyme(s) may be in the form of one single enzyme or a mixture of enzymes. The conditions of the hydrolysis should match the optimal hydrolytic conditions of the added enzyme(s) and the selection of optimal conditions for the selected exogenous hydrolytic enzyme(s) is known to the person skilled in the art. As an example, the exogenous enzyme alkalase having a pH optimum of about 8, a temperature optimum of 60° C and a hydrolysis time of 40-120 minutes. The selected enzymes, or combination of enzymes, should also be chosen for reducing emulsions caused by high content of phospholipids in the raw material.

An efficient amount of proteolytic enzyme(s) will be set after a process- and product optimization process that depends upon the efficiency of a specific chosen commercial enzyme or mix of enzymes. A typical amount by weight of commercial enzymes, as a ratio of the amount of the weight of the disintegrated raw material, are preferably between 0.5% and 0.05%, more preferably between 0.3% and 0.07% and most preferable between 0.2% and 0.09%. This hydrolysis step is aided by endogenous (natural) enzymes because rapid and uncontrolled autolysis is well known in fresh caught crustaceans.

In one embodiment, the exogenous enzymes breakdown the proteinaceous material in the disintegrated substance as well as speed up and/or accelerate the hydrolysis of the material to avoid and/or preclude the leaking of fluoride from the shell, carapace and crust. These hydrolytic enzymes, or a combination of hydrolytic enzymes, should also be carefully chosen to reduce emulsion in the separation process. For example, such enzymes may be selected from exo- and/or endopeptidases. If a mixture of enzymes is used, such a mixture may also include one or more chitinases for subsequently making the chitin-containing fraction(s) more amenable to further downstream processing. If chitinases are used, care must be taken for not increasing the leakage of fluoride from the shell/crust/carapace of the crustacean into the other fractions. However, since such fluoride leakage takes time, it is possible to perform such an enzymatic treatment within the preferred time parameters. A

more convenient alternative to including chitinases in the enzyme mix of the initial hydrolysis step will be to process the separated chitin-containing fraction subsequently to the separation step.

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In one embodiment, the leaking of fluoride from the milled exoskeletal material into the milled fleshy material is avoided by completing the disintegration/hydrolozing steps within a time interval of 100 minutes, preferably within 60 minutes, most preferred within 45 minutes calculated from the addition of the endogenous enzyme(s). The amount of enzyme(s) added is related to the type of enzyme product used. As an example it may be mentioned that the enzyme alkalase may be added in an amount of 0.1-0.5% (w/w) of the raw material. This should be taken into context with the added endogenous enzymes since the addition of more enzymes will reduce the time interval of the hydrolytic step. Although it is not necessary to understand the mechanism of an invention, it is believed that a short hydrologysis duration reduces the diffusion time of fluoride from particles of the exoskeleton into the proteinaceous material.

Subsequent to, or together with, the hydrolytic processing step the hydrolyzed and distintegrated crustacean material is passed through a particle removal device operating through a gravitational force such as a longitudinal centrifuge (i.e., for example, a decanter). This first separation step removes the fine particles containing a considerable amount of the fluoride from the hydrolysed or hydrolysing crustacean material to create a solids fraction. The centrifuge is operated with a g force between 1,000 and 1,800 g, more preferably between 1,200 and 1,600 g and most preferably between 1,300 and 1,500 g. Through this particle removal step a substantial amount of fluoride is removed from the proteinaceous crustacean fraction. The reduction of fluoride on a dry weight basis as compared to conventional crustacean meal, with a typical fluoride content of 1,500 ppm, may be up to 50%, even more preferred up to 85%, most preferred up to 95%.

The enzymatic hydrolysis may be terminated by heating of the hydrolysing material (incubate) to a temperature over 90° C, preferably between 92-98° C and most preferred between 92-95° C, prior to, during or after the separation step, as long as the hydrolysis duration lies within the above given boundaries. The hydrolysis is terminated before, during, or after the fine particle removal step, most preferred after the fine particle removal step. The temperature of the first centrifugation particle removal step, in one embodiment, depend on the optimal activity temperature of the enzyme (in the case where the enzymatic hydrolysis step is terminated by heating after the fine particle separation step).

The fluoride content in the prior art processed krill protein material (e.g., ~1,500 ppm) has limited applications and are less suitable for food or feed or corresponding food or feed additives. In one embodiment, removal of the fluoride content from the exoskeletal material may be followed by a further separation/purification of materials such as chitin, chitosan and astaxanthin. Such isolation procedures are known within the art. Steps may also be taken to further reduce the fluoride content from the isolated exoskeletal material using techniques including, but not limited to, dialysis, nanofiltration, electrophoresis or other appropriate technologies.

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Hydrolytic enzyme(s) deactivation may be performed in different ways, such as adding inhibitors, removing co-factors (e.g., crucial ions through dialysis), through thermal inactivation and/or by any other deactivating means. Among these, thermal inactivation, is preferred by heating the proteinaceous material to a temperature where the hydrolytic enzymes become denatured and deactivated. However, if a product where the relevant native proteins are not denatured is wanted, other means than heating for deactivating the hydrolytic enzymes should be selected.

A first centrifugation forms a de-fluoridated hydrolyzed and disintegrated crustacean material fraction and a solids fraction (e.g., containing high fluoride exoskeleton particles). As described below, the low fluoride hydrolyzed and disintegrated crustacean material fraction may be subsequently separated (e.g., by a second centrifugation) to form a low fluoride phospholipid-peptide complex (PPC) composition fraction and a lean low fluoride concentrated hydrolysate fraction (CHF) fraction that can be used as a food and/or feed additives, and a lipid fraction mainly consisting of neutral lipids. The PPC composition subfraction is rich in lipids, like a smooth cream with no particles, wherein the lipids are well suspended within the peptide components. This suspension results in small density differences between the different PPC composition components thereby making it difficult to further separate the PPC composition with common centrifugal separators and/or decanters. This is especially accentuated with crustacean catches during the second half of the fishing season.

Ordinary disc centrifugal separators (i.e., generating rotational force in the X and Y plane) do not work properly to separate a PPC composition subfraction into its respective components since emptying and necessary cleaning cycles with water will disturb separation zones. Conventional centrifugation separation processes result in the formation of unwanted emulsion products having a high phospholipid (PL) content and low dry matter concentrations. Standard decanters cannot separate the PPC composition subfraction into its

respective components due to a low g force limitation, short separation zone and an intermixing of light and heavy phases at the discharge of heavy phase from the machine.

In one embodiment, the present invention contemplates a method comprising separating a low fluoride PPC material into subfractions using a horizontal decanter centrifuge with an extended separation path. See, Figure 2. Horizontal centrifuges (e.g., generating a rotational force in the Z plane) are useful for the present invention comprise modified convention decanter centrifuges. For example, a PPC composition subfraction would enter an ordinary decanter from a bowl through a central placed feed pipe in the middle of the separation zone. In contrast, when using horizontal centrifuges as contemplated herein, the PPC composition subfraction enters at the end and at the opposite side of the outlet (1). This modification provides a significant improvement in the separation process by providing a considerably longer clarification/separation zone than ordinary decanters and utilizes the total available separation length (2) of the machine. The drive is able to impart high g-forces: 10,000 g for small machines and 5,000 to 6,000 g for high capacity machines, facilitating the separation of very fine, slow-settling PPC composition subfractions without the complications of emulsification. The PPC composition subfraction will be subjected to the highest g-force just before entering under the baffle (3). The different liquid layers separated from PPC composition subfraction are concentrated gradually along the axis of the horizontal centrifuge thereby exiting the machine under baffle (3) by the g force pressure generated by the machine (4). The separation of the PPC composition subfraction into a layer comprising about 27-30% dry matter makes the downstream processing efficient in terms of operating/robustness and as well economically considering both yield and costs of preparing the dry matter into a meal composition. The PPC composition subfraction separation also creates a layer comprising a lean hydrolysate that can be evaporated into a concentrated hydrolysate of greater than 60%.

#### B. Processing Of Krill

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One embodiment according to the invention is depicted as a flow diagram for the processing of krill. See, Figure 1. The function according to the method, or the process according to the invention is initiated immediately as a krill catch has been raised to the ship. Although it is not necessary to understand the mechanism of an invention, it is believed that fluoride immediately starts to leak/diffuse from the chitinous exoskeleton into the flesh and juices of the dead krills. "Immediately" means here a period at most 60 minutes, in practice, for example 15 minutes. During this period the krill catch is transferred from the trawl/net to a suitable disintegrator. In the disintegrator the krill material is crushed to relatively small

particles. The disintegrating can be performed by any convenient method: pulping, milling, grinding or shredding. The temperature in the disintegration process is around the ambient temperature of the water, i.e. between -2°C and +10°C, preferably between +0°C and +6°C. The disintegration produces large amount of chitinous debris among the rest of the krill material, thereby contributing to a high fluoride content.

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The particle size distribution of the disintegrated krill material is significant because of the above-mentioned fluoride leak from the chitinous debris and to the rest of the raw material. It is believed that the smaller particle sizes results in a more complete separation of the solids fraction from the disintegrated krill material. For this reason the preferable range of the particle size is 1.0 - 8 mm. However, the leaking process is relatively slow and has not time to be realized during the following process phases.

Next, fresh water is added to the disintegrated krill material (step 11). The volume/L of the water added is, for example, same as the weight/kg of the disintegrated krill material to be processed during the subsequent process phase of enzymatic hydrolysis. The temperature of the disintegrated krill material with the added water is increased such that it is suitable for the hydrolysis and enzyme(s) are added. The heating is carried out fast, within at most five minutes, after the disintegrating step to reduce the processing time and thereby to prevent diffusion of fluoride and to prepare the material for the enzymatic hydrolysis. The enzyme(s) can be added directly to the disintegrated krill material, or through the added water or both, before, during or after the heating step.

The term "hydrolysis" as used herein, means that breaks are made in the protein structure in the disintegrated substance, and the protein chains become shorter. This process is controlled by hydrolytic enzyme(s). For example, one or more exogenous proteolytic enzymes (e.g. alkalase, neutrase, and enzymes derived from microorganisms or plant species) may be used in the process. Co-factors such as specific ions can be added depending on the used enzymes. The selected enzyme(s) can also be chosen for reducing emulsions caused by high content of phospholipids in the raw material. Besides the temperature, the hydrolysis takes place within optimal or near-optimal pH and sufficient time (e.g., for example, the exogenous enzyme alkalase the optimum pH is about 8, optimum temperature about 60°C and the hydrolysis time 40-120 minutes).

The amount of proteolytic enzyme(s) can be set after a process/product optimization, and depends naturally on the efficiency of the chosen enzyme or mix of enzymes. A typical ratio of the weight of added commercial enzymes to the weight of the disintegrated krill material is between 0.05% and 0.5%, preferably between 0.1% and 0.2%. Fresh caught krill

is known for rapid and uncontrolled autolysis, or the destruction of the cells by endogenous (natural) enzymes, for which reason the treatment described here has to be proceeded without delays when the catch is not frozen.

The enzymatic hydrolysis also causes removing the bindings between the soft tissue of the krill and the exoskeleton. If a mixture of enzymes is used, the mixture may also include one or more chitinases to facilitate the further processing of the chitin-containing fractions. Chitinases are enzymes that break down glycosidic bonds in chitin.

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The enzymatic hydrolysis is finished within 100 minutes from the addition of the endogenous enzyme(s). The preferred duration Δt of the hydrolysis is shorter, for example 45 minutes (step 12). Relatively short hydrolysis duration is important, because in that case the diffusion of the fluoride from the exoskeleton particles to the other material is reduced.

The hydrolysis is stopped by deactivating the hydrolytic enzyme(s) (step 13). There are many ways to deactivate the enzymes. Here it is used the thermal one: the temperature of the enzymatically processed material is increased over 90°C, preferably between 92-98°C, in which case the hydrolytic enzymes become denatured. In practice the deactivating of the hydrolytic enzyme(s) can be performed also during or after the solid particle removal.

The solid particles (e.g., krill exoskeleton) are removed from the enzymatically hydrolyzed and disintegrated krill material by passage through a device based on the centrifugal force such as a conventional horizontal centrifuge and/or decanter (step 14). Although it is not necessary to understand the mechanism of an invention, it is believed that these solid particles, or solids, originate from the exoskeleton of krills and, as mentioned, contain a considerable amount of the fluoride. The decanter is operated with a force between 1,000 and 1,800 g, preferably between 1,300 and 1,500 g. Through this particle removal step a substantial amount of fluoride, more than 90 %, is removed from the krill material. The temperature in the decanter is for example 90°C, and if the deactivation of the enzyme(s) is done after the removal of solids, the temperature in the decanter is then increased to e.g. 93°C.

Next, the hydrolyzed and disintegrated krill material with low fluoride content is modified by passage through an extended separation path horizontal centrifuge (i.e., for example, a sedicanter). See, Figure 1 step 15, and Figure 2. In the sedicanter, the hydrolyzed and disintegrated krill material, is separated into the valuable fatty portion, or PPC (phospholipid-peptide complex) material fraction, and a CHF portion (concentrated hydrolysate fraction).

The separation of hydrolyzed and disintegrated krill material into the PPC material is difficult because of the small density differences within the krill material. The sedicanter is a modified horizontal centrifuge including a long horizontal clarification/ separation zone and generating high centrifugal forces (5,000 to 6,000 g). These features facilitate the separation of fine, slow-settling PPC without emulsification. The latter is a problem in the ordinary centrifuges with short separation zone and lower forces, and in which water is used in emptying and cleaning cycles. The dry matter concentration of PPC material, pressured out from the sedicanter, is about 27-30%.

The PPC material may be then dried to a meal to avoid the lipid oxidation. Figure 1, step 16. The drying process is gentle with low temperature (0-15°C, preferably 2-8°C) and inert conditions, which give a reduced oxidative stress on the long-chain poly-unsaturated omega-3 fatty acids. A lyophilisation process would also be suitable since this avoids an over-heating of the product.

The PPC krill meal, or more briefly PPC, is then packed in air tight bags under nitrogen atmosphere for later direct use and continuation process.

A typical mass balance of the processed raw lean Antarctic krill is shown below in Table I:

Table I: Typical Mass Balance Of Antarctic Krill

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20	Matter F	rom 500kg raw krill + water	Dry weight
	Wet PPC material	80 kg	28%
	PPC meal	25 kg	97%
	Hydrolysate	770 kg	6%
	CHF	78 kg	60%
25	Fluoride-containing	g particles 45 kg	40%
	Neutral oils	<5 kg	

The fluoride content, prior to separation, in hydrolyzed and disintegrated krill material is 1.2 g/kg, whereas, after separation, the PPC is at most 0.5 g/kg and typically 0.3 g/kg. Thus, about two thirds of the fluoride has been removed.

When the PPC is further processed, components may be isolated by an extraction. In this phase, a solvent may be used. Figure 1, step 17. For example, to obtain krill oil from the PPC, supercritical CO<sub>2</sub> and/or ethanol may be utilized, either separately or in combination. The extraction process yields, in addition to the krill oil, a protein hydrolysate (step 18).

Compressing and heating a material (e.g., for example, carbon dioxide or dimethyl ether) to above its critical temperature and pressure results in a supercritical fluid. The

density is intermediate between a liquid and a gas and can be varied as a function of temperature and pressure. Hence, the solubility of supercritical fluids can be tuned so that selective extractions can be obtained. Due to the gas like properties, rapid extractions can be accomplished compared to liquid extractions as the diffusion rates are higher. CO<sub>2</sub> is a commonly utilized supercritical fluid as its critical parameters can easily be reached. For example, one report has demonstrated a low yield of krill phospholipids by using supercritical fluid extraction at a pressure of 500 bar and a temperature of 100°C. Yamaguchi (1986). A second report provides data on specific process conditions, which include pressure and temperature ranges (e.g., 300 to 500 bar and 60 to 75°C). These data are from a pilot scale process wherein an extraction of 84 to 90% of krill total lipids was achieved. Bruheim et al., United States Patent Application Publication Number 2008/0274203 (herein incorporated by reference).

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Supercritical CO<sub>2</sub> is also non-flammable, cheap and inert, wherein such factors are relevant when considering industrial applicability. The inertness results in low grade of oxidation of labile compounds during extraction. CO<sub>2</sub> also has a low surface tension which is an advantage so that the extraction medium can penetrate the material efficiently. In order to extract more polar substances, the CO<sub>2</sub> can be mixed with a polar solvent such as ethanol. The level of modifier can be varied to provide extra selectivity as well.

Consequently, currently available industrial scale supercritical fluid extraction processes using high temperatures and pressures has resulted in a low extraction efficiency of conventional krill meal thereby providing an insufficient oil yield to provide a commercially feasible solution for krill extraction. Further, these currently available extraction processes do not solve the problems discussed herein regarding providing improved low fluoride meal and/or oil compositions.

Therefore, the improved solvent extraction methods described herein have been developed. In one embodiment, co-solvents are used with supercritical CO2 or supercritical dimethyl ether either alone or in various combinations of ethanol, hexane, acetone. For example, if ethanol is used alone as an extraction solvent, it has been observed that krill material is less selective than extraction with supercritical CO<sub>2</sub>. Pronova et al., WO 2008/060163 A1. As a result, undesirable substances are extracted into the krill oil resulting in a need for additional post-extraction clean-up/processing. Further, ethanol-only extracted krill oil tends to have higher viscosity and darker color which is independent of astaxanthin content of the oil.

In some embodiments, the present invention contemplates methods that have unexpected findings including but not limited to: i) PPC was extracted using low pressures (i.e., for example, between approximately 177 to 300 bar) and low temperatures (i.e., for example, between approximately 33 and 60°C); and ii) high yield of lipid extract was produced (data available). It appears that krill meal comprising hydrolyzed protein allows for easier extraction of the associated lipids in particular the phospholipid rich fraction of krill oil.

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The data presented herein demonstrates that supercritical CO<sub>2</sub> was found to be a selective extraction method as it produced high purity extracts containing triglycerides, phospholipids and astaxanthin with minimal brown color and superior organoleptic quality as compared to krill oils produced by ethanol-only extraction and/or acetone + ethanol extraction. Brown color of krill oil is considered to be undesirable. The exact origin of the brown color is unknown but it is believed to be associated with oxidation of krill lipids during the manufacture of krill meal phospholipids and/or degradation of the carotenoid astaxanthin.

The properties of such a supercritical fluid can be altered by varying the pressure and temperature, allowing selective component extraction. Extraction conditions for supercritical CO<sub>2</sub> are above the critical temperature of 31°C and critical pressure of 74 bar. Addition of modifiers may slightly alter these values. For example, neutral lipids and cholesterol can be extracted from egg yolk with CO<sub>2</sub> pressures up to 370 bar and temperature up to 45°C, while using higher temperature, e.g. 55°C, would result in increased rate of phospholipid extraction. CO<sub>2</sub> has a high industrial applicability because it is non-flammable, cheap and inert. The inertness results in low oxidation of labile compounds during extraction.

As mentioned, either supercritical CO<sub>2</sub> or supercritical dimethyl ether is fluid. Its density is intermediate between a liquid and a gas and can be varied as a function of temperature and pressure. Hence, the solubility of supercritical fluids can be tuned so that selective extractions can be obtained. Due to the gas-like properties, rapid extractions can be accomplished compared to liquid-extractions. In the present method the extraction is effective; even 95% of the krill oil existing in the PPC is separated. Although it is not necessary to understand the mechanism of an invention, it is believed that the phospholipids of the feed material are embbded in a matrix of hydrolyzed protein which means that the close association between the phospholipids and hydrophobic/phosphorylated proteins is broken thus facilitating the extraction of the lipids. In addition, a minimal amount of fluoride content is transferred to oil during the CO<sub>2</sub> extraction process. For example, the fluoride

content of PPC is about 0.3 g/kg, but after the CO<sub>2</sub> extraction the fluoride content of the krill oil is less than 0.5 mg/kg.

Alternatively, when using only supercritical CO<sub>2</sub> as solvent, triglycerides and/or neutral oil may be separated from the PPC composition subfraction. Figure 1, step 19. In one embodiment, supercritical CO<sub>2</sub>-only extraction also generates a low fluoride 'de-oiled PPC' composition. Although it is not necessary to understand the mechanism of an invention, it is believed that de-oiled PPC is the most valuable portion of the PPC composition subfraction. When thereafter, the de-oiled PPC composition may be extracted using ethanol as a solvent, wherein a phospholipid subfraction and a protein hydrolysate fraction is also generated. See, Figure 1, step 1A.

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In one embodiment, the present invention contemplates a system comprising an extraction plant, including but not limited to, a solvent unit 21, vertical tank 22, separators 23 and adsorbents 24. See, Figure 3. Normal CO<sub>2</sub> and possible co-solvent are fed to the solvent unit, which comprises a pump to generate a certain pressure (p) and a heater to generate a certain temperature (T). The supercritical CO<sub>2</sub> with possible co-solvent are then fed to the lower end of the tank 22. The feed material, in this case the PPC, is fed to the tank by means of a pump. Material affected by the solvent flows out of the upper end of the tank. The separators 22 separate the extract result, for example krill oil, to output of the system. If ethanol is used as co-solvent, it follows the extract proper and has to be evaporated away. The CO<sub>2</sub> continues its circulation to adsorbents 23, where it is cleaned, and thereafter back to the solvent unit 21.

In one embodiment, the present invention contemplates low fluoride PPC compositions including, but not limited to, polar lipids (~ 43% w/w) and/or neutral lipids (~ 46% w/w). For example, the PPC neutral lipids may range between approximately 40 – 50% (w/w). In one embodiment, the polar lipids include, but are not limited to, phosphatidylethanoamine (~ 3% w/w), phosphatidylinositol (~ < 1% w/w), phosphatidylserine (~ 1% w/w), phosphatidylcholine (~ 38% w/w) and/or lysophosphatidylcholine (~ 2% w/w). In one embodiment, the neutral lipids include, but are not limited to triacylglycerol (~ 40% w/w), diacylglycerol (~ 1.6% w/w), monoacylglycerol (~ 1% w/w), cholesterol (~ 2% w/w), cholesterol esters (~ 0.5% w/w), free fatty acids (~ 2% w/w) and fat (~ 48% w/w). In one embodiment, the neutral lipid fat comprises approximately 75% fatty acids. In one embodiment, the neutral lipid fat fatty acids include, but are not limited to, saturated fatty acids (~ 28% w/w), monenoic fatty acids (~ 22% w/w),

n-6 polyunsaturated fatty acids ( $\sim 2\%$  w/w) and/or n-3 polyunsaturated fatty acids ( $\sim 26\%$  w/w). See, Example XIII.

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Phospholipid profiles have been created to evaluate low fluoride krill oil extracted by the methods described herein. For example, nuclear magnetic resonance technology has determined that phosphatidylcholine is the largest phospholipid component of krill oil and its proportion is relatively stable. Several krill oil samples underwent independent analysis. See, Example XII. In one embodiment, the present invention contemplates a low fluoride krill oil comprising approximately 39 - 52% (w/w) phospholipids. In one embodiment, the phospholipids comprise phosphatidylcholine ranging between approximately 65 - 80% (w/w). In one embodiment, the phospholipids comprise alkyl acyl phosphatidylcholine ranging between approximately 6-10% (w/w). In one embodiment, the phospholipids comprise phosphatidylinositol ranging between approximately 0.3 - 1.6% (w/w). In one embodiment, the phospholipids comprise phosphatidylserine ranging between approximately 0.0-0.7 % (w/w). In one embodiment, the phospholipids comprise lysophosphatidylcholine ranging between approximately 2.4 – 19% (w/w). In one embodiment, the phospholipids comprise lyso acyl alkyl phosphatidylcholine ranging between approximately 0.6 - 1.3% (w/w). In one embodiment, the phospholipids comprise phosphatidylethanolamine ranging between approximately 1.4 – 4.9% (w/w). In one embodiment, the phospholipids comprise alkyl acyl phosphatidylethanolamine ranging between approximately 0.0 - 2.1 % (w/w). In one embodiment, the phospholipids comprise a combination of cardiolipin and Nacylphosphatidylethanolamine ranging between approximately 1-3% (w/w). In one embodiment, the phospholipids comprise lysophosphatidylethanolamine ranging between approximately 0.5 - 1.3% (w/w). In one embodiment, the phospholipids comprise lyso alkyl acyl phosphatidylethanolamine ranging between approximately 0.0 and 0.3% (w/w).

As described above, the non-polar solvent extraction of a low fluoride crustacean oil results in the production of a low fluoride de-oiled phospholipid-protein complex composition (de-oiled PPC). Although it is not necessary to understand the mechanism of an invention, it is believed that the low fluoride de-oiled phospholipid-protein complex comprises a fluoride content similar to the low fluoride PPC complex (e.g., between approximately 200 - 500 ppm). A component analysis of de-oiled PPC includes, but is not limited to, polar lipids ( $\sim 69\%$  w/w) and/or neutral lipids ( $\sim 20\%$  w/w). In one embodiment, the polar lipids include, but are not limited to, phosphatidylethanoamine ( $\sim 4.2\%$  w/w), phosphatidylinositol ( $\sim < 1\%$  w/w), phosphatidylcholine ( $\sim < 1\%$  w/w) and/or lysophosphatidylcholine ( $\sim 2\%$  w/w). In one embodiment, the neutral

lipids include, but are not limited to triacylglycerol ( $\sim 17\%$  w/w), diacylglycerol ( $\sim 0.6\%$  w/w), monoacylglycerol ( $\sim < 1\%$  w/w), cholesterol ( $\sim 1\%$  w/w), cholesterol esters ( $\sim 0.5\%$  w/w), free fatty acids ( $\sim 1\%$  w/w) and fat ( $\sim 35\%$  w/w). In one embodiment, the neutral lipid fat comprises approximately 69% fatty acids. In one embodiment, the neutral lipid fat fatty acids include, but are not limited to, saturated fatty acids ( $\sim 21\%$  w/w), monenoic fatty acids ( $\sim 13\%$  w/w), n-6 polyunsaturated fatty acids ( $\sim 2\%$  w/w) and/or n-3 polyunsaturated fatty acids ( $\sim 31\%$  w/w). See, Example IX.

#### III. Production Of Low Trimethyl Amine Crustacean Materials

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Trimethylamine (TMA) is an organic compound comprising a chemical formula of N(CH<sub>3</sub>)<sub>3</sub>. TMA is a colorless, hygroscopic, and flammable tertiary amine that may have a strong "fishy" odor in low concentrations and an ammonia-like odor at higher concentrations. TMA may be produced commercially and is also a natural by-product of plant and/or animal decomposition. It is the substance mainly responsible for the odor often associated with rotting fish, some infections, and bad breath. It is also associated with taking large doses of choline and carnitine.

Chemically, TMA comprises a nitrogenous base and can be readily protonated to give trimethylammonium cation. Trimethylammonium chloride is a hygroscopic colorless solid prepared from hydrochloric acid. Trimethylamine is a good nucleophile, and this reaction is the basis of most of its applications.

Trimethylamine N-oxide (TMAO) is an organic compound comprising a formula (CH<sub>3</sub>)<sub>3</sub>NO. This colorless solid is usually encountered as the dihydrate. TMAO is an oxidation product of TMA, a common metabolite in animals. TMAO is also an osmolyte found in saltwater fish, sharks and rays, molluses, and crustaceans. Further, TMAO may function as a protein stabilizer that may serve to counteract urea, the major osmolyte of sharks, skates and rays. TMAO has high concentration in deep-sea fishes and crustaceans, where it may counteract the protein-destabilizing effects of pressure. Yancey, P. "Organic osmolytes as compatible, metabolic, and counteracting cytoprotectants in high osmolarity and other stresses" *J. Exp. Biol.* 208(15):2819–2830 (2005). TMAO decomposes to trimethylamine (TMA), which is the main odorant that is characteristic of degrading seafood.

Removal of TMA/TMAO compounds from crustacean products confers a useful advantage in that these compounds contribute to the strong, unpleasant smell of crustacean oils. Consequently, low TMA/TMAO compounds have an improved industrial applicability as compared to traditionally prepared crustacean oils.

In one embodiment, the present invention contemplates a method comprising extracting a low fluoride protein peptide complex (PPC) is a suitable raw material for krill oil production by extraction with any combination of solvents including, but not limited to, ethanol, acetone, ethyl acetate, carbon dioxide, or dimethyl ether to produce a low fluoride-low trimethyl amine crustacean product. In one embodiment, the low fluoride-low trimethyl amine crustacean product comprises an oil. In one embodiment, the low fluoride-low trimethyl amine crustacean produce comprises a de-oiled PPC.

Dimethyl ether (DME) has been previously reported as an extraction solvent for polyunsaturated fatty, but not for the preparation of low TMA products. Catchpole et al. "Extraction Of Highly Unsaturated Lipids With Liquid Dimethyl Ether" WO 2007/136281. When DME is in a supercritical form, the solvent has sufficient solvent power to extract phospholipids resulting in rapid and gentle extractions. DME can be used on wet raw materials and can be operated at low pressures as compared to other supercritical fluids such as CO<sub>2</sub>. In one embodiment, the present invention contemplates a crustacean extraction product comprising krill oils with a low TMA/TMAO crustacean oil. In one embodiment, the low TMA/TMAO crustacean oil is a krill oil.

# IV. Formulated Compositions

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In some embodiments, the present invention contemplates compositions comprising low fluoride crustacean PPC or compositions comprising low fluoride crustacean de-oiled PPC compositions and/or protein hydrolysates as described herein. In one embodiment, the compositions comprises mixtures of the crustacean PPC complex, crustacean de-oiled PPC and the protein hydrolysates in any combination. Although it is not necessary to understand the mechanism of an invention, it is believed that the mixed ratio can be any ratio but is preferably a ratio of approximately 1:1. In one embodiment, the mixture comprises a milled fine powder. In one embodiment, the powder has a particle size of approximately 250  $\mu$ m. In one embodiment, the compositions have improved stability because of lower peroxide (e.g., < 0.1 %; mEq/kg) and/or aniside levels (< 0.1 %; w/w). In one embodiment, the compositions have improved stability because of lower microbiological contamination. In one embodiment, the composition further comprises microencapsulated polyunsaturated Omega-3 fatty acids. In one embodiment, the composition further comprises marine peptides. In one embodiment, the composition further comprises marine peptides. In one embodiment, the composition further comprises marine peptides. In one embodiment, the composition further comprises marine peptides.

In some embodiments, the present invention contemplates a method for formulating a composition comprising a low fluoride crustacean PPC and/or a low fluoride crustacean deoiled PPC and/or a protein hydrolysate as described herein. In one embodiment, the composition is a powder. In one embodiment, the composition is a tablet. In one embodiment, the composition is a capsule. In one embodiment, the method further comprises mixing the powder with a food product. In one embodiment, the mixing further comprises a microencapsulated polyunsaturated Omega-3 fatty acids. In one embodiment, the mixing further comprises marine peptides. In one embodiment the mixing further comprises at least one supplemental amino acid.

#### **EXPERIMENTAL**

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#### Example I

15 <u>Production Of Low Fluoride Krill Oil</u>

The feed material, 'Emerald krill meal' granules (Olymeg® or low fluoride PPC prepared as described herein), were supplied in a sealed plastic bag containing approximately 25kg. The feed material was kept frozen until used in extractions. The granules have a size distribution typically in the range 2 to 5mm, but a number of fine fragments were also present. The granules are greasy to the touch but still break up under compression rather than smear.

5 kg batches of feed material in granular form, as processed using supercritical CO<sub>2</sub> as solvent and azeotropic food grade ethanol as co-solvent, the weight of the ethanol being 23% of the weight of CO<sub>2</sub>. The plant was pre-pressurised to operating pressure with CO<sub>2</sub> only, and ethanol was added when CO<sub>2</sub> circulation started. Solvent to feed material ratio was 25:1 or greater and co-solvent to feed material ratio was 5:1. Runs were carried out under two extraction conditions; 300 bar at 60°C, and 177 bar at 40°C. See, Table II.

Table II -Krill Oil Extraction Conditions

30		<u>Run 1</u>	<u>Run 2</u>
	Feed Mass (g, as received)	5000.5	5000.9
	Extraction pressure (bar)	300	177
	Extraction temperature (°C)	60	33
35	First separator pressure (bar)	90	90
	First separator temperature (°C)	41	41

	Second separator pressure (bar)	48-50	48-50
	Second separator temperature (°C)	39	39
	CO <sub>2</sub> used with ethanol co-solvent (kg)	132.6	134.9
	Additional CO <sub>2</sub> at end of run (kg)	33.1	44.5
5	Total ethanol used (kg)	31.65	32.19

The extracted krill oil material was passed through two separation vessels in series, held at 90 bar and 45-50 bar respectively. The final krill oil material collected from both separators was pooled together and the ethanol was evaporated. The residual feed material comprises a deoiled feed material (e.g., for example, de-oiled PPC) having a reduced lipid content in comparison to the starting feed material. *See, Example IX*.

After ethanol evaporation, krill oil cumulative extraction curves were generated for both Run 1 and Run 2 by independently analyzing each sample taken during the extraction runs. *See, Table III.* 

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Table III – Progressive krill oil extraction sample points and yields.

	Sample Number	1	2	3	4	5	6	Total
20	Run 1 Cumulative CO <sub>2</sub> (kg/kg feed) Extracted oil (g, dry)	5.5 1137	9.1 398	13.4 282	17.8 135	22.0 78	33.1 86	33.1 2115
25	Run 2 Cumulative CO <sub>2</sub> (kg/kg feed) Extracted oil (g, dry)	5.6 715	9.1 496	13.5 368	17.5 220	21.5 149	34.4 129	34.4 2077

A total yield of 41-42 wt% of the feed material was achieved for all runs. The runs carried out at 300 bar and 60°C had a higher initial rate of extraction. The curves indicate that the extraction is virtually complete at Sample Number 5 after a cumulative CO<sub>2</sub> use ranging between 21.5 - 22.0 kg per kg of feed material. Estimated maximum extraction is achieved at a point where the CO<sub>2</sub>:feed ratio is 26.5:1. See, Figure 3 (estimated maximum extraction is marked by an arrow). The ratio of azeotropic ethanol to CO<sub>2</sub> was 0.24:1 for the 300 bar runs, and slightly higher at 0.26:1 for the lower pressure run.

This method of krill oil production resulted in the near complete extraction of total lipids from the krill meal (e.g., for example, approximately 95% of neutral lipids and 90% of phospholipids. The final yield was similar for both the high and low pressure runs, but neutral lipids were more rapidly extracted at higher pressure. The phospholipid extraction rate was similar under both extraction conditions. As detailed below, in this extraction process, the

pooled krill oil total lipid had an overall phospholipid level of just over 40 wt% and both phosphatidyl inositol and phosphatidyl serine were poorly extracted.

Phospholipid profiles of the various krill material compositions were then determined using traditional column chromatography techniques. *See, Table IV*.

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Table IV – Comparative Phospholipid Profiles Of Krill Compositions (run 1)

Sample	Olymeg 10071199	Extract 1	Extract 2	Extract 3	Extract 4	Extract 5	Extract 6	Residue (Top)	Residue (Bottom)
		•	W	t% of total PL					
PC	70.1	80.4	77.1	76.9	75.9	73.5	72.7	40.2	32.5
AAPC	8.5	8.0	9.0	9.8	9.1	10.6	9.0	7.5	7.8
PI	1.8				0.7	0.6	0.6	6.2	10.1
P8	1.0							5.5	8.1
LPC	6,9	4.6	5.6	5.7	6.0	6.8	7.5	13.4	8.9
LAAPC	1.7	1.2	1.2	1.0	1.3	1.2	1.4	3.2	2.6
PE	5.3	3.6	4.0	3.5	3.8	3.5	4.5	9.4	9.4
EPLAS	0.6	0.0	0.5	0.5	0.5	0.5	0.3	1.0	2.2
AAPE	2.0	1.1	1.5	1.3	1.6	1.6	2.0	4.4	4.9
LPS								0.7	1.9
CL/NAPE	1.0	0.9	0.7	8.0	0.8	1.2	1.6	4.2	5.7
LPE	8.0	0.3	0.4	0.4	0.4	0.4	0.4	3.2	4.5
Total PL (wt% of lipid)	40.88				<del>  -</del>			81.46	80.96
Lipid yield (wt%)	44.7							4.9	5.9
Total Pi. (wt% of sample)	18.3	26.68	46.03	57.94	71 34	76.13	78.50	4.0	4.8

The first column shows the specific phospolipids that were analyzed. The second column show the phospholipid profile of the starting feed material (e.g., a low fluoride PPC prepared as described herein, or 'Olymeg<sup>®</sup>). Columns three – eight (Extracts 1-6) show the phospholipid profile of each krill oil sample taken during the extraction process as described above. The last two columns show the phospholipid profile of the residual extracted feed material sampled from either the top and/or the bottom of the phospholipid extraction column.

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The data show that the major phospholipid in the extracted krill oil samples is phosphatidyl choline (PC), ranging approximately from 72.7% to 80.4% of total phospholipids, including contributions from both alkyl acyl phosphatidyl choline (AAPC) and lyso phosphatidyl cholines (e.g., for example, LPC and/or LAAPC). Smaller amounts of phosphatidyl ethanolamine (PE) are present in both the feed material (column 1, ~ 5.3%) and in the krill oil extract samples (columns 3 - 8), ~ 3.5 - 4.5%). Alkyl acyl and lyso forms of PE (AAPE, LPE) are also present in the feed material and krill oil extracts. Phosphatidyl inositol (PI) and phosphatidyl serine (PS) are present in the feed material, but because they are poorly soluble in ethanol, these phospholipids are poorly extracted and are therefore concentrated in the extracted feed material residue (e.g., having a higher level in the residual PPC in comparison to the feed material, see columns 9 and 10).

Further analysis determined the overall relative lipid component proportions of the extracted krill oil. *See, Figure V*.

Table V – Main Lipid Components Of Extracted Krill Oil (%w/w)

	TAG	Polar lipid	Sterols	FFA	Astaxanthin	Total lipid
Run 1	40,3	46,9	1,9	ND	0,05	92,2
Run 2	42,1	50,2	2	ND	0,05	95,3

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The data show: i) a relative absence of free fatty acids (FFAs); ii) less than 2% of sterols; iii) 40 wt% of triacylglycerides (TAGs); and iv) approxiately 50% phospholipids (e.g., polar lipids). While FFA's were not detected (ND) in this particular example, it is believed that extracted krill oils may comprise between approximately 0.01 – 0.1 % FFA of total lipids. As described above, the extraction process results a yield of between approximately 92.2 – 95.3% of the feed material total lipid.

The method and products according to the invention has been described above. The method can naturally vary in its details from those presented. The inventive idea may be applied in different ways within the limits as described herein.

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# Example II Lipid Extraction Efficiency

This example demonstrates an exemplary analytical lipid extraction with the Soxhlet method comparing conventional krill meal with a low fluoride krill meal (e.g. low fluoride PPC) as described herein. Soxhlet method is a standard method in quantitative determination of fat content of foods and feeds and thus it can be used as a reference method to determine the extractability of various krill meals. For example, the Soxhlet method may be carried out as below using petroleum ether (boiling point 30–60 °C). Conventional krill meal was prepared as described in US 2008/0274203 (Aker Biomarine ASA, Bruheim et al.) and the low fluoride PPC was prepared according to the present invention.

The neutral lipids are often part of large aggregates in storage tissues, from which they are relatively easily extracted. The polar lipids, on the other hand, are present as constituents of membranes, where they occur in a close association with proteins and polysaccharides, with which they interact, and therefore are not extracted so readily. Furthermore, the phospholipids are relatively tightly bound with hydrophobic proteins and in particular with

the phosphorylated proteins.

The data show that partial hydrolysis of the protein matrix in the preparation of a low fluoride PPC composition as described herein improves the extraction efficiency of total lipid by use of non-polar organic solvents (e.g., for example, supercritical CO<sub>2</sub>, ethanol, and/or petroleum ether).

Briefly, a 10 g sample of either conventional milled krill meal or low fluoride PPC was weighed and placed in a Soxhlet apparatus and then continuously extracted for approximately eight (8) hours using 300 mL petroleum ether. After extraction, the solvent was evaporated at 60 °C under a nitrogen stream. Soxhlet F., "Die gewichtsanalytische bestimmung des milchfettes" *Dingler's Polytech. J.* 232:461–465 (1879).

The results show that the proportion of residual (e.g., un-extracted) lipid was twice as large in the conventional krill meal compared to the low fluoride krill meal. See, Table VI.

Table VI: Lipid Extraction Efficiency Of Low Fluoride Krill Meals

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Source material	Extracted krill oil lipid	Source Material Residual lipid (e.g., de-oiled meal)
Conventional krill meal	79.6%	20.4%
Low fluoride krill meal	88.9%	11.1%

15 Consequently, the lipid extraction methods described herein have provided an unpredictable and surprising result that provides a superior product because of a greatly improved extraction efficiency.

#### Example III

# 20 <u>Determination Of Fluoride Content</u>

This example presents one method of determining fluoride content of krill products as fluoride by chemical analysis using an ion selective electrode.

A low fluoride PPC krill meal was prepared as described herein and extracted in accordance with Example I to create a low fluoride krill oil were analyzed for fluoride content and compared with conventional preparation processes. Briefly, the method disclosed herein removes, in most part, the krill exoskeleton from the krill meal thereby reducing the fluoride content. In contrast, the krill exoskeleton is included in the conventional krill meal thereby having relatively high levels of fluoride. Conventional processes are, for example, described in WO 2002/102394 (Neptune Technologies & Bioresources) and US 2008/0274203 (Aker Biomarine ASA).

The krill meals analyzed for fluoride content were produced by: i) a low fluoride method of present invention; and ii) a whole krill material produced by a conventional process. *See, Table VII*.

#### 5 Table VII: Fluoride Content Comparison To Conventional Processes

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Analyzed Material	Low Fluoride Preparation	Conventional Preparation
Krill meal	200 - 500 ppm	1300 ppm
Krill oil	< 0.5 ppm	~ 3 - 5 ppm

The data demonstrate that by removing the exoskeleton in the process of producing krill meal (e.g., the low fluoride preparation as disclosed herein), the fluoride content of the krill meal and the krill oil produced from the meal have a markedly reduced fluoride content (e.g., 3 – 10 fold reduction).

### Example IV

# Krill Oil Color Comparison

Krill oil has typically a strong red colour arising from the carotenoid astaxanthin present in the oil at levels varying from 50 ppm to 1500 ppm. Color of krill oil can be determined with a LabScan® XE spectrophotometer (Hunter Associates Laboratory, INC. Resbon, VA, USA) and reported in CIELAB colour scales (L\*, a\* and b\* values). Deviation from the red colour of astaxanthin can occur when the krill biomass is processed at high temperature and under conditions that induce oxidation. Typical oxidation induced deviation in krill oil color is an increase in the brownish hue. Brown color in krill oil arises from oxidation of lipids and formation of secondary and tertiary oxidation products with amino residues. This process is also called non-enzymatic browning.

Strecker degradation products and pyrroles are products of non-enzymatic browning that have been characterized in samples of krill oil. For example, polymerization of pyrroles results in formation of brown, melatonin like macromolecules. Furthermore, pyrrole content of krill oil can be determined spectroscopically with absorbance at 570 nm.

Samples of three krill oils will be examined for color. One produced by the method of the present invention, one produced from frozen krill by a method described in WO 2002/102394 (Neptune Technologies & Bioresources) and one extracted from dried krill meal with ethanol alone as described in US 2008/0274203 (Aker Biomarine ASA). It is to be found that krill oil produced by the method of the present invention has the lowest level of

brown color determined spectrophotometrically by using CIELAB colour scales (L\*, a\* and b\* values) and/or the lowest level of pyrroles determined spectroscopically.

#### Example V

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#### Organoleptic Krill Oil Quality Determination

Organoleptic quality of krill oil is conventionally determined by chemical analysis of volatile nitrogenous compounds arising from the decomposition of krill proteins and trimethyl amine oxide (TMAO). Nitrogenous compounds analyzed are total volatile nitrogen (TVN) and trimethylamine (TMA). In simplified terms the level of nitrogenous compounds correlate with the level of spoilage in the raw material i.e. krill biomass used for extraction of the oil.

It has become evident that, in addition to the volatile nitrogenous compounds, a large number of volatile components with distinct odour contribute to the sensory properties of krill oil. Many of the volatile components arise from the oxidation of lipid and proteinaceous compounds of krill biomass. Thus, a method that limits the level of oxidative degradation in the krill biomass, will reduce the amount of volatile components in krill oil.

Assessment of the organoleptic quality of different types of krill oil is to be performed by a panel of trained individuals. The sensory properties to be determined include several pre-defined parameters of smell and taste. It is to be found that the novel krill oil has an improved sensory profile compared to the other oils tested. The other oils to be tested include one extracted from frozen krill by a method described in WO 2002/102394 (Neptune Technologies & Bioresources) and one extracted from dried krill meal with ethanol alone as described in US 2008/0274203 (Aker Biomarine ASA).

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#### Example VI

#### Production Of Low Trimethyl Amine Crustacean Products

This example describes one method to produce low TMA crustacean products using a krill meal material composition. One having ordinary skill in the art, upon reading this specification would understand that this krill meal material composition may have variable fluoride content, including fluoride contents below 0.5 ppm, in addition to the basic components described below. *See, Table VIII*.

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Table VIII: Unextracted Krill Meal Composition

Eicosapentaenoic Acid (EPA)	11 g/100g (11 % w/w)
Docosahexaenoic acid (DHA)	7 g/100g (7 % w/w)
Omega-3 Fatty Acids	22.7 g/100g (22.7% w/w)
Phospholipids (PLs)	45 g/100g (45% w/w)
Trimethylamine (TMA)	44 mg N/100g (0.044% w/w)
Trimethylamine oxide (TMAO)	354 mg N/100g (0.354% w/w)

A krill oil may then be prepared from the krill meal using ethanol extraction as described above that has the basic components described below. See, Table IX.

Table IX. Krill Oil Components After Conventional Ethanol Extraction Of Krill Meal

Parameter	Value
EPA	11.5 g/100g (11.5% w/w)
DHA	6,5 g/100g (6.5% w/w)
Omega-3 Fatty Acides	22,1 g/100g (22.1% w/w)
Phospholipids	44 g/100g (44 % w/w)
Trimethylamine	50 mg N/100g (0.05 % w/w)
Trimethylamineoxide	216 mg N/100g (0.216% w/w)

Alternatively, krill oil was prepared by krill meal extraction at 40 bars and 40°C using supercritical dimethyl ether (SC DME). The DME extract composition was dried on a Rotavapor® and then flushed with nitrogen. The components of the resultant dried composition is listed below. See, Table X.

15 Table X: Krill Oil Components After SC DME Extraction Of Krill Meal

Parameter	Value
EPA	10,4 g/100g (10.4% w/w)
DHA	6,8 g/100g (6.8% w/w)
Omega-3 Fatty Acids	21,7 g/100g (21.7% w/w)
Phospholipids	45,7 g/100g (45.7% w/w)
Trimethyl amine	<1 mg N/100 g (< 0.001% w/w)
Trimethylamine oxide	20 mg N/100 g (0.02% w/w)

These data clearly show that supercritical DME extraction of krill meal compositions result in a preferential 10-100 fold reduction of TMA and TMAO levels.

# Example VII

# Nuclear Magnetic Resonance Phospholipid Profiles Of Low Fluoride Krill Oil

This example presents representative data of the phospholipid composition of low fluoride krill oils prepared by the methods described herein. *See, Table XI.* 

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Table XI: Phospholipids in Low fluoride krill oil analyzed using <sup>31</sup>P NMR. Sample #1 (color; orange)

Phospholipid (PL)		wt% of total PL	g/100g sample
Phosphatidylcholine	PC	79,7	31.1
Alkyl acyl phosphatidylcholine	AAPC	9.9	3.9
Phosphalidylinositol	PI	0.8	0.3
Phosphatidylserine	PS	0.7	0.3
Lysophosphatidylcholine	LPC	2.4	1.0
Lyso alkyl acyl phosphatidylcholine	LAAPC	0.6	0.2
Phosphatidylethanolamine	PE	3.5	1.4
Alkyl acyl phosphatidylethanolamine	AAPE	0,5	0.2
Cardiolipin + N-acylphosphatidylethanolamine	CL/NAPE	1.1	0.4
Lysophosphatidylethanolamine	LPE	0.6	0.2
Lyso alkyl acyl phosphatidylethanolamine	LAAPE	G.2	0.1

Total phospholipid content\*

39.0 g/100g sample 39.5 g/100g solids

n.d. = not detected

\* Sum of the identified phospholipid classes

10 Sample #2 (color; orange)

Phospholipid (PL)		wt% of total PL	g/100g sample
Phosphatidylcholine	PC	66.7	27.0
Alkyl acyl phosphatidylcholine	AAPC	6.9	2.8
Phosphatidylinositol	Pl	0.9	0.4
Phosphatidylserine	PS		n.d.
Lysophosphatidylcholine	LPC	18.9	7.7
Lyso alkyl acyl phosphatidylcholine	LAAPC	0.8	0.3
Phosphatidylethanolamine	PE	1.4	0.6
Alkyl acyl phosphatidylethanolamine**	AAPE	CONTROL CONTROL OF THE PROPERTY OF THE PROPERT	van vir van vir vir vir vir vir vir vir vir vir vir
Cardiolipin + N-acylphosphatidylethanolamine	CL/NAPE	3.0	1.2
Lysophosphatidylethanolamine	LPE	1.2	0.5
Lyso alkyl acyl phosphatidylethanolamine	LAAPE	0.2	0.1

Total phospholipid content\*

40.5 g/100g sample 42.2 g/100g solids

n.d. = not detected

\* Sum of the identified phospholipid classes

# Sample #3 (color; orange)

Phospholipid (PL)		wt% of total PL	g/100g sample
Phosphatidylcholine	PC	72.3	31.1
Alkyl acyl phosphatidylcholine	AAPC	6.1	2.6
Phosphatidylinositol	PI	0,3	0.1
Phosphatidylserine	PS	0.2	0.1
Lysophosphatidylcholine	LPC	16.1	6.9
Lyso alkyl acyl phosphatidylcholine	LAAPC	0.8	0.3
Phosphatidylethanolamine	PE	1.8	0.8
Alkyl acyl phosphatidylethanolamine**	AAPE	The house of the state of the s	The second secon
Cardiolipin + N-acylphosphatidylethanolamine	CL/NAPE	1.2	0.5
Lysophosphatidylethanolamine	LPE	1.1	0.5
Lyso alkyl acyl phosphatidylethanolamine	LAAPE		n.d.

Total phospholipid content\* 43.0 g/100g sample 45.1 g/100g solids

5

# Sample #4 (color; orange)

Phospholipid (PL)		wt% of total PL	g/100g sample
Phosphatidylcholine	PC	77.4	39.5
Alkyl acyl phosphatidylcholine	AAPC	8.9	4.6
Phosphatidylinositol	Pl	0.9	0.5
Phosphatidylserine	PS	0.4	0,2
Lysophosphatidylcholine	LPC	5.5	2.8
Lyso alkyl acyl phosphatidylcholine	LAAPC	0.6	0.3
Phosphatidylethanolamine	PE	2.6	1.3
Alkyl acyl phosphatidylethanolamine**	AAPE	1.3	0.7
Cardiolipin + N-acylphosphatidylethanolamine	CL/NAPE	1.8	0.9
Lysophosphatidylethanolamine	LPE	0.5	0.3
Lyso alkyl acyl phosphatidylethanolamine	LAAPE	0.2	0.1

Total phospholipid content\* 51.1 g/100g sample 52.8 g/100g solids

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n.d. = not detected

<sup>\*</sup> Sum of the identified phospholipid classes

n.d. = not detected

<sup>\*</sup> Sum of the identified phospholipid classes

<sup>\*\*</sup> May contain some glycerophosphocholine (GPC)

Sample #5 (color; orange)

Phospholipid (PL)		wt% of total PL	g/100g sample
Phosphatidylcholine	PC	65.5	26.8
Alkyl acyl phosphatidylcholine	AAPC	9.4	3.9
Phosphatidylinositol	PI	1,6	0.6
Phosphatidylserine	PS	0.7	0.3
Lysophosphatidylcholine	LPC	10.1	4.2
Lyso alkyl acyl phosphatidylcholine	LAAPC	1,3	0.5
Phosphatidylethanolamine	PE	4.9	2.0
Alkyl acyl phosphatidylethanolamine	AAPE	2.1	0.9
Cardiolipin + N-acylphosphatidylethanolamine	CL/NAPE	2.8	1.2
Lysophosphatidylethanolamine	LPE	1.3	0.5
Lyso alkyl acyl phosphatidylethanolamine	LAAPE	0.3	0.1

Total phospholipid content*	41.0 g/100g sample
	43.0 a/100a solids

5 These data are consistent with those obtained using traditional column chromatography techniques shown in Example I.

#### Example VIII

#### Lipid Compositional Analysis Of Low Fluoride PPC Material

The example presents data showing the lipid compositional analysis of a low fluoride phospholipid-protein complex composition created by the methods described herein.

Consequently, it would be expected that the fluoride content of the compositions described below are less than 500 ppm.

The PPC comprises approximately 46.7 g/100 g (e.g.,  $\sim$  47%) total fat, 11.8 g/100 g (e.g.,  $\sim$  12%) eicosapentaenoic Acid (EPA) and 6.7 g/ 100 g (e.g.,  $\sim$ 7%) docosahexaenoic acid (DHA). The total lipid content of the PPC total fat was approximately 87.7 % (w/w) and comprises between approximately 115 - 260 mg/kg astaxanthin and between approximately 35.2% - 46.7% unextracted oil.

20

n.d. = not detected

Sum of the identified phospholipid classes

Table XII: Low Fluoride Krill PPC Fat: Neutral Lipid Content (45.2% w/w of total fat):

Sample Number 1MG

Components	% (w/w)
	neutral
	lipid
Triacylglycerol	38
Diacylglycerol	1.7
Monoacylglycerol	< 1
Free fatty acids	2.2
Cholesterol	2.4
Cholesterol Esters	< 0.5

5 Table XIII: Low Fluoride Krill PPC Fat: Neutral Lipid Content (46.6% w/w of total fat):

Sample Number 2MG

Components	% (w/w)
	neutral
	lipid
Triacylglycerol	41
Diacylglycerol	1.5
Monoacylglycerol	< 1
Free fatty acids	1.6
Cholesterol	1.8
Cholesterol Esters	0.6

Table IXV: Low Fluoride Krill PPC Neutral Lipids: Fatty Acid Content (49.7% w/w of

neutral lipids): Sample Number 1MG

Components	% (w/w)
	neutral
	lipid
Saturated	27.4
Monoenoic	21.9
N-6 Polyunsaturated	1.8
N-3 Polyunsaturated	22.7
Total	74.4

Table XV: Low Fluoride Krill PPC Neutral Lipids: Fatty Acid Content (46.7% w/w of

neutral lipid): Sample Number 2MG

Components	% (w/w)
	neutral
	lipid
Saturated	29.2
Monoenoic	21.6
N-6 Polyunsaturated	2.1
N-3 Polyunsaturated	23.3
Total	76.9

Table XVI: Low Fluoride Krill PPC Polar Lipid Content (42.6% w/w of total lipids): Sample Number 1MG

Components	% (w/w)
	polar lipid
Phosphatidylethanolamine	3.4
Phosphatidylinositol	< 1
Phosphatidylserine	< 1
Phosphatidylcholine	37
Lyso Phosphatidylcholine	2.3

# 5 Table XVII: Low Fluoride Krill PPC Polar Lipid Content (42.8% w/w of total lipids): Sample Number 2MG

Components	% (w/w)
	polar lipid
Phosphatidylethanolamine	2.5
Phosphatidylinositol	< 1
Phosphatidylserine	< 1
Phosphatidylcholine	39
Lyso Phosphatidylcholine	1.8

#### Example IX

# Lipid Compositional Analysis Of Low Fluoride De-Oiled PPC Material

The example presents data showing the lipid compositional analysis of a low fluoride de-oiled phospholipid-protein complex composition created by the methods described herein. Consequently, it would be expected that the fluoride content of the compositions described below are less than 500 ppm. The de-oiled PPC comprises approximately 35 g/ 100 g (e.g.,  $\sim$  35%) total fat, 16.6 g/100 g (e.g.,  $\sim$  17%) eicosapentaenoic Acid (EPA) and 10.0 g/ 100 g (e.g.,  $\sim$ 10%) docosahexaenoic acid (DHA). The total lipid content of the de-oiled PPC total fat was approximately 87.7 % (w/w) and comprises approximately 115 mg/kg astaxanthin and approximately 35.2% unextracted oil.

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Table XVIII: Low Fluoride Krill De-Oiled PPC Fat: Neutral Lipid Content (20.1% w/w of

total fat): Sample Number 3MG

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Components	% (w/w)
	Neutral
	Lipid
Triacylglycerol	17
Diacylglycerol	0.6
Monoacylglycerol	< 1
Free fatty acids	1.1
Cholesterol	1.3
Cholesterol Esters	< 0.5

5 Table IXX: Low Fluoride Krill De-Oiled PPC Neutral Lipids: Fatty Acid Content (35.2% w/w of neutral lipids): Sample Number 3MG

Components	% (w/w)
	Neutral lipid
Monoenoic	13.9
N-6 Polyunsaturated	2.1
N-3 Polyunsaturated	31.2

10 Table XX: Low Fluoride Krill PPC De-Oiled Polar Lipid Content (68.9% w/w of total fat): Sample Number 3MG

Components	% (w/w)
	polar lipid
Phosphatidylethanolamine	4.2
Phosphatidylinositol	< 1
Phosphatidylserine	< 1
Phosphatidylcholine	62
Lyso Phosphatidylcholine	2.2

# Example X

15 Compositional Analysis Of PPC/Protein Hydrolysate Mixtures

The example presents data showing the lipid compositional analysis of a low fluoride phospholipid-protein complex mixed with a protein hydrolysate composition created by the methods described herein in an approximate 60/40 ratio. It would be expected that the fluoride content of the compositions described below are less than 500 ppm. The mixture comprises between approximately 28-30 g/100 g (e.g., ~ 30%) total fat, approximately 98

mg/kg astaxantine esters, approximately less than 1 mg/kg astaxanthine, a peroxide level of less than 0.1 %;(mEq/kg) and/or an ananiside level of less than 0.1 % (w/w).

Table XXI: Low Fluoride PPC/Protein Mixture Fat: Neutral Lipid Content (28% w/w of total fat)

Components	% (w/w)
	Neutral
	Lipid
Triacylglycerol	34
Diacylglycerol	1.1
Monoacylglycerol	< 1
Free fatty acids	1.0
Cholesterol	1.9
Cholesterol Esters	< 0.5

Table XXII: Low Fluoride PPC/Protein Mixture Neutral Lipids: Fatty Acid Content

y Acid Comen
% (w/w)
Neutral
lipid
25.1
19.2
2.0
24.9

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Table XXIII: Low Fluoride PPC/Protein Mixture Polar Lipid Content

Components	% (w/w)
	polar lipid
Phosphatidylethanolamine	5.0
Phosphatidylinositol	< 1
Phosphatidylserine	< 1
Phosphatidylcholine	41
Lyso Phosphatidylcholine	1.4

#### **CLAIMS**

We claim:

5 1. A crustacean oil composition comprising phospholipids and less than approximately 0.5 ppm fluoride.

2. The crustacean oil composition of Claim 1, further comprising less than approximately 0.001% (w/w) trimethyl amine.

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- 3. The crustacean oil composition of Claim 1, further comprising less than approximately 0.02% (w/w) trimethyl amine oxide.
- 4. The crustacean oil composition of Claim 1, wherein said phospholipids are between
   approximately 39-52 wt%, wherein said phospholipids comprise at least approximately 65% phosphatidylcholine and at least approximately 2.4 wt% lysophasphatidylcholine.
  - 5. The crustacean oil composition of Claim 1, further comprising triglycerides, neutral lipids, approximately 20 26 wt% Omega-3 fatty acids, and at least approximately 0.8 wt% free fatty acids.
    - 6. The crustacean oil composition of Claim 1, wherein said oil is krill oil.
- 7. A crustacean phospholipid-peptide complex (PPC) composition comprising a matrix of
   25 hydrolyzed protein, phospholipids and between approximately 300-500 ppm fluoride.
  - 8. The crustacean PPC composition of Claim 7, wherein said phospholipids are at least 40 wt%.
- The crustacean PPC composition of Claim 7, further comprising approximately 0.044%
   (w/w) trimethyl amine and approximately 0.354% (w/w) trimethyl amine oxide.
  - 10. The crustacean PPC composition of Claim 7, further comprising at least 40% (w/w) triglycerides.

11. A crustacean de-oiled phospholipid-peptide complex (PPC) composition comprising a matrix of hydrolyzed protein, between approximately 200-500 ppm fluoride, approximately 35% total fat, approximately 16.6% eicosapentaenoic acid, approximately 10.0% docosahexaenoic acid and at least 0.1 wt% free fatty acids.

- 12. The crustacean de-oiled PPC composition of Claim 11, wherein said total fat comprises less than 20% triglycerides and approximately 69% other lipid components.
- 10 13. The crustacean de-oiled PPC composition of Claim 12, wherein said total fat comprises approximately 35.2% fatty acids, wherein approximately 30 wt% of said fatty acids are n-3 fatty acids.
- 14. The crustacean de-oiled PPC composition of Claim 12, wherein said total lipids further15 comprise at least 68% phospholipids.
  - 15. The crustacean de-oiled PPC composition of Claim 12, further comprising approximately 2.2% lysophosphatidyl choline.
- 20 16. The crustacean de-oiled PPC composition of Claim 12, further comprising approximately 115 mg/kg astaxanthin.
  - 17. A dried protein hydrolysate composition comprising approximately 70 80% protein, approximately 1.5 3.0% lipids, and approximately 5 7 % ash.

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- 18. A method for creating low fluoride crustacean compositions, comprising:
  - a) disintegrating a crustacean catch into a material having a particle size ranging between approximately 1-25 millimeters; and
- b) separating said disintegrated crustacean material into a phospholipid-peptide complex (PPC) composition subfraction, wherein said subfraction comprises a fluoride content of less than 500 ppm.

19. The method according to Claim 18, wherein said method further comprises extracting said PPC composition subfraction with a fluid comprising a solvent wherein a low fluoride oil is created, said oil having a fluoride content of less than 0.5 ppm.

- 5 20. The method according to Claim 19, wherein said extracting further creates a low trimethyl amine/trimethyl amine oxide oil, wherein said trimethyl amine is less than approximately 0.001% (w/w) and said trimethyl amine oxide is less than approximately 0.02% (w/w).
- 10 21. The method according to Claim 18, wherein said separating is performed without emulsification.
  - 22. The method according to Claim 18, wherein said solvent comprises a non-polar solvent.
- 15 23. The method according to Claim 18, wherein said solvent comprises at least one polar solvent.
  - 24. The method according to Claim 18, wherein said solvent comprises said non-polar solvent and said at least one polar solvent.

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- 25. The method according to Claim 22, wherein said non-polar solvent is selected from at least one of the group consisting of supercritical carbon dioxide and supercritical dimethyl ether.
- 26. The method according to Claim 23, wherein said polar solvent is selected from at least one of the group consisting of ethanol and acetone.
  - 27. The method according to Claim 18, wherein said method further comprises hydrolyzing said crustacean material before said separating.
  - 28. The method according to Claim 22, wherein said extracting further creates a de-oiled PPC composition.

29. The method according to Claim 23, wherein said polar solvent separates a phospholipid composition and a protein hydrolysate composition from said de-oiled PPC composition.

- 30. The method according to Claim 19, wherein said extracting comprises less than ten bours.
  - 31. The method according to Claim 19, wherein said extracting comprises less than five hours.
- 10 32. The method according to Claim 19, wherein said extracting comprises less than two hours.
  - 33. The method according to Claim 18, wherein said crustacean material is krill material.
- 15 34. The method according to Claim 18, wherein said separating comprises a centrifugal force of between approximately 1,000 1,800 g.
  - 35. The method according to Claim 18, wherein said separating comprises a centrifugal force of between approximately 5,000 10,000 g.
  - 36. A composition comprising a mixture of a low fluoride crustacean PPC and a low fluoride de-oiled PPC, wherein said fluoride level ranges between approximately 200 500 ppm.
  - 37. The composition of Claim 36, wherein said crustacean PPC is krill PPC.

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- 38. The composition of Claim 36 wherein said crustacean de-oiled PPC is krill de-oiled PPC.
- 39. The composition of Claim 36, wherein said crustacean PPC and crustacean de-oiled PPC are in a 1:1 ratio.
- 30 40. The composition of Claim 36, wherein said mixture comprises a milled fine powder.
  - 41. The composition of Claim 40, wherein said powder comprises a particle size of approximately 250  $\mu m$ .

42. The composition of Claim 36, wherein said composition comprises a peroxide level of less than 0.1 % (mEq/kg).

- 43. The composition of Claim 36, wherein said composition comprises ananiside level of less than 0.1 % (w/w).
  - 44. The composition of Claim 36, wherein said composition further comprises microencapsulated polyunsaturated Omega-3 fatty acids.
- 10 45. The composition of Claim 36, wherein said composition further comprises zinc oxide.
  - 46. The composition of Claim 36, wherein said composition further comprises marine peptides.
- 15 47. The composition of Claim 36, wherein said composition further comprises at least one supplemental amino acid.
  - 48. A method, comprising formulating a composition comprising a low fluoride crustacean PPC and a low fluoride crustacean de-oiled PPC, wherein said fluoride level ranges between approximately 200 500 ppm.
    - 49. The method of Claim 48, wherein said method further comprises milling said composition into a powder.

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- 25 50. The method of Claim 48, wherein said method further comprises tabletting said composition into a tablet.
  - 51. The method of Claim 48, wherein said method further comprises encapsulating said composition into a capsule.
  - 52. The method of Claim 48, wherein said method further comprises mixing said powder with a food product.

53. The method of Claim 48, wherein said formulating further comprises microencapsulated polyunsaturated Omega-3 fatty acids.

54. The method of Claim 48, wherein said formulating further comprises zinc oxide.

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- 55. The method of Claim 48, wherein said formulating further comprises marine peptides.
- 56. The method of Claim 48, wherein said formulating further comprises at least one supplemental amino acid.

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- 57. A composition comprising a mixture of a low fluoride crustacean PPC and a crustacean protein hydrolysate, wherein said fluoride level ranges between approximately 200 500 ppm.
- 15 58. The composition of Claim 57, wherein said crustacean PPC is krill PPC.
  - 59. The composition of Claim 57, wherein said crustacean protein hydrolysate is a krill protein hydrolysate.
- 20 60. The composition of Claim 57, wherein said crustacean PPC and said crustacean protein hydrolysate are in a 1:1 ratio.
  - 61. The composition of Claim 57, wherein said mixture comprises a milled fine powder.
- 25 62. The composition of Claim 61, wherein said powder comprises a particle size of approximately 250 μm.
  - 63. The composition of Claim 57, wherein said composition comprises a peroxide level of less than 0.1 % (mEq/kg).

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64. The composition of Claim 57, where said composition comprises ananiside level of less than 0.1 wt%.

65. The composition of Claim 57, wherein said composition further comprises microencapsulated polyunsaturated Omega-3 fatty acids.

- 66. The composition of Claim 57, wherein said composition further comprises zinc oxide.
  - 67. The composition of Claim 57, wherein said composition further comprises marine peptides.
- 68. The composition of Claim 57, wherein said composition further comprises at least one supplemental amino acid.
  - 69. A method, comprising formulating a composition comprising a low fluoride crustacean PPC and a crustacean protein hydrolysate, wherein said fluoride level ranges between approximately 200 500 ppm.

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- 70. The method of Claim 69, wherein said method further comprises milling said composition into a powder.
- 71. The method of Claim 69, wherein said method further comprises tabletting said20 composition into a tablet.
  - 72. The method of Claim 69, wherein said method further comprises encapsulating said composition into a capsule.
- 25 73. The method of Claim 69, wherein said method further comprises mixing said powder with a food product.
  - 74. The method of Claim 69, wherein said formulating further comprises microencapsulated polyunsaturated Omega-3 fatty acids.

- 75. The method of Claim 69, wherein said formulating further comprises zinc oxide.
- 76. The method of Claim 69, wherein said formulating further comprises marine peptides.

77. The method of Claim 69, wherein said formulating further comprises at least one supplemental amino acid.

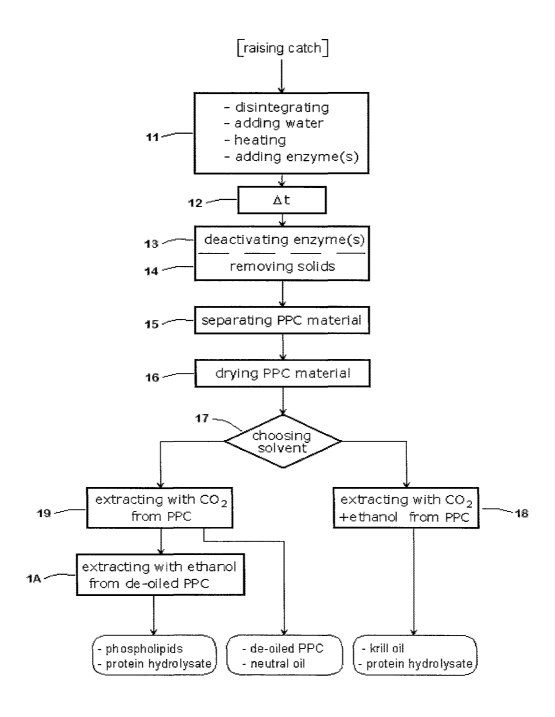


Fig. 1

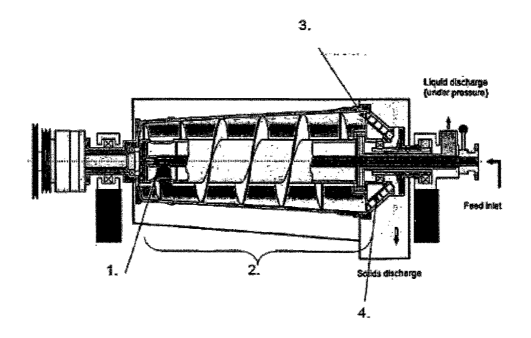


Fig. 2

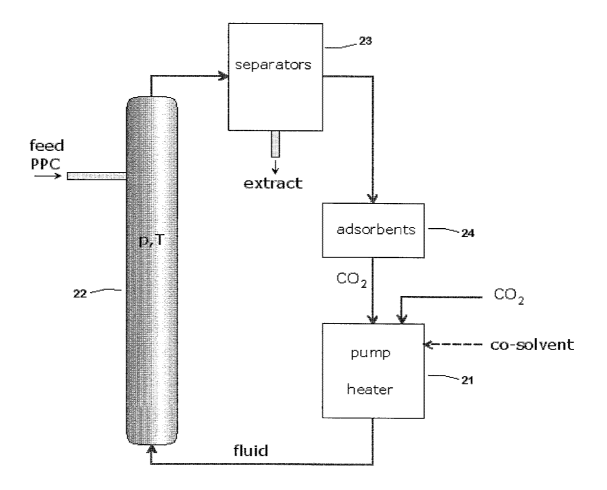


Fig. 3

WO 2013/102792 PCT/IB2012/003004

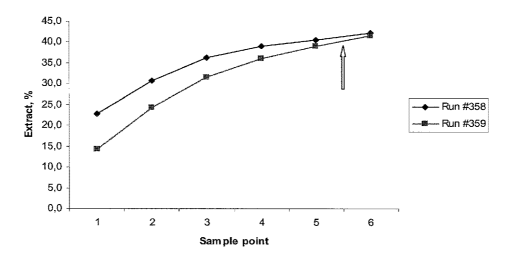


FIG. 4

#### **PATENT COOPERATION TREATY**

## **PCT**

#### INTERNATIONAL SEARCH REPORT

(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference	FOR FURTHER	see Form PCT/ISA/220
AKBM34345WO	ACTION	as well as, where applicable, item 5 below.
International application No.	International filing date (day/month/	(Earliest) Priority Date (day/month/year)
PCT/IB2016/000208	10 February 2016 (10-02-2016)	11 February 2015 (11-02-2015)
Applicant		
AKER BIOMARINE ANTARCTIC AS		
This international search report has been according to Article 18. A copy is being tra		ng Authority and is transmitted to the applicant
This international search report consists o	f a total of sheets	3.
X It is also accompanied by	a copy of each prior art document cite	ed in this report.
Basis of the report     a. With regard to the language, the i	nternational agersh was carried out a	n the basis of
	pplication in the language in which it	
a translation of the	e international application into	, which is the language al search (Rules 12.3(a) and 23.1(b))
b. This international search r	eport has been established taking int	account the rectification of an obvious mistake
	o this Authority under Rule 91 (Rule 4 htide and/or amino acid sequence o	ვ.ი <i>ხი</i> (a)). isclosed in the international application, see Box No. t.
	·	
2. Certain claims were fou	nd unsearchable (See Box No. II)	
3. Unity of invention is lact	ding (see Box No III)	
4. With regard to the title,		
X the text is approved as su	bmitted by the applicant	
the text has been establish	ned by this Authority to read as follow	s:
5. With regard to the abstract,		
X the text is approved as suf	• • • •	11 9 9 15 1 15 Al 104 W. 15 1
		uthority as it appears in Box No. IV. The applicant nal search report, submit comments to this Authority
6. With regard to the <b>drawings,</b>		
a. the figure of the <b>drawings</b> to be po	ublished with the abstract is Figure No	o,
as suggested by t	he applicant	
	s Authority, because the applicant fail	
	Authority, because this figure better	characterizes the invention
b none of the figures is to be	published with the abstract	

#### INTERNATIONAL SEARCH REPORT

International application No

PCT/IB2016/000208

a. classification of subject matter INV. A23J7/00 A61K3

C11C1/02

A61K35/612 C11C1/08

C11B1/10

C11B3/00

C11C1/00

ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

#### **B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

A61K C11B A23J C11C

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal, WPI Data

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
(	WO 2014/013335 A2 (HOEM NILS [NO]; TILSETH SNORRE [NO]; AKER BIOMARINE AS [NO]) 23 January 2014 (2014-01-23) claims; examples	1-56
(	US 2014/370115 A1 (HOEM NILS [NO] ET AL) 18 December 2014 (2014-12-18) claims; examples 2-4	1-56
(	WO 2012/139588 A2 (TRIPLENINE PHARMA AS [DK]; SOERENSEN HANS OTTO [DK]; JENSEN NILS CHRIS) 18 October 2012 (2012-10-18) claims; example 2	1-56
(	CN 102 746 941 B (UNIV SHANDONG NORMAL) 15 January 2014 (2014-01-15) claims; example 2 	1-56

Х	Further documents are listed in the	continuation of Box C.
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X See patent family annex.

- Special categories of cited documents :
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- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
- "&" document member of the same patent family

Date of mailing of the international search report Date of the actual completion of the international search 6 May 2016 13/05/2016

Name and mailing address of the ISA/

European Patent Office, P.B. 5818 Patentlaan 2 NL · 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016

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Vernier, Frédéric

1

#### INTERNATIONAL SEARCH REPORT

International application No
PCT/IB2016/000208

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Refevant to claim No.
(	JP 2 909508 B2 (TAIYO FISHERY CO LTD) 23 June 1999 (1999-06-23) claims; examples 	1-56

1

#### INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No
PCT/IB2016/000208

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
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CN 102746941 B	15-01-2014	NONE	
JP 2909508 B2	23-06-1999	JP 2909508 B2 JP H02215351 A	23-06-1999 28-08-1990

# Information on Search Strategy - Pilot phase (see OJ 2015, A86) The type of information contained in this sheet may change during the pilot for Improving the usefulness of this new service.

Application Number

PCT/IB2016/000208

TITLE: LIPID EXTRACTION PROCESSES

APPLICANT: AKER BIOMARINE ANTARCTIC AS

IPC CLASSIFICATION: A23J7/00, A61K35/612, C11B1/10, C11B3/00, C11C1/00,

C11C1/02, C11C1/08

EXAMINER: Vernier, Frédéric

CONSULTED DATABASES: EPODOC, WPI, INET, DOSYS

CLASSIFICATION SYMBOLS DEFINING EXTENT OF THE SEARCH:

IPC:

CPC: A23L1/3006, A23J7/00, A61K35/612, C11B1/10, C11B3/006, C11C1/007, C11C1/02, C11C1/08

FI/F-TERMS:

KEYWORDS OR OTHER ELEMENTS FEATURING THE INVENTION: Process for the extraction of lipids from marine biomass, in particular krill, using ethanol. Fractionation into an astaxanthin-rich fraction and a phospholipid-rich fraction.

#### PATENT COOPERATION TREATY



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Jones, J. Mitchell CASIMIR JONES, S.C.

CASIMIR JUNES, S.C.
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INVITATION TO PAY ADDITIONAL FEES AND, WHERE APPLICABLE, PROTEST FEE

M	TATS-UNIS D'AMERIQUE	(PCT Article 17(3)(a) and Rule 40.1 and 40.2(e))
		Date of mailing (day/month/year) 15 June 2016 (15-06-2016)
	cant's or agent's file reference	PAYMENT DUE within ONE MONTH from the above date of mailing
	national application No.	International filing date (day/month/year) 10 February 2016 (10-02-2016)
Appli	cant CASIMIR JONES 5:0	***
AK	ER BIOMARINE ANTARCTIC AS	
1.	This International Searching Authority	
	(i) considers that there are 3 (no by the claims indicated on an extra sheet:	umber of) inventions claimed in the international application covered
	(ii) therefore considers that <b>the international application o</b> (Rules 13.1, 13.2 and 13.3) for the reasons indicated on	loes not comply with the requirements of unity of invention an extra sheet:
	(iii) X has carried out a partial international search (see A on those parts of the international application which related	
	see extra sheet	
	(iv) will establish the international search report on the other to which, additional fees are paid.	parts of the international application only if, and to the extent
2.	Consequently, the applicant is hereby invited to pay, within	the time limit indicated above, the amount indicated below:
	EUR 1.875,00 x 2	= EUR 3.750,00
	Fee per additional invention number of additional	inventions currency/total amount of additional lees
3.	The applicant is informed that, according to Rule 40.2(c), the i.e., a reasoned statement to the effect that the international a or that the amount of the required additional fee is excessive, where the applicant pays additional fees under protest, the apto pay a protest fee (Rule 40.2(e)) in the amount of where the applicant has not, within the time limit indicated about to have been made and the International Searching Authorities.	pplication complies with the requirement of unity of invention where applicable, subject to the payment of a protest fee.  plicant is hereby invited, within the time limit indicated above,  EUR 875,00  (currency/amount)  ve, paid the required protest fee, the protest will be considered
4.	Claim(s) Nos.  Article 17(2)(b) because of defects under Article 17(2)(a)	have been found to be unsearchable under and therefore have not been included with any invention.
Name	and mailing address of the International Searching Authority	Authorized officer

European Patent Office, P.B. 5818 Patentlaan 2 NL-2280 HV Rijswijk Tel. (+31-70) 340-2040 Fax: (+31-70) 340-3016

BENINCA CORDES, Carmelita Tel: +49 (0)89 2399-2806

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. claims: 1-30, 46-52(completely); 54-59(partially)

i/ A desalted krill lipid composition, a krill phospholipid concentrate, a krill lipid composition (with a phospholipid content 30-50% or 50-85% and a triglyceride content 32-52%, 5-35% and 26-46% and at least one of the properties mentioned in the respective claims); and ii/ an oral delivery vehicle comprising said compositions, a functional food comprising said compositions. a nutritional

functional food comprising said compositions, a functional food comprising said compositions, a nutritional supplement, dietary supplement or medical food comprising said compositions, a composition comprising said compositions and a second nutraceutical ingredient and a composition comprising said compositions and a pharmaceutically acceptable carrier (as far as as the ii/refer back to the claims 1-15, 16-30 and 46).

2. claims: 31-45(completely); 54-59(partially)

i/ A krill lipid composition (with a phospholipid content 2-20% and a triglyceride content 70-95%, and at least one of the properties mentioned in the respective claims); and ii/ an oral delivery vehicle comprising said compositions, a functional food comprising said compositions, a nutritional supplement, dietary supplement or medical food comprising said compositions, a composition comprising said compositions and a second nutraceutical ingredient and a composition comprising said composition and a pharmaceutically acceptable carrier (as far as as the ii/ refer back to the claims 31-45).

3. claims: 53(completely); 54-59(partially)

i/ A concentrated krill astaxanthin composition comprising 400-4000 ppm astaxanthin esters and more than 98% krill neutral lipids;

ii/ an oral delivery vehicle comprising said composition, a functional food comprising said composition, a nutritional supplement, dietary supplement or medical food comprising said composition, a composition comprising said composition and a second nutraceutical ingredient and a composition comprising said composition and a pharmaceutically acceptable carrier (as far as as the ii/ refer back to the claim 53).

In the first invention, the Special Technical Feature (STF) is the

combination of the concentrations of phospholipids and triglycerides as well as at least one of the properties. This STF solves the technical problem of providing a krill oil composition that has a high quality with respect to purity, smell and taste (see also page 7 of the present invention).

In the second invention, a possible STF is the combination of the (completely different from those of the 1st invention) concentrations of phospholipids and triglycerides as well as at least one of the properties as in the claims of the 2nd invention. This STF does not appear to solve any technical problem.

In the third invention, a possible STF is the combination of the concentration of the astaxanthin esters and krill neutral lipids (no mention of any additional technical features / properties). Again, this STF does not appear to solve any technical problem.

Since the three aforementioned inventions do not have the same STF, the present application lacks unity.

Only the subject-matter of the 1st invention is searched.

# Annex to Form PCT/ISA/206 COMMUNICATION RELATING TO THE RESULTS OF THE PARTIAL INTERNATIONAL SEARCH

International Application No PCT/IB2016/000326

1.The present communication is an Annex to the invitation to pay additional fees (Form PCT/ISA/206). It shows the results of the international search established on the parts of the international application which relate to the invention first mentioned in claims Nos.:

see 'Invitation to pay additional fees' 2. This communication is not the international search report which will be established according to Article 18 and Rule 43.

3.If the applicant does not pay any additional search fees, the information appearing in this communication will be considered as the result of the international search and will be included as such in the international search report.

4.If the applicant pays additional fees, the international search report will contain both the information appearing in this communication and the results of the international search on other parts of the international application for which such fees will have been paid.

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	DATABASE FSTA [Online] INTERNATIONAL FOOD INFORMATION SERVICE (IFIS), FRANKFURT-MAIN, DE; SHIBATA N: "Effect of fishing season on lipid content and composition of Antarctic krill. (translated)", XP002758417, Database accession no. FS-1985-04-R-0091 abstract	1-30, 46-52, 54-59
Х	WO 2010/097701 A1 (AKER BIOMARINE ASA [NO]; TILSETH SNORRE [NO]) 2 September 2010 (2010-09-02) * page 1, lines 5-8; claims 1-17 *	1-30, 46-52, 54-59
х	US 2010/226977 A1 (TILSETH SNORRE [NO]) 9 September 2010 (2010-09-09) * claims 1-15 *	1-30, 46-52, 54-59
X	US 2014/274968 A1 (BERGE KJETIL [NO] ET AL) 18 September 2014 (2014-09-18)  * claims 1-20 *	1-30, 46-52, 54-59

° Special categories of cited documents :

<sup>&</sup>quot;A" document defining the general state of the art which is not considered to be of particular relevance

<sup>&#</sup>x27;E" earlier document but published on or after the international filing date

<sup>&</sup>quot;L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

<sup>&</sup>quot;O" document referring to an oral disclosure, use, exhibition or other means

<sup>°</sup>P° document published prior to the international filing date but later than the priority date claimed

<sup>&</sup>quot;T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

<sup>&</sup>quot;X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

<sup>&</sup>quot;Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

<sup>&</sup>quot;&" document member of the same patent family

### **Patent Family Annex**

Information on patent family members

International Application No
PCT/IB2016/000326

	ent document in search report		Publication date	Patent family Publication member(s) date	
WO	2010097701	A1	02-09-2010	NONE	
US	2010226977	A1	09-09-2010	NONE	
US	2014274968	A1	18-09-2014	AU 2014229540 A1 08-10-2015 US 2014274968 A1 18-09-2014 WO 2014140873 A2 18-09-2014	

# **Information on Search Strategy** - Pilot phase (see OJ 2015, A86) The type of information contained in this sheet may change during the pilot for improving the usefulness of this new service.

**Application Number** 

PCT/IB2016/000326

TITLE: LIPID COMPOSITIONS

APPLICANT: AKER BIOMARINE ANTARCTIC AS

IPC CLASSIFICATION: C11B1/10, A23K10/22, A23K20/158

EXAMINER: Georgopoulos, N

CONSULTED DATABASES: DOSYS, EPODOC, WPI, TXPJPEA, NPL, XPESP, FSTA, BIOSIS,

MEDLINE

CLASSIFICATION SYMBOLS DEFINING EXTENT OF THE SEARCH:

IPC:

CPC: A23L1/3008, C11B1/10, A23K10/22, A23K20/158

FI/F-TERMS:

KEYWORDS OR OTHER ELEMENTS FEATURING THE INVENTION: Krill compositions and products (food products, feed products, nutraceutical products, etc.) containing the same.

Electronic Acknowledgement Receipt			
EFS ID:	26338763		
Application Number:	15180431		
International Application Number:			
Confirmation Number:	2763		
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS		
First Named Inventor/Applicant Name:	Inge Bruheim		
Customer Number:	72960		
Filer:	John Mitchell Jones/Mallory Checkett		
Filer Authorized By:	John Mitchell Jones		
Attorney Docket Number:	AKBM-14409/US-12/CON		
Receipt Date:	13-JUL-2016		
Filing Date:	13-JUN-2016		
Time Stamp:	17:04:11		
Application Type:	Utility under 35 USC 111(a)		

## **Payment information:**

Submitted with Payment	no
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## File Listing:

Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1	Transmittal Letter	14409US12CON_IDS_Letter_7- 13-2016.pdf	81994 	no	1

Warnings: RIMFROST EXHIBIT 1111 Page 0445

Information:					
			1036089		
2	Information Disclosure Statement (IDS) Form (SB08)	14409US12CON_IDS_7-13-201 6.pdf	f0097c951af962b98a26731ab9b962ed567 d1e25	no	5
Warnings:					
Information:					
			853239		
3	Foreign Reference	WO2014013335A2.pdf	3e490302d63906df9fc52606012318173c35 c450	no	26
Warnings:	-				
Information:					
			326292		
4	Foreign Reference	CN102746941B.pdf	6dc7f0e11f95489f6ce0f024dcde7e92e97e6 11c	no	6
Warnings:	<u> </u>				
Information:					
			345403		
5	Foreign Reference	JP2909508.pdf	46b28b22bd7163560ec1610f83d98a24369 c890f	no	7
Warnings:					
Information:					
			2949290		
6	Foreign Reference	WO2010097701.pdf	78c8095a597e883a8a6045da7ccf3f4cacb7 b23d	no	71
Warnings:	-				
Information:					
			3225439		
7	Foreign Reference	WO2013102792.pdf	0591d9ba8291dcb2418e73f5917754e8607 06089	no	60
Warnings:	-				<u> </u>
Information:					
			150964		
8	Other Reference-Patent/App/Search documents	ISR_34345_PCTIB2016000208.  pdf  5ef07fa22609ea10dfee4c72980008b61e0df f52		no	5
Warnings:	+				
Information:					

			194408		
9	Other Reference-Patent/App/Search documents	Partial_ISR_34344_PCTIB20160 00326.pdf	3593f5f7fe5ef309b27fbca9fb81acd7c40e0 7c6	no	6
Warnings:	-				
Information	<b>:</b>				
			32615		
10	Non Patent Literature	FSTA_Shibata_abstract.pdf	e29068a81bef6233f4d9abe88f6c9299b776 4a91	no	1
Warnings:					
Information	1				
		StatemenofGrounds_33382AU	477111		
11	Other Reference-Patent/App/Search documents	_Rimfrost_6-10-2016_USPTO. pdf	a4061e996de9c51044fea80ccb522adb0f33 7134	no	21
Warnings:					
Information	•				
		Total Files Size (in bytes)	96	72844	

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

#### 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.

#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: Inge Bruheim Confirmation: 2763 Serial No.: 15/180,431 Group No.: 1672

Filing Date: 13-Jun-2016 Examiner: CUTLIFF, YATE

Entitled: BIOEFFECTIVE KRILL OIL COMPOSITIONS

#### INFORMATION DISCLOSURE STATEMENT LETTER

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

#### **Examiner Cutliff:**

The citations listed in the attached **IDS Form SB08A** 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.

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.

This Information Disclosure Statement is being filed under 37 C.F.R. §1.97 (b)(3) before the mailing of a first Office Action on the merits. Therefore, applicant holds that no fee is due.

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-12/CON.

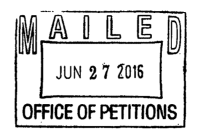
Dated: July 13, 2016 /J. Mitchell Jones/

J. Mitchell Jones Registration No. 44174 CASIMIR JONES, S.C. 2275 Deming Way, Suite 310 Middleton, WI 53562 608.662.1277





Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON WI 53562



Commissioner for Patents United States Patent and Trademark Office P.O. Box 1450 Alexandria, VA 22313-1450 www.uspto.gov

Doc Code: TRACK1.GRANT

	Prior	Granting Request for itized Examination ck I or After RCE)	Application No.: 15/180,431
1.	THE R	EQUEST FILEDJune 13, 20	16 IS <b>GRANTED</b> .
	The above- A. B.	for an original nonprovisiona	requirements for prioritized examination I application (Track I). g continued examination (RCE).
2.			indergo prioritized examination. The application will be course of prosecution until one of the following occurs:
	A.	filing a <b>petition for extension of</b>	f time to extend the time period for filing a reply;
	B.	filing an amendment to amend	the application to contain more than four independent
		claims, more than thirty total c	laims, or a multiple dependent claim;
	C.	filing a request for continued e	xamination;
	D.	filing a notice of appeal;	
	E.	filing a request for suspension of	action;
	F.	mailing of a notice of allowance;	
	G.	mailing of a final Office action;	
	H.	completion of examination as de	fined in 37 CFR 41.102; or
	I.	abandonment of the application.	
	Telephone	inquiries with regard to this decision	on should be directed to Brian W. Brown at 571-272-5338.
	/Brian W. [Signatu		Petitions Examiner, Office of Petitions (Title)

U.S. Patent and Trademark Office PTO-2298 (Rev. 02-2012)



#### UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

FILING or GRP ART APPLICATION FIL FEE REC'D NUMBER 371(c) DATE UNIT ATTY.DOCKET.NO TOT CLAIMS IND CLAIMS 15/180,431 06/13/2016 1600 AKBM-14409/US-12/CON 20 2

CONFIRMATION NO. 2763

72960 Casimir Jones, S.C. 2275 DEMING WAY, SUITE 310 MIDDLETON, WI 53562 FILING RECEIPT

Date Mailed: 06/24/2016

Receipt is acknowledged of this non-provisional patent application. The application will be taken up for examination in due course. Applicant will be notified as to the results of the examination. Any correspondence concerning the application must include the following identification information: the U.S. APPLICATION NUMBER, FILING DATE, NAME OF APPLICANT, and TITLE OF INVENTION. Fees transmitted by check or draft are subject to collection. Please verify the accuracy of the data presented on this receipt. If an error is noted on this Filing Receipt, please submit a written request for a Filing Receipt Correction. Please provide a copy of this Filing Receipt with the changes noted thereon. If you received a "Notice to File Missing Parts" for this application, please submit any corrections to this Filing Receipt with your reply to the Notice. When the USPTO processes the reply to the Notice, the USPTO will generate another Filing Receipt incorporating the requested corrections

Inventor(s)

Inge Bruheim, Volda, NORWAY; Snorre Tilseth, Bergen, NORWAY; Daniele Mancinelli, Orsta, NORWAY;

Applicant(s)

AKER BIOMARINE ANTARCTIC AS, Stamsund, NORWAY;

**Power of Attorney:** The patent practitioners associated with Customer Number <u>72960</u>

Domestic Priority data as claimed by applicant

This application is a CON of 14/020,162 09/06/2013 PAT 9375453 which is a CON of 12/057,775 03/28/2008 PAT 9034388 which claims benefit of 60/920,483 03/28/2007 and claims benefit of 60/975,058 09/25/2007 and claims benefit of 60/983,446 10/29/2007 and claims benefit of 61/024.072 01/28/2008

**Foreign Applications** for which priority is claimed (You may be eligible to benefit from the **Patent Prosecution Highway** program at the USPTO. Please see <a href="http://www.uspto.gov">http://www.uspto.gov</a> for more information.) - None. Foreign application information must be provided in an Application Data Sheet in order to constitute a claim to foreign priority. See 37 CFR 1.55 and 1.76.

Permission to Access Application via Priority Document Exchange: Yes

Permission to Access Search Results: Yes

Applicant may provide or rescind an authorization for access using Form PTO/SB/39 or Form PTO/SB/69 as appropriate.

If Required, Foreign Filing License Granted: 06/22/2016

The country code and number of your priority application, to be used for filing abroad under the Paris Convention,

is **US 15/180,431** 

Projected Publication Date: 09/29/2016

Non-Publication Request: No Early Publication Request: No

Title

BIOEFFECTIVE KRILL OIL COMPOSITIONS

**Preliminary Class** 

Statement under 37 CFR 1.55 or 1.78 for AIA (First Inventor to File) Transition Applications: No

#### PROTECTING YOUR INVENTION OUTSIDE THE UNITED STATES

Since the rights granted by a U.S. patent extend only throughout the territory of the United States and have no effect in a foreign country, an inventor who wishes patent protection in another country must apply for a patent in a specific country or in regional patent offices. Applicants may wish to consider the filing of an international application under the Patent Cooperation Treaty (PCT). An international (PCT) application generally has the same effect as a regular national patent application in each PCT-member country. The PCT process **simplifies** the filing of patent applications on the same invention in member countries, but **does not result** in a grant of "an international patent" and does not eliminate the need of applicants to file additional documents and fees in countries where patent protection is desired.

Almost every country has its own patent law, and a person desiring a patent in a particular country must make an application for patent in that country in accordance with its particular laws. Since the laws of many countries differ in various respects from the patent law of the United States, applicants are advised to seek guidance from specific foreign countries to ensure that patent rights are not lost prematurely.

Applicants also are advised that in the case of inventions made in the United States, the Director of the USPTO must issue a license before applicants can apply for a patent in a foreign country. The filing of a U.S. patent application serves as a request for a foreign filing license. The application's filing receipt contains further information and guidance as to the status of applicant's license for foreign filing.

Applicants may wish to consult the USPTO booklet, "General Information Concerning Patents" (specifically, the section entitled "Treaties and Foreign Patents") for more information on timeframes and deadlines for filing foreign patent applications. The guide is available either by contacting the USPTO Contact Center at 800-786-9199, or it can be viewed on the USPTO website at http://www.uspto.gov/web/offices/pac/doc/general/index.html.

For information on preventing theft of your intellectual property (patents, trademarks and copyrights), you may wish to consult the U.S. Government website, http://www.stopfakes.gov. Part of a Department of Commerce initiative, this website includes self-help "toolkits" giving innovators guidance on how to protect intellectual property in specific countries such as China, Korea and Mexico. For questions regarding patent enforcement issues, applicants may call the U.S. Government hotline at 1-866-999-HALT (1-866-999-4258).

#### LICENSE FOR FOREIGN FILING UNDER

#### Title 35, United States Code, Section 184

#### Title 37, Code of Federal Regulations, 5.11 & 5.15

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This license is to be retained by the licensee and may be used at any time on or after the effective date thereof unless it is revoked. This license is automatically transferred to any related applications(s) filed under 37 CFR 1.53(d). This license is not retroactive.

The grant of a license does not in any way lessen the responsibility of a licensee for the security of the subject matter as imposed by any Government contract or the provisions of existing laws relating to espionage and the national security or the export of technical data. Licensees should apprise themselves of current regulations especially with respect to certain countries, of other agencies, particularly the Office of Defense Trade Controls, Department of State (with respect to Arms, Munitions and Implements of War (22 CFR 121-128)); the Bureau of Industry and Security, Department of Commerce (15 CFR parts 730-774); the Office of Foreign AssetsControl, Department of Treasury (31 CFR Parts 500+) and the Department of Energy.

#### **NOT GRANTED**

No license under 35 U.S.C. 184 has been granted at this time, if the phrase "IF REQUIRED, FOREIGN FILING LICENSE GRANTED" DOES NOT appear on this form. Applicant may still petition for a license under 37 CFR 5.12, if a license is desired before the expiration of 6 months from the filing date of the application. If 6 months has lapsed from the filing date of this application and the licensee has not received any indication of a secrecy order under 35 U.S.C. 181, the licensee may foreign file the application pursuant to 37 CFR 5.15(b).

#### SelectUSA

The United States represents the largest, most dynamic marketplace in the world and is an unparalleled location for business investment, innovation, and commercialization of new technologies. The U.S. offers tremendous resources and advantages for those who invest and manufacture goods here. Through SelectUSA, our nation works to promote and facilitate business investment. SelectUSA provides information assistance to the international investor community; serves as an ombudsman for existing and potential investors; advocates on behalf of U.S. cities, states, and regions competing for global investment; and counsels U.S. economic development organizations on investment attraction best practices. To learn more about why the United States is the best country in the world to develop technology, manufacture products, deliver services, and grow your business, visit <a href="http://www.SelectUSA.gov">http://www.SelectUSA.gov</a> or call +1-202-482-6800.

	PATI	ENT APPL		N FEE DE		ION RECORI	)	Applica 15/18	tion or Docket Num 0,431	ber
	APPI	LICATION A	S FILEI		umn 2)	SMALL	ENTITY	OR	OTHER SMALL I	
	FOR	NUMBE	R FILE	) NUMBE	NUMBER EXTRA RATE(\$) FEE(\$)				RATE(\$)	FEE(\$)
	IC FEE FR 1.16(a), (b), or (c))		I/A	N	I/A	N/A		1	N/A	280
SEA	RCH FEE FR 1.16(k), (i), or (m))		I/A	<u> </u>	I/A	N/A		1	N/A	600
EXA	MINATION FEE FR 1.16(o), (p), or (q))		I/A	<u> </u>	I/A	N/A		1	N/A	720
TOT	AL CLAIMS FR 1.16(i))	20	minus	20= *				OR	x 80 =	0.00
INDE	EPENDENT CLAIN FR 1.16(h))	<sup>IS</sup> 2	minus	3 = *				1	x 420 =	0.00
APF FEE	LICATION SIZE	sheets of   \$310 (\$15 50 sheets	paper, th 5 for sma or fractio	and drawings e e application si all entity) for ea on thereof. See CFR 1.16(s).	ze fee due is ch additional					0.00
MUL	TIPLE DEPENDE	NT CLAIM PRE	SENT (37	7 CFR 1.16(j))						0.00
* If th	ne difference in co	lumn 1 is less th	nan zero,	enter "0" in colur	mn 2.	TOTAL		1	TOTAL	1600
AMENDMENT A	Total (37 CFR 1.16(i)) Independent	CLAIMS REMAINING AFTER AMENDMENT *	Minus	HIGHEST NUMBER PREVIOUSLY PAID FOR	PRESENT EXTRA	RATE(\$)	ADDITIONAL FEE(\$)	OR	RATE(\$)	ADDITIONAL FEE(\$)
<u> </u>	(37 CFR 1.16(h))					X =		OR	X =	
₹	Application Size Fe							┨		
	FIRST PRESENTA	TION OF MULTIP	LE DEPEN	DENT CLAIM (37 (	CFR 1.16(j))			OR		
		(Caluma 1)		(Caluman 0)	(Caluman 2)	TOTAL ADD'L FEE		OR	TOTAL ADD'L FEE	
NT B		(Column 1)  CLAIMS  REMAINING  AFTER  AMENDMENT		(Column 2) HIGHEST NUMBER PREVIOUSLY PAID FOR	(Column 3) PRESENT EXTRA	RATE(\$)	ADDITIONAL FEE(\$)		RATE(\$)	ADDITIONAL FEE(\$)
ME	Total (37 CFR 1.16(i))	*	Minus	**	=	X =		OR	x =	
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## SCORE Placeholder Sheet for IFW Content

Application Number: 15180431 Document Date: 06/13/2016

The presence of this form in the IFW record indicates that the following document type was received in electronic format on the date identified above. This content is stored in the SCORE database.

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## Drawing

At the time of document entry (noted above):

- USPTO employees may access SCORE content via eDAN using the Supplemental Content tab, or via the SCORE web page.
- External customers may access SCORE content via PAIR using the Supplemental Content tab

Form Revision Date: August 26, 2013

**Document Description: TrackOne Request** 

PTO/AIA/424 (04-14)

# CERTIFICATION AND REQUEST FOR PRIORITIZED EXAMINATION UNDER 37 CFR 1.102(e) (Page 1 of 1)

First Named Inventor:	Inge Bruheim	Nonprovisional Application Number (if known):	
Title of Invention:	BIOEFFECTIVE KRILL OIL	COMPOSITIONS	

APPLICANT HEREBY CERTIFIES THE FOLLOWING AND REQUESTS PRIORITIZED EXAMINATION FOR THE ABOVE-IDENTIFIED APPLICATION.

- 1. The processing fee set forth in 37 CFR 1.17(i)(1) and the prioritized examination fee set forth in 37 CFR 1.17(c) have been filed with the request. The publication fee requirement is met because that fee, set forth in 37 CFR 1.18(d), is currently \$0. The basic filing fee, search fee, and examination fee are filed with the request or have been already been paid. I understand that any required excess claims fees or application size fee must be paid for the application.
- 2. I understand that the application may not contain, or be amended to contain, more than four independent claims, more than thirty total claims, or any multiple dependent claims, and that any request for an extension of time will cause an outstanding Track I request to be dismissed.
- 3. The applicable box is checked below:
  - I. V Original Application (Track One) Prioritized Examination under § 1.102(e)(1)
- i. (a) The application is an original nonprovisional utility application filed under 35 U.S.C. 111(a).
   This certification and request is being filed with the utility application via EFS-Web.
  - (b) The application is an original nonprovisional plant application filed under 35 U.S.C. 111(a). This certification and request is being filed with the plant application in paper.
- ii. An executed inventor's oath or declaration under 37 CFR 1.63 or 37 CFR 1.64 for each inventor, <u>or</u> the application data sheet meeting the conditions specified in 37 CFR 1.53(f)(3)(i) is filed with the application.
  - II. Request for Continued Examination Prioritized Examination under § 1.102(e)(2)
- i. A request for continued examination has been filed with, or prior to, this form.
- ii. If the application is a utility application, this certification and request is being filed via EFS-Web.
- iii. The application is an original nonprovisional utility application filed under 35 U.S.C. 111(a), or is a national stage entry under 35 U.S.C. 371.
- iv. This certification and request is being filed prior to the mailing of a first Office action responsive to the request for continued examination.
- v. No prior request for continued examination has been granted prioritized examination status under 37 CFR 1.102(e)(2).

Signature/J. Mitchell Jones/	<sub>Date</sub> June 13, 2016
Name (Print/Typed) J. Mitchell Jones	Practitioner 44174 Registration Number
Note: This form must be signed in accordance with 37 CFR 1.33. See 37 CFR 1.4(d) for Submit multiple forms if more than one signature is required.*	
*Total of forms are submitted.	

#### **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:

- 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).
- 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 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.

Applicati	on Data Ch	not 27 CED 4 76	Attorney	Docke	et Number	AKBM-144	109/US-12/CON	
Application Data Sheet 37 CFR 1.			Applicati	on Nu	mber			
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Applica	tion Data	Shee	et 37 CFR	1.76	Attorney	Dock	et Number	AKBM-14	409/US-12/CON	
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Only complete this section when filing an application by reference under 35 U.S.C. 111(c) and 37 CFR 1.57(a). Do not complete this section if application papers including a specification and any drawings are being filed. Any domestic benefit or foreign priority information must be provided in the appropriate section(s) below (i.e., "Domestic Benefit/National Stage Information" and "Foreign Priority Information").  For the purposes of a filing date under 37 CFR 1.53(b), the description and any drawings of the present application are replaced by this eference to the previously filed application, subject to conditions and requirements of 37 CFR 1.57(a).										
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Application Data Sheet 37 CFR 1.76					y D	Oocket Number AKBM-14409/US-12/CON					
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Title of Invention	Title of Invention BIOEFFECTIVE KRILL OIL COMPOSI										
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Claims benefit of provisional

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Page 0459

Add

U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE 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.

Application Data Sheet 37 CFR 1.76			Attorney	/ D	ocket Number	AKBM-1440	9/US-12/CON
Application Bata offect of Of K 1.70			Applicat	Application Number			
Title of Invention	OMPOSITI	ION	IS				
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### **Foreign Priority Information:**

by selecting the Add button.

This section allows for the applicant to claim priority to a foreign application. Providing this information in the application data sheet constitutes the claim for priority as required by 35 U.S.C. 119(b) and 37 CFR 1.55. When priority is claimed to a foreign application that is eligible for retrieval under the priority document exchange program (PDX)<sup>i</sup> the information will be used by the Office to automatically attempt retrieval pursuant to 37 CFR 1.55(i)(1) and (2). Under the PDX program, applicant bears the ultimate responsibility for ensuring that a copy of the foreign application is received by the Office from the participating foreign intellectual property office, or a certified copy of the foreign priority application is filed, within the time period specified in 37 CFR 1.55(g)(1).

Additional Domestic Benefit/National Stage Data may be generated within this form

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Application Number	Country <sup>i</sup>	Filing Date (YYYY-MM-DD)	Access Code <sup>i</sup> (if applicable)
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# Statement under 37 CFR 1.55 or 1.78 for AIA (First Inventor to File) Transition Applications

This application (1) claims priority to or the benefit of an application filed before March 16, 2013 and (2) also
contains, or contained at any time, a claim to a claimed invention that has an effective filing date on or after March
16, 2013.
NOTE: By providing this statement under 37 CFR 1.55 or 1.78, this application, with a filing date on or after March
16, 2013, will be examined under the first inventor to file provisions of the AIA.

Application Da	ta Sheet 37 CFR 1.76	Attorney Docket Number	AKBM-14409/US-12/CON
Application Da	ita Sileet Si Ci K 1.70	Application Number	
Title of Invention	BIOEFFECTIVE KRILL OIL C	OMPOSITIONS	

### Authorization or Opt-Out of Authorization to Permit Access:

When this Application Data Sheet is properly signed and filed with the application, applicant has provided written authority to permit a participating foreign intellectual property (IP) office access to the instant application-as-filed (see paragraph A in subsection 1 below) and the European Patent Office (EPO) access to any search results from the instant application (see paragraph B in subsection 1 below).

Should applicant choose not to provide an authorization identified in subsection 1 below, applicant <u>must opt-out</u> of the authorization by checking the corresponding box A or B or both in subsection 2 below.

**NOTE**: This section of the Application Data Sheet is **ONLY** reviewed and processed with the **INITIAL** filing of an application. After the initial filing of an application, an Application Data Sheet cannot be used to provide or rescind authorization for access by a foreign IP office(s). Instead, Form PTO/SB/39 or PTO/SB/69 must be used as appropriate.

- 1. Authorization to Permit Access by a Foreign Intellectual Property Office(s)
- A. <u>Priority Document Exchange (PDX)</u> Unless box A in subsection 2 (opt-out of authorization) is checked, the undersigned hereby <u>grants the USPTO authority</u> to provide the European Patent Office (EPO), the Japan Patent Office (JPO), the Korean Intellectual Property Office (KIPO), the State Intellectual Property Office of the People's Republic of China (SIPO), the World Intellectual Property Organization (WIPO), and any other foreign intellectual property office participating with the USPTO in a bilateral or multilateral priority document exchange agreement in which a foreign application claiming priority to the instant patent application is filed, access to: (1) the instant patent application-as-filed and its related bibliographic data, (2) any foreign or domestic application to which priority or benefit is claimed by the instant application and its related bibliographic data, and (3) the date of filing of this Authorization. See 37 CFR 1.14(h) (1).
- B. <u>Search Results from U.S. Application to EPO</u> Unless box B in subsection 2 (opt-out of authorization) is checked, the undersigned hereby <u>grants the USPTO authority</u> to provide the EPO access to the bibliographic data and search results from the instant patent application when a European patent application claiming priority to the instant patent application is filed. See 37 CFR 1.14(h)(2).

The applicant is reminded that the EPO's Rule 141(1) EPC (European Patent Convention) requires applicants to submit a copy of search results from the instant application without delay in a European patent application that claims priority to the instant application.

2.	Opt-Out of Authorizations to Permit Access by a Foreign Intellectual Property Office(s)
	A. Applicant <b>DOES NOT</b> authorize the USPTO to permit a participating foreign IP office access to the instant application-as-filed. If this box is checked, the USPTO will not be providing a participating foreign IP office with any documents and information identified in subsection 1A above.
	B. Applicant <u>DOES NOT</u> authorize the USPTO to transmit to the EPO any search results from the instant patent application. If this box is checked, the USPTO will not be providing the EPO with search results from the instant application.
ı	OTE: Once the application has published or is otherwise publicly available, the USPTO may provide access to the plication in accordance with 37 CFR 1.14.

Application Da	ta Sheet 37 CFR 1.76	Attorney Docket Number	AKBM-14409/US-12/CON
Application Da	ita Sileet Si Ci K 1.70	Application Number	
Title of Invention	BIOEFFECTIVE KRILL OIL COMPOSITIONS		

## **Applicant Information:**

Providing assignment information in this section does not substitute for compliance with any requirement of part 3 of Title 37 of CFR to have an assignment recorded by the Office.				
Applicant 1 Remove				
If the applicant is the inventor (or the remaining joint inventor or inventors under 37 CFR 1.45), this section should not be completed. The information to be provided in this section is the name and address of the legal representative who is the applicant under 37 CFR 1.43; or the name and address of the assignee, person to whom the inventor is under an obligation to assign the invention, or person who otherwise shows sufficient proprietary interest in the matter who is the applicant under 37 CFR 1.46. If the applicant is an applicant under 37 CFR 1.46 (assignee, person to whom the inventor is obligated to assign, or person who otherwise shows sufficient proprietary interest) together with one or more joint inventors, then the joint inventor or inventors who are also the applicant should be identified in this section.				
<ul> <li>Assignee</li> </ul>	Legal Representative un	der 35 U.S.C. 117	Joint Inventor	
Person to whom the inventor is	obligated to assign.	Person who shows	sufficient proprietary interest	
If applicant is the legal represen	ntative, indicate the authority to t	file the patent application	, the inventor is:	
			•	
Name of the Deceased or Lega	ally Incapacitated Inventor:		_	
If the Applicant is an Organiza	ation check here.			
Organization Name	BIOMARINE ANTARCTIC AS			
Mailing Address Information For Applicant:				
Address 1	I.M. Johansens vei 99			
Address 2				
City	tamsund	State/Province		
Country NO		Postal Code	340	
Phone Number		Fax Number		
Email Address				
Additional Applicant Data may	be generated within this form by	selecting the Add button	. Add	

## **Assignee Information including Non-Applicant Assignee Information:**

Providing assignment information in this section does not substitute for compliance with any requirement of part 3 of Title 37 of CFR to have an assignment recorded by the Office.

Application Data Sheet 37 CFR 1.76			Attorney Doo	ket Number	AKBM-14	AKBM-14409/US-12/CON		
Application Da	ala Sileel	37 CFK 1.76	Application N	lumber				
Title of Invention BIOEFFECTIVE KRILL OIL CO			OMPOSITIONS					
Assignee 1								
application publicatior publication as an appl	Complete this section if assignee information, including non-applicant assignee information, is desired to be included on the patent application publication. An assignee-applicant identified in the "Applicant Information" section will appear on the patent application publication as an applicant. For an assignee-applicant, complete this section only if identification as an assignee is also desired on the patent application publication.							
If the Assignee or	Non-Annlica	ant Assignad is ar	Organization	check here		<u> </u>	Remove	
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Additional Assignee or Non-Applicant Assignee Data may be generated within this form by selecting the Add button.								
Signature:							Remove	
NOTE: This Application Data Sheet must be signed in accordance with 37 CFR 1.33(b). However, if this Application Data Sheet is submitted with the INITIAL filing of the application and either box A or B is not checked in subsection 2 of the "Authorization or Opt-Out of Authorization to Permit Access" section, then this form must also be signed in accordance with 37 CFR 1.14(c).  This Application Data Sheet must be signed by a patent practitioner if one or more of the applicants is a juristic entity (e.g., corporation or association). If the applicant is two or more joint inventors, this form must be signed by a patent practitioner, all joint inventors who are the applicant, or one or more joint inventor-applicants who have been given power of attorney (e.g., see USPTO Form PTO/AIA/81) on behalf of all joint inventor-applicants.  See 37 CFR 1.4(d) for the manner of making signatures and certifications.								
Signature /J. Mitchell Jones/				Date (Y	YYY-MM-DI	D) 2016-0	6-13	
First Name J. M	itchell	Last Name	Jones		Registra	tion Numbe	r 44174	
Additional Signatu	Additional Signature may be generated within this form by selecting the Add button.							

PTO/AIA/14 (11-15)

Approved for use through 04/30/2017. OMB 0651-0032 U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

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.

Application Da	ata Shoot 37 CED 1 76	Attorney Docket Number	AKBM-14409/US-12/CON
Application Data Sheet 37 CFR 1.76		Application Number	
Title of Invention	BIOEFFECTIVE KRILL OIL C	OMPOSITIONS	

This collection of information is required by 37 CFR 1.76. 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 23 minutes to complete, including gathering, preparing, and submitting the completed application data sheet 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.** 

### **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 the Freedom of Information Act requires disclosure of these records.
- 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.
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## DECLARATION (37 CFR 1.63) FOR UTILITY OR DESIGN APPLICATION USING AN APPLICATION DATA SHEET (37 CFR 1.76)

Title of Invention	BIOEFFECTIVE KRILL OIL COMPOSITIONS
As the belo	w named inventor, I hereby declare that:
This declar is directed	
	filed on
The above-	dentified application was made or authorized to be made by me.
I believe tha	It I am the original inventor or an original joint inventor of a claimed invention in the application.
	nowledge that any willful false statement made in this declaration is punishable under 18 U.S.C. 1001 prisonment of not more than five (5) years, or both.
	WARNING:
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	AME OF INVENTOR
Inventor: Signature:	Inge Bruheim Date (Optional): 20/5-17
Note: An appl	ication data sheet (PTO/SB/14 or equivalent), including naming the entire inventive entity, must accompany this form or must have sly filed. Use an additional PTO/AIA/01 form for each additional inventor.

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## DECLARATION (37 CFR 1.63) FOR UTILITY OR DESIGN APPLICATION USING AN APPLICATION DATA SHEET (37 CFR 1.76)

Title of Invention	BIOEFFECTIVE KRILL OIL COMPOSITIONS					
As the belo	As the below named inventor, I hereby declare that:					
This declar						
	United States application or PCT international application number 14/020,162  filed on 06-Sep-2013					
The above-i	identified application was made or authorized to be made by me.					
I believe tha	et I am the original inventor or an original joint inventor of a claimed invention in the application.					
	knowledge that any willful false statement made in this déclaration is punishable under 18 U.S.C. 1001 aprisonment of not more than five (5) years, or both.					
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contribute to (other than a to support a petitioners/a USPTO. Pe application ( patent. Furt referenced in	oplicant is cautioned to avoid submitting personal information in documents filed in a patent application that may be identity theft. Personal information such as social security numbers, bank account numbers, or credit card numbers a check or credit card authorization form PTO-2038 submitted for payment purposes) is never required by the USPTO petition or an application. If this type of personal information is included in documents submitted to the USPTO, applicants should consider redacting such personal information from the documents before submitting them to the etitioner/applicant is advised that the record of a patent application is available to the public after publication of the unless a non-publication request in compliance with 37 CFR 1.213(a) is made in the application) or issuance of a thermore, the record from an abandoned application may also be available to the public if the application is n a published application or an issued patent (see 37 CFR 1.14). Checks and credit card authorization forms ubmitted for payment purposes are not retained in the application file and therefore are not publicly available.					
LEGAL NA	AME OF INVENTOR					
Inventor:	Daniele Mancinelli  Date (Optional): 17/9/2013					
	ication data sheet (PTO/SB/14 or equivalent), including naming the entire inventive entity, must accompany this form or must have sly filed. Use an additional PTO/AIA/01 form for each additional inventor.					

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## DECLARATION (37 CFR 1.63) FOR UTILITY OR DESIGN APPLICATION USING AN APPLICATION DATA SHEET (37 CFR 1.76)

Title of Invention	BIOEFFECTIVE KRILL OIL COMPOSITIONS					
As the below named inventor, I hereby declare that:						
This declar	to:					
	United States application or PCT international application number 14/020,162  filed on 06-Sep-2013					
The above-	dentified application was made or authorized to be made by me.					
I believe tha	it I am the original inventor or an original joint inventor of a claimed invention in the application.					
I hereby acknowledge that any willful faise statement made in this declaration is punishable under 18 U.S.C: 1001 by fine or imprisonment of not more than five (5) years, or both.						
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LEGAL N	AME OF INVENTOR					
Inventor: Signature;	Snorre Tilseth Date (Optional): <u>\(\frac{1}{2}\) \(\frac{1}{2}\)</u>					
Note: An appl been previous	ication data sheet (PTO/SB/14 or equivalent), including naming the entire inventive entity, must accompany this form or must have sly filed. Use an additional PTO/AIA/01 form for each additional inventor.					

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# BIOEFFECTIVE KRILL OIL COMPOSITIONS

### CROSS-REFERENCE TO RELATED APPLICATIONS

This application is a continuation of U.S. Patent Application No. 14/020,162, filed September 6, 2013, which is a continuation of U.S. Patent Application No. 12/057,775, filed March 28, 2008, now U.S. Patent No. 9,034,388, which claims the benefit of expired U.S. Provisional Patent Application No. 60/920,483, filed March 28, 2007, expired U.S. Provisional Patent Application No. 60/975,058, filed September 25, 2007, expired U.S. Provisional Patent Application No. 60/983,446, filed October 29, 2007, and expired U.S. Provisional Patent Application No. 61/024,072, filed January 28, 2008, all of which are incorporated by reference herein in their entirety.

#### FIELD OF THE INVENTION

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This invention relates to extracts from Antarctic krill that comprise bioactive fatty acids.

#### BACKGROUND OF THE INVENTION

In the Southern Ocean, off the coast of Antarctica, Antarctic krill (*Euphausia superba*) can be found in large quantities, ranging from 300-500 million metric tons of biomass. It feeds on phytoplankton during the short Antarctic summer. During winter, however, its food supply is limited to ice algae, bacteria, marine detritus as well as depleting body protein for energy.

In order to isolate the krill oil from the krill, solvent extraction methods have been used. See, e.g., WO 00/23546. Krill lipids have been extracted by placing the material in a ketone solvent (e.g. acetone) in order to extract the lipid soluble fraction. This method involves separating the liquid and solid contents and recovering a lipid rich fraction from the liquid fraction by evaporation. Further processing steps include extracting and recovering by evaporation the remaining soluble lipid fraction from the solid contents by using a solvent such as ethanol. See, e.g., WO 00/23546. The compositions produced by these methods are characterized by containing at least 75 μg/g astaxanthin, preferably 90 μg/g astaxanthin. Another krill lipid extract disclosed contained at least 250 μg/g canastaxanthin, preferably 270 μg/g canastaxanthin.

Krill oil compositions have been described as being effective for decreasing cholesterol, inhibiting platelet adhesion, inhibiting artery plaque formation, preventing hypertension,

controlling arthritis symptoms, preventing skin cancer, enhancing transdermal transport, reducing the symptoms of premenstrual symptoms or controlling blood glucose levels in a patient. See, e.g., WO 02/102394. In yet another application, a krill oil composition has been disclosed comprising a phospholipid and/or a flavonoid. The phospholipid content in the krill lipid extract could be as high as 60% w/w and the EPA/DHA content as high as 35% (w/w). See, e.g., WO 03/011873.

Furthermore, nutraceuticals, pharmaceuticals and cosmetics comprising the phospholipid extract were disclosed. Previously, it was also shown that supercritical fluid extraction using neat CO<sub>2</sub> could be used to prevent the extraction of phospholipids in order to extract the neutral lipid fraction from krill, which comprised of esterified and free astaxanthin. See, e.g., Yamaguchi et al., *J. Agric. Food Chem.* (1986), 34(5), 904-7. Supercritical fluid extraction with solvent modifier has previously been used to extract marine phospholipids from salmon roe, but has not been previously used to extract phospholipids from krill meal. See, e.g., Tanaka et al., J. Oleo Sci. (2004), 53(9), 417-424.

The methods described above rely on the processing of frozen krill that are transported from the Southern Ocean to the processing site. This transportation is both expensive and can result in degradation of the krill starting material. Data in the literature showing a rapid decomposition of the oil in krill explains why some krill oil currently offered as an omega-3 supplement in the marketplace contains very high amounts of partly decomposed phosphatidylcholine and also partly decomposed glycerides. Saether et al., Comp. Biochem Phys. B 83B(1): 51-55 (1986). The products offered also contain high levels of free fatty acids.

What is needed in the art are methods for processing krill that do not require transport of frozen krill material over long distances and the products produced by those methods.

## SUMMARY OF THE INVENTION

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In a first aspect of the invention is a composition characterized by comprising at least 65% (w/w) phospholipids.

In another aspect of the invention is a composition obtained from aquatic or marine sources, characterized by comprising 65% (w/w) phospholipids.

In yet another aspect of the invention is a composition obtained from krill, characterized by comprising at least 65% (w/w) phospholipids.

In another aspect of the invention is a composition obtained from krill, characterized by comprising at least 65% (w/w) phospholipids and at least 39% omega-3 fatty acids (w/w).

In yet another aspect of the invention is a composition obtained from krill, characterized by comprising at least 65% (w/w) phospholipids, at least 39% omega-3 fatty acids (w/w) and at least 580 mg/kg astaxanthin esters.

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In another aspect of the invention is a composition obtained from krill, characterized by comprising at least 39% omega-3 fatty acids (w/w) and at least 580 mg/kg astaxanthin esters.

In yet another aspect of the invention is a composition obtained from krill, characterized by comprising at least 65% (w/w) phospholipids and at least 580mg/kg astaxanthin esters.

In yet another aspect, the present invention provides a krill oil effective for reducing insulin resistance, improving blood lipid profile, reducing inflammation or reducing oxidative stress.

In some embodiments, the present invention provides compositions 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. In some embodiments, the ether phospholipids are selected from the group consisting of alkylacylphosphatidylcholine, lyso-alkylacylphosphatidylcholine, alkylacylphosphatidylethanolamine, and combinations thereof. In some embodiments, the ether lipids are greater than 90% alkylacylphosphatidylcholine. In some embodiments, the non-ether phospholipids are selected from the group consisting of phosphatidylcholine, phosphatidylserine, phosphatidylethanolamine and combinations thereof. In some embodiments, krill oil composition comprises a blend of lipid fractions obtained from krill. In some preferred embodiments, krill is Euphausia superba, although other krill species also find use in the present invention. Other krill species include, but are not limited to E. pacifica, E. frigida, E. longirostris, E. triacantha, E. vallentini, Meganyctiphanes norvegica, Thysanoessa raschii and Thysanoessa inermis. In some embodiments, the compositions comprise 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. In some embodiments, the present invention provides a capsule containing the foregoing compositions.

In further embodiments, the present inventions provide compositions comprising: from about 3% to 10% ether phospholipids on a w/w basis; and from about 400 to about 2500 mg/kg astaxanthin. In some embodiments, the compositions further comprise from about 35% to 50% non-ether phospholipids on w/w basis, so that the total amount of ether phospholipids and nonether phospholipids in the composition is from about 38% to 60% on a w/w basis. In some embodiments, the compositions further comprise from about 20% to 45% triglycerides on a w/w basis. In some embodiments, the ether phospholipids are selected from the group consisting of alkylacylphosphatidylcholine, lyso-alkylacylphosphatidylcholine, alkylacylphosphatidylethanolamine, and combinations thereof. In some embodiments, the ether lipids are greater than 90% alkylacylphosphatidylcholine. In some embodiments, the non-ether phospholipids are selected from the group consisting of phosphatidylcholine, phosphatidylserine, phosphatidylethanolamine and combinations thereof. In some embodiments, krill oil composition comprises a blend of lipid fractions obtained from krill. In some preferred embodiments, krill is Euphausia superba, although other krill species also find use in the present invention. Other krill species include, but are not limited to E. pacifica, E. frigida, E. longirostris, E. triacantha, E. vallentini, Meganyctiphanes norvegica, Thysanoessa raschii and Thysanoessa inermis. In some embodiments, the compositions comprise 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. In some embodiments, the present invention provides a capsule containing the foregoing compositions.

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In some embodiments, the present invention provides a composition comprising at least 65% (w/w) of phospholipids, said phospholipids characterized in containing at least 35% omega-3 fatty acid residues. In some preferred embodiments, the composition is derived from a marine or aquatic biomass. In some further preferred embodiments, the composition is derived from krill. In some embodiments, the composition comprises less than 2% free fatty acids. In some embodiments, composition comprises less than 10% triglycerides. In some preferred embodiments, the phospholipids comprise greater than 50% phosphatidylcholine. In some embodiments, the composition comprises at least 500 mg/kg astaxanthin esters. In some embodiments, the composition comprises at least 500 mg/kg astaxanthin esters and at least 36% (w/w) omega-3 fatty acids. In some embodiments, the composition comprises less than about

0.5g/100g total cholesterol. In some embodiments, the composition comprises less than about 0.45% arachidonic acid (w/w).

In some embodiments, the present invention provides a krill lipid extract comprising at least 500, 100, 1500, 2000, 2100, or 2200 mg/kg astaxanthin esters and at least 36% (w/w) omega-3 fatty acids. In further embodiments, the present invention provides 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).

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In some embodiments, the present invention provides methods comprising administering the foregoing compositions to a subject in an amount effective for reducing insulin resistance, reducing inflammation, improving blood lipid profile and reducing oxidative stress.

In some embodiments, the present invention provides a krill lipid extract comprising greater than about 80% triglycerides and greater than about 90, 100, 500, 1000, 1500, 200, 2100 or 2200 mg/kg astaxanthin esters. In some embodiments, the krill lipid extract is characterized in containing from about 5% to about 15% omega-3 fatty acid residues. In some embodiments, the krill lipid extract is characterized in containing less than about 5% phospholipids. In some embodiments, the krill lipid extract is characterized in comprising from about 5% to about 10% cholesterol.

In some embodiments, the present invention provides a krill meal composition comprising less than about 50g/kg total fat. In some embodiments, the krill meal composition comprises from about 5 to about 20 mg/kg astaxanthin esters. In some embodiments, the krill meal composition comprises greater than about 65% protein. In some embodiments, the krill meal composition of comprises greater than about 70% protein. In some further embodiments, the present invention provides an animal feed comprising the krill meal composition.

In some embodiments, the present invention provides methods of increasing flesh coloration in an aquatic species comprising feeding said aquatic species a composition comprising the krill meal described above. In some embodiments, the present invention provides methods of increasing growth and overall survival rate of aquatic species by feeding the krill meal described above.

In some embodiments, the present invention provides methods of producing krill oil comprising: a) providing krill meal; and b) extracting oil from said krill meal. In some embodiments, the krill meal is produced by heat-treating krill. In some embodiments, the krill

meal is stored prior to the extraction step. In some embodiments, the extracting step comprises extraction by supercritical fluid extraction. In some embodiments, the supercritical fluid extraction is a two step process comprising a first extraction step with carbon dioxide and a low concentration of a co-solvent (e.g., from about 1-10% co-solvent) and a second extraction step with carbon dioxide and a high concentration of a co-solvent (e.g., from about 10-30% co-solvent). In preferred embodiments, the co-solvent is a C<sub>1</sub>-C<sub>3</sub> monohydric alcohol, preferably ethanol. In some embodiments, the present invention provides oil produced by the foregoing method.

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In some embodiments, the present invention provides methods 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. In some embodiments, the denaturation step comprises heating of said fresh krill. In some embodiments, the denaturation step comprises heating said fresh krill after grinding. In some embodiments, the methods further comprise storing said denatured krill product at room temperature or below between the denaturation step and the extraction step. In some embodiments, the enzyme denaturation step is achieved by application of heat. In some embodiments, the extraction step comprises use of supercritical carbon dioxide, with or without use of a polar modifier. In some embodiments, the extraction step comprises use of ethanol. In some embodiments, the extraction step is comprises ethanol extraction followed by acetone to precipitation of phospholipids. In some embodiments, the denatured krill product is a meal. In some embodiments, the present invention provides oil produced by the foregoing method.

In some embodiments, the present invention provides a composition comprising oil extracted from krill having a phosphatidylcholine content of greater then about 50% (w/w). In some embodiments, the oil has a phosphatidylcholine content of greater then about 70% (w/w). In some embodiments, the oil has a phosphatidylcholine content of greater then about 80% (w/w). In some embodiments, the composition comprises less than 2% free fatty acids. In some embodiments, the composition comprises less than 10% triglycerides. In some embodiments, the composition comprises at least 500 mg/kg astaxanthin esters. In some embodiments, the composition comprises less than about 0.45% arachidonic acid (w/w).

In some embodiments, the present invention provides composition comprising odorless krill oil. In some embodiments, the odorless krill oil comprises less than about 10 mg/kg (w/w)

trimethylamine. In some further embodiments, the present invention provides 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.

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In some embodiments, the present invention provides a composition comprising krill oil containing less than about 70 micrograms/kilogram (w/w) astaxanthin esters. In some embodiments, the compositions comprise less than about 50 micrograms/kilogram (w/w) astaxanthin esters. In some embodiments, the compositions comprise less than about 20 micrograms/kilogram (w/w) astaxanthin esters. In some embodiments, the compositions comprise less than about 5 micrograms/kilogram (w/w) astaxanthin esters.

In some embodiments, the present invention provides 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.

In further embodiments, the present invention provides 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. In some embodiments, the blended krill oil product comprises a blend of lipid fractions obtained from *Euphausia superba*. In some embodiments, the 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.

In still other embodiments, the present invention provides a *Euphausia superba* krill oil composition comprising: from about 30% to 60% w/w phospholipids; from about 20% to 50% 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.

In still further embodiments, the present invention provides a dietary supplement comprising encapsulated *Euphausia superba* krill oil comprising from about 30% to 60% w/w phospholipids; from about 20% to 50% 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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In some embodiments, the present invention provides methods of making a *Euphausia superba* krill oil composition comprising: contacting *Euphausia superba* with a polar solvent to provide a polar extract comprising phospholipids; contacting *Euphausia 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 comprising from about 30% to 60% w/w phospholipids; from about 20% to 50% 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. In some embodiments, the methods further comprise the step of encapsulating the *Euphausia superba* krill oil. In some embodiments, the present invention provides a *Euphausia superba* krill oil produced by the methods described above.

In some embodiments, the present invention provides methods of producing a dietary supplement comprising; contacting *Euphausia superba* with a polar solvent to provide an polar extract comprising phospholipids; contacting *Euphausia* 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 comprising from about 30% to 60% w/w phospholipids; from about 20% to 50% 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; and encapsulating said *Euphausia superba* krill oil.

In some embodiments, the present invention provides methods 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. The present invention is not limited to any particular krill oil composition. In some embodiments, the krill oil composition is a *Euphausia superba* krill oil composition. The present invention is not limited to any particular formulation of krill oil. In some embodiments, the krill

oil composition is encapsulated. In some preferred embodiments, the effective amount of a krill oil composition is from 0.2 grams to 10 grams of said krill oil composition. In some embodiments, the krill oil composition comprises: 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. In some embodiments, the krill oil composition comprises a blend of lipid fractions obtained from *Euphausia superba*. In some embodiments, the krill oil 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. In some embodiments, the krill oil composition comprises from about 30% to 60% w/w phospholipids; from about 20% to 50% 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, and wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

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In some embodiments, the present invention provides methods 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. The present invention is not limited to any particular krill oil composition. In some embodiments, the krill oil composition is a Euphausia superba krill oil composition. The present invention is not limited to any particular formulation of krill oil. In some embodiments, the krill oil composition is encapsulated. In some preferred embodiments, the effective amount of a krill oil composition is from 0.2 grams to 10 grams of said krill oil composition. In some embodiments, the krill oil composition comprises: 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. In some embodiments, the krill oil composition comprises a blend of lipid fractions obtained from Euphausia superba. In some embodiments, the krill oil 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. In some embodiments, the krill oil composition comprises from about 30% to 60% w/w phospholipids; from about 20% to 50% 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, and wherein from about 70% to 95% of said omega-3 fatty acids are attached to said phospholipids.

In some embodiments, the present invention provides methods 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. In some embodiments, the present invention provides methods 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. In some embodiments, the present invention provides methods 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. In some embodiments, the present invention provides methods 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. In some embodiments, the present invention provides methods 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.

#### **DESCRIPTION OF THE FIGURES**

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- Figure 1. 31P NMR analysis of polar lipids in krill oil.
- Figure 2. Blood lipid profiles in Zucker rats fed different forms of omega-3 fatty acids (TAG = FO, PL1 = NKO and PL2 = Superba).
- Figure 3. Plasma glucose concentration in Zucker rats fed different forms of omega-3 fatty acids.
- Figure 4. Plasma insulin concentration in Zucker rats fed different forms of omega-3 fatty acids.
- Figure 5. Estimated HOMA-IR values in Zucker rats fed different forms of omega-3 fatty acids.
- Figure 6. The effect of dietary omega-3 fatty acids on TNF□ production by peritoneal macrophages.
  - Figure 7. The effect of dietary omega-3 fatty acids on lipid accumulation in the liver.

Figure 8. The effect of dietary omega-3 fatty acids on lipid accumulation in the muscle.

Figure 9. The effect of dietary omega-3 fatty acids on lipid accumulation in the heart.

Figure 10. Relative concentrations of DHA in the brain in Zucker rats supplemented with omega-3 fatty acids.

Figure 11. Mean group body weights (g) in the collagen-induced male DBA/1 arthritic mice. B - PL2 is the krill oil group. \* p<0.05, significantly different from Group A (Positive Control - Fish Oil) and Group C (Control).

Figure 12. Body weight for the various treatment groups.

Figure 13. Muscle weight for the various treatment groups.

Figure 14. Muscle to body weight ratio for the various treatment groups.

Figure 15. Serum adiopnectin levels (ng/ml) for the various treatment groups.

Figure 16. Serum insulin levels for the various treatment groups.

Figure 17. Blood glucose (mmol/l) levels in the various treatment groups.

Figure 18. HOMA-IR values for the various treatment groups.

Figure 19. Liver triglyceride levels (µmol/g) for the various treatment groups.

## **DEFINITIONS**

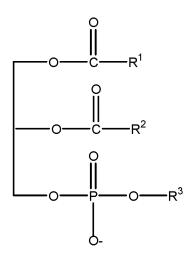
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As used herein, "phospholipid" refers to an organic compound having the following general structure:



wherein R1 is a fatty acid residue, R2 is a fatty acid residue or –OH, and R3 is a –H or nitrogen containing compound choline (HOCH<sub>2</sub>CH<sub>2</sub>N<sup>+</sup>(CH<sub>3</sub>)<sub>3</sub>OH<sup>-</sup>), ethanolamine (HOCH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub>), inositol or serine. R1 and R2 cannot simultaneously be OH. When R3 is an –OH, the compound is a diacylglycerophosphate, while when R3 is a nitrogen-containing compound, the compound is a phosphatide such as lecithin, cephalin, phosphatidyl serine or plasmalogen.

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An "ether phospholipid" as used herein refers to a phospholipid having an ether bond at position 1 the glycerol backbone. Examples of ether phospholipids include, but are not limited to, alkylacylphosphatidylcholine (AAPC), lyso-alkylacylphosphatidylcholine (LAAPC), and alkylacylphosphatidylethanolamine (AAPE). A "non-ether phospholipid" is a phospholipid that does not have an ether bond at position 1 of the glycerol backbone.

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, 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).

As used herein, astaxanthin refers to the following chemical structure:

As used herein, astaxanthin esters refer to the fatty acids esterified to OH group in the astaxanthin molecule.

As used herein, the term w/w (weight/weight) refers to the amount of a given substance in a composition on weight basis. For example, a composition comprising 50% w/w phospholipids means that the mass of the phospholipids is 50% of the total mass of the composition (i.e., 50 grams of phospholipids in 100 grams of the composition, such as an oil).

#### DETAILED DESCRIPTION OF THE INVENTION

This invention discloses novel krill oil compositions characterized by containing high levels of astaxanthin, phospholipids, included an enriched quantities of ether phospholipids, and omega-3 fatty acids. The krill oils compositions are extracted from krill meal using supercritical fluid extraction (SFE) with a co-solvent modifier. The krill meal has been processed on board a ship in Antarctica using live krill as starting material in order to ensure the highest possible quality of the krill meal. The krill oils are extracted from the krill meal in two stages, in step 1 the neutral fraction is extracted using neat supercritical CO<sub>2</sub> or in combination with 5% ethanol. The neutral fraction consisted mostly of triglycerides and cholesterol. In stage 2, the polar lipids (phospholipids) are extracted by adding at least 20% ethanol to the supercritical CO<sub>2</sub> extraction medium.

The present invention provides methods to avoid decomposition of glycerides and phospholipids in krill oil and compositions produced by those methods. The product obtained by these new methods is virtually free of enzymatically decomposed oil constituents. The solution to the problem is to incorporate a protein denaturation step on fresh krill prior to use of any extraction technology. Denaturation can be achieved by thermal stress or by other means. After denaturation, the oil can be extracted by an optional selection of nonpolar and polar solvents including use of supercritical carbon dioxide. Krill is adapted to a very efficient nutrient digestion at very low temperatures. Therefore the enzymes are sensitive to heat and the step of applying thermal denaturation of lipases and phospholipases does not imply use of very high temperatures. Surprisingly, it has been found that the use of mild denaturation conditions can greatly enhance the quality of krill oil.

Additionally, a major obstacle of several processes of extraction is the cost of removing water. This is particularly true for methods feasible for extraction of highly unsaturated lipids where freeze drying has been regarded as the method of choice to avoid oxidative breakdown of lipids. However, the lipids in krill are surprisingly stable against oxidative deterioration. Therefore, a process including moderate use of heat in the water removing process is feasible provided that the enzymes have been inactivated.

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## A. Krill Processing

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The present invention provides methods for processing freshly caught krill at the site of capture and preferably on board a ship. After processing on board, the krill can be further subjected to extraction processes on board the ship or at a remote location away from the ship. The processing steps described herein also allow for the storage of krill material, preferably a krill meal for from about 1,2, 3, 4, 5, 6, 8, 9, 10, 11, or 12 months to about 24 to 36 months prior to processing.

In some preferred embodiments, freshly caught krill is first subjected to a protein denaturation step. The present invention is not limited to any particular method of protein denaturation. In some embodiments, the denaturation is accomplished by application of chemicals, heat, or combinations thereof. In some embodiments, freshly caught krill is wet pressed to obtain oil and meal. In some embodiments, the meal is then heated to a temperature of about 50°C to about 100°C for about 20 minutes to about an hour, preferably about 40 minutes to denature the proteins. In some embodiments, this material is then pressed to yield a press cake. When this method is used on krill, only a small amount of oil is released. Most of the oil is still present in the denatured meal. In some embodiments, antioxidants such as ethoxyquin or Vitamin E are added to the meal. However, as shown in the examples, the resulting meal is surprisingly stable. The stability can only partly be explained by addition of an antioxidant to the meal. This antioxidant can, after extraction of the oil from denatured meal, be removed by further processing steps. Alternatively the oil can be extracted rather shortly after production of the meal without any addition of antioxidant in the process. Further, storage conditions at a low to very low temperature can be applied if addition of antioxidant is not desired.

Krill oil extracted from denatured krill meal by supercritical fluid extraction even 19 months after the production of the meal contained virtually no decomposed phospholipids. This product turned out to be substantially different from samples of krill oil available in the market today. Previously described commercial krill processing procedures utilize krill that has been frozen immediately after catching followed by freeze drying and extraction at low temperatures. However, these processes only yield a suitable product if the time the krill is kept frozen is very short or the temperature is extremely low (-60°to -80°C). However, data provided herein clearly shows that if a step of denaturation of the proteins is added in front of an optional extraction method, an excellent krill oil can be produced even after a long time of storage. This

methodology also opens up for use of alternative methods to remove water prior to extraction, which in turn has a great impact on costs in full scale operation. If a long time of storage is desired, the denatured material should preferably be stored at low temperature preferably at - 20°C.

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In some embodiments, krill oil is extracted from the denatured krill meal. In some embodiments, the krill oil is extracted by contacting the krill meal with ethanol. In some embodiments, krill is then extracted with a ketone solvent such as acetone. In other embodiments, the krill oil is extracted by one or two step supercritical fluid extraction. In some embodiments, the supercritical fluid extraction uses carbon dioxide and neutral krill oil is produced. In some embodiments, the supercritical fluid extraction uses carbon dioxide with the addition of a polar entrainer, such as ethanol, to produce a polar krill oil. In some embodiments, the krill meal is first extracted with carbon dioxide followed by carbon dioxide with a polar entrainer, or vice versa. In some embodiments, the krill meal is first extracted with CO<sub>2</sub> supplemented with a low amount of a polar co-solvent (e.g., from about 1% to about 10%, preferably about 5%) such a C<sub>1</sub>-C<sub>3</sub> monohydric alcohol, preferably ethanol, followed by extraction with CO<sub>2</sub> supplemented with a high amount of a polar co-solvent (from about 10% to about 30%, preferably about 23%) such as such a C<sub>1</sub>-C<sub>3</sub> monohydric alcohol, preferably ethanol, or vice versa. Surprisingly, it has been found that use of a low amount of polar solvent in the CO<sub>2</sub> as an entrainer facilitates the extraction of neutral lipid components and astaxanthin in a single step. Use of the high of polar solvent as an entrainer in the other step facilitates extraction of ether phospholipids, as well as non-ether phospholipids.

The present invention is distinguished from previously described krill oil products, such as those described in U.S. Pat. No. 6,800,299 or WO 03/011873 and Neptune brand krill oil, by having substantially higher levels of non-ether phospholipids, ether phospholipids, and astaxanthin. The krill oils of the present invention also have unexpected and superior properties as compared to previously available krill oils. In particular, the krill oil of the present invention has been demonstrated to reduce blood LDL cholesterol levels, improve DHA transfer to the brain as well as reduce lipid accumulation in the liver and muscle while the previously described krill oil compositions do not have such a properties. Accordingly, in some embodiments, the present invention provides a krill oil composition, preferably a *Euphausia superba* krill oil composition, comprising from about 40% to about 60% w/w phospholipids, preferably from

about 45% to 55% w/w phospholipids and from about 300 mg/kg astaxanthin to about 2500 mg/kg astaxanthin, preferably from about 1000 to about 2200 mg/kg astaxanthin, more preferably from about 1500 to about 2200 mg/kg astaxanthin. In some preferred embodiments, the compositions comprise greater than about 1000, 1500, 1800, 1900, 2000, or 2100 mg/kg astaxanthin. In some preferred embodiments, the krill oil compositions of the present invention comprise from about 1%, 2%, 3% or 4% to about 8%, 10%, 12% or 15% w/w ether phospholipids or greater than about 4%, 5%, 6%, 7%, 8%, 9% or 10% ether phospholipids. In some embodiments the ether phospholipids are preferably alkylacylphosphatidylcholine, lysoalkylacylphosphatidylcholine, alkylacylphosphatidyl-ethanolamine or combinations thereof. In some embodiments, the krill oil compositions comprise from about 1%, 2%, 3% or 4% to about 8%, 10%, 12% or 15% w/w ether phospholipids and from about 30%, 33%, 40%, 42%, 45%, 48%, 50%, 52%, 54%, 55% 56%, 58% to about 60% non-ether phospholipids so that the total amount of phospholipids (both ether and non-ether phospholipids) ranges from about 40% to about 60%. One of skill in the art will recognize that the range of 40% to 60% total phospholipids, as well as the other ranges of ether and non-ether phospholipids, can include other values not specifically listed within the range.

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In further embodiments, the compositions comprise from about 20% to 45% w/w triglycerides; and from about 400 to about 2500 mg/kg astaxanthin. In some embodiments, the compositions comprise from about 20% to 35%, preferably from about 25% to 35%, omega-3 fatty acids as a percentage of total fatty acids in the composition, wherein from about 70% to 95%, or preferably from about 80% to 90% of the omega-3 fatty acids are attached to the phospholipids. In some embodiments, the present invention provides encapsulated *Euphausia superba* krill oil compositions. In some embodiments, the present invention provides a method of making a *Euphausia superba* krill oil composition comprising contacting *Euphausia superba* with a polar solvent to provide an polar extract comprising phospholipids, contacting *Euphausia superba* with a neutral solvent to provide a neutral extract comprising triglycerides and astaxanthin, and combining said polar extract and said neutral extract to provide the *Euphausia superba* krill oils described above. In some embodiments, fractions from polar and non-polar extractions are combined to provide a final product comprising the desired ether phospholipids, non-ether phospholipids, omega-3 moieties and astaxanthin. In other embodiments, the present invention provides methods of making a *Euphausia superba* (or other krill species) krill oil

comprising contacting a *Euphausia superba* preparation such as *Euphausia superba* krill meal under supercritical conditions with CO<sub>2</sub> containing a low amount of a polar solvent such as ethanol to extract neutral lipids and astaxanthin; contacting meal remaining from the first extraction step under supercritical conditions with CO<sub>2</sub> containing a high amount of a polar solvent such as ethanol to extract a polar lipid fraction containing ether and non-ether phospholipids; and then blending the neutral and polar lipid extracts to provide the compositions described above.

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The krill oil extracted by the methods of the present invention contains few enzymatic breakdown products. Examples of the krill oil compositions of the present invention are provided in Tables 9-24. In some embodiments, the present invention provides a polar krill oil comprising at least 65% (w/w) of phospholipids, wherein the phospholipids are characterized in containing at least 35% omega-3 fatty acid residues. The present invention is not limited to the presence of any particular omega-3 fatty acid residues in the krill oil composition. In some preferred embodiments, the krill oil comprises EPA and DHA residues. In some embodiments, the krill oil compositions comprise less than about 5%, 4%, 3% or preferably 2% free fatty acids on a weight/weight (w/w) basis. In some embodiments, the krill oil compositions comprise less than about 25%, 20%, 15%, 10% or 5% triglycerides (w/w). In some embodiments, the krill oil compositions comprise greater than about 30%, 40%, 45%, 50%, 55%, 60%, or 65% phosphatidyl choline (w/w). In some embodiments, the krill oil compositions comprise greater than about 100, 200, 300, 400, or 500 mg/kg astaxanthin esters and up to about 700 mg/kg astaxanthin esters. In some embodiments, the present invention provides krill oil compositions comprising at least 500, 1000, 1500, 2000, 2100, or 2200 mg/kg astaxanthin esters and at least 36% (w/w) omega-3 fatty acids. In some embodiments, the krill oil compositions of the present invention comprise less than about 1.0g/100g, 0.5g/100g, 0.2g/100g or 0.1g/100g total cholesterol. In some embodiments, the krill oil compositions of the present invention comprise less than about 0.45

In some embodiments, the present invention provides a neutral krill oil extract comprising greater than about 70%, 75% 80%, 85% or 90% triglycerides. In some embodiments, the krill oil compositions comprise from about 50 to about 2500 mg/kg astaxanthin esters. In some embodiments, the krill oil compositions comprise from about 50, 100, 200, or 500 to about 750, 1000, 1500 or 2500 mg/kg astaxanthin esters. In some embodiments, the compositions comprise

from about 1% to about 30% omega-3 fatty acid residues, and preferably from about 5%-15% omega-3 fatty acid residues. In some embodiments, the krill oil compositions comprise less than about 20%, 15%, 10% or 5% phospholipids.

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In some embodiments, the present invention provides krill oil containing less than about 70, 60, 50, 40, 30, 20, 10, 5 or 1 micrograms/kilogram (w/w) astaxanthin esters. In some embodiments, the krill oil is clear or only has a pale red color. In some embodiments, the low-astaxanthin krill oil is obtained by first extracting a krill material, such as krill oil, by supercritical fluid extraction with neat carbon dioxide. It is contemplated that this step removes astaxanthin from the krill material. In some embodiments, the krill material is then subjected to supercritical fluid extraction with carbon dioxide and a polar entrainer such as ethanol, preferably about 20% ethanol. The oil extracted during this step is characterized in containing low amounts of astaxanthin. In other embodiments, krill oil comprising astaxanthin is extracted by countercurrent supercritical fluid extraction with neat carbon dioxide to provide a low-astaxanthin krill oil.

In some embodiments, the present invention provides krill oil that is substantially odorless. By substantially odorless it is meant that the krill oil lacks an appreciable odor as determined by a test panel. In some embodiments, the substantially odorless krill oil comprises less than about 10, 5 or 1 milligrams/kilogram trimethylamine. In some preferred embodiments, the odorless krill oil is produced by first subjecting krill material to supercritical fluid extraction with neat carbon dioxide to remove odor causing compounds such as trimethylamine, followed by extraction with carbon dioxide with a polar entrainer such as ethanol.

In some embodiments, the present invention provides a delipidated krill meal produced after extraction of lipids from the krill meal. In some embodiments, the delipidated krill meal comprises krill protein. In some embodiments, the delipidated krill meal comprises less than about 200, 150, 120, 100, 75, 65, 60, 55, or 50 g/kg total fat. In some embodiments, the delipidated krill meal comprises from about 1 to about 100 mg/kg astaxanthin esters, and preferably from about 5 to about 20 mg/kg astaxanthin esters. In some embodiments, the delipidated krill meal comprises greater than about 60%, 65%, 70% or 75% krill protein. In some embodiments, the present invention provides animal feeds comprising the delipidated krill meal. In some embodiments, the animal feed is a fish feed or aquatic organism feed, such as shrimp feed, crab feed, or crawfish feed. In preferred embodiments, the krill meal is incorporated into

complete ration for the target organism. In preferred embodiments, the feed is provided in pelleted form. In many instances, compounds such as astaxanthin are removed during delipidation. The methods of the present invention provide a delipidated krill meal that retains significant amounts of astaxanthin. Accordingly, in some embodiments, the present invention provides methods of feeding aquatic organisms, comprising providing to the aquatic organism a feed comprising the delipidated krill meal described above. In other embodiments, the present invention provides methods of increasing flesh coloration in an aquatic species comprising feeding the aquatic species a comprising the delipidated krill meal described above.

## 10 B. Compositions Containing Krill Oil

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In some embodiments, the compositions of this invention (such as those described in the preceding sections) are contained in acceptable excipients and/or carriers for oral consumption. The actual form of the carrier, and thus, the composition itself, is not critical. The carrier may be a liquid, gel, gelcap, capsule, powder, solid tablet (coated or non-coated), tea, or the like. The composition is preferably in the form of a tablet or capsule and most preferably in the form of a soft gel capsule. Suitable excipient and/or carriers include maltodextrin, calcium carbonate, dicalcium phosphate, tricalcium phosphate, microcrystalline cellulose, dextrose, rice flour, magnesium stearate, stearic acid, croscarmellose sodium, sodium starch glycolate, crospovidone, sucrose, vegetable gums, lactose, methylcellulose, povidone, carboxymethylcellulose, corn starch, and the like (including mixtures thereof). Preferred carriers include calcium carbonate, magnesium stearate, maltodextrin, and mixtures thereof. The various ingredients and the excipient and/or carrier are mixed and formed into the desired form using conventional techniques. The tablet or capsule of the present invention may be coated with an enteric coating that dissolves at a pH of about 6.0 to 7.0. A suitable enteric coating that dissolves in the small intestine but not in the stomach is cellulose acetate phthalate. Further details on techniques for formulation for and administration may be found in the latest edition of Remington's Pharmaceutical Sciences (Maack Publishing Co., Easton, PA).

The dietary supplement may comprise one or more inert ingredients, especially if it is desirable to limit the number of calories added to the diet by the dietary supplement. For example, the dietary supplement of the present invention may also contain optional ingredients including, for example, herbs, vitamins, minerals, enhancers, colorants, sweeteners, flavorants,

inert ingredients, and the like. For example, the dietary supplement of the present invention may contain one or more of the following: ascorbates (ascorbic acid, mineral ascorbate salts, rose hips, acerola, and the like), dehydroepiandosterone (DHEA), Fo-Ti or Ho Shu Wu (herb common to traditional Asian treatments), Cat's Claw (ancient herbal ingredient), green tea (polyphenols), inositol, kelp, dulse, bioflavinoids, maltodextrin, nettles, niacin, niacinamide, rosemary, selenium, silica (silicon dioxide, silica gel, horsetail, shavegrass, and the like), spirulina, zinc, and the like. Such optional ingredients may be either naturally occurring or concentrated forms.

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In some embodiments, the dietary supplements further comprise vitamins and minerals including, but not limited to, calcium phosphate or acetate, tribasic; potassium phosphate, dibasic; magnesium sulfate or oxide; salt (sodium chloride); potassium chloride or acetate; ascorbic acid; ferric orthophosphate; niacinamide; zinc sulfate or oxide; calcium pantothenate; copper gluconate; riboflavin; beta-carotene; pyridoxine hydrochloride; thiamin mononitrate; folic acid; biotin; chromium chloride or picolonate; potassium iodide; sodium selenate; sodium molybdate; phylloquinone; vitamin D3; cyanocobalamin; sodium selenite; copper sulfate; vitamin A; vitamin C; inositol; potassium iodide. Suitable dosages for vitamins and minerals may be obtained, for example, by consulting the U.S. RDA guidelines.

In further embodiments, the compositions comprise at least one food flavoring such as acetaldehyde (ethanal), acetoin (acetyl methylcarbinol), anethole (parapropenyl anisole), benzaldehyde (benzoic aldehyde), N butyric acid (butanoic acid), d or l carvone (carvol), cinnamaldehyde (cinnamic aldehyde), citral (2,6 dimethyloctadien 2,6 al 8, gera nial, neral), decanal (N decylaldehyde, capraldehyde, capric aldehyde, caprinaldehyde, aldehyde C 10), ethyl acetate, ethyl butyrate, 3 methyl 3 phenyl glycidic acid ethyl ester (ethyl methyl phenyl glycidate, strawberry aldehyde, C 16 aldehyde), ethyl vanillin, geraniol (3,7 dimethyl 2,6 and 3,6 octadien 1 ol), geranyl acetate (geraniol acetate), limonene (d, 1, and dl), linalool (linalol, 3,7 dimethyl 1,6 octadien 3 ol), linalyl acetate (bergamol), methyl anthranilate (methyl 2 aminobenzoate), piperonal (3,4 methylenedioxy benzaldehyde, heliotropin), vanillin, alfalfa (Medicago sativa L.), allspice (Pimenta officinalis), ambrette seed (Hibiscus abelmoschus), angelic (Angelica archangelica), Angostura (Galipea officinalis), anise (Pimpinella anisum), star anise (Illicium verum), balm (Melissa officinalis), basil (Ocimum basilicum), bay (Laurus nobilis), calendula (Calendula officinalis), (Anthemis nobilis), capsicum (Capsicum frutescens), caraway (Carum carvi), cardamom (Elettaria cardamomum), cassia, (Cinnamomum cassia), cayenne pepper

(Capsicum frutescens), Celery seed (Apium graveolens), chervil (Anthriscus cerefolium), chives (Allium schoenoprasum), coriander (Coriandrum sativum), cumin (Cuminum cyminum), elder flowers (Sambucus canadensis), fennel (Foeniculum vulgare), fenugreek (Trigonella foenum graecum), ginger (Zingiber officinale), horehound (Marrubium vulgare), horseradish (Armoracia lapathifolia), hyssop (Hyssopus officinalis), lavender (Lavandula officinalis), mace (Myristica fragrans), marjoram (Majorana hortensis), mustard (Brassica nigra, Brassica juncea, Brassica hirta), nutmeg (Myristica fragrans), paprika (Capsicum annuum), black pepper (Piper nigrum), peppermint (Mentha piperita), poppy seed (Papayer somniferum), rosemary (Rosmarinus officinalis), saffron (Crocus sativus), sage (Salvia officinalis), savory (Satureia hortensis, Satureia montana), sesame (Sesamum indicum), spearmint (Mentha spicata), tarragon (Artemisia dracunculus), thyme (Thymus vulgaris, Thymus serpyllum), turmeric (Curcuma longa), vanilla (Vanilla planifolia), zedoary (Curcuma zedoaria), sucrose, glucose, saccharin, sorbitol, mannitol, aspartame. Other suitable flavoring are disclosed in such references as Remington's Pharmaceutical Sciences, 18th Edition, Mack Publishing, p. 1288-1300 (1990), and Furia and Pellanca, Fenaroli's Handbook of Flavor Ingredients, The Chemical Rubber Company, Cleveland, Ohio, (1971), known to those skilled in the art.

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In other embodiments, the compositions comprise at least one synthetic or natural food coloring (e.g., annatto extract, astaxanthin, beet powder, ultramarine blue, canthaxanthin, caramel, carotenal, beta carotene, carmine, toasted cottonseed flour, ferrous gluconate, ferrous lactate, grape color extract, grape skin extract, iron oxide, fruit juice, vegetable juice, dried algae meal, tagetes meal, carrot oil, corn endosperm oil, paprika, paprika oleoresin, riboflavin, saffron, tumeric, tumeric and oleoresin).

In still further embodiments, the compositions comprise at least one phytonutrient (e.g., soy isoflavonoids, oligomeric proanthcyanidins, indol 3 carbinol, sulforaphone, fibrous ligands, plant phytosterols, ferulic acid, anthocyanocides, triterpenes, omega 3/6 fatty acids, conjugated fatty acids such as conjugated linoleic acid and conjugated linolenic acid, polyacetylene, quinones, terpenes, cathechins, gallates, and quercitin). Sources of plant phytonutrients include, but are not limited to, soy lecithin, soy isoflavones, brown rice germ, royal jelly, bee propolis, acerola berry juice powder, Japanese green tea, grape seed extract, grape skin extract, carrot juice, bilberry, flaxseed meal, bee pollen, ginkgo biloba, primrose (evening primrose oil), red clover,

burdock root, dandelion, parsley, rose hips, milk thistle, ginger, Siberian ginseng, rosemary, curcumin, garlic, lycopene, grapefruit seed extract, spinach, and broccoli.

In still other embodiments, the compositions comprise at least one vitamin (e.g., vitamin A, thiamin (B1), riboflavin (B2), pyridoxine (B6), cyanocobalamin (B12), biotin, ascorbic acid (vitamin C), retinoic acid (vitamin D), vitamin E, folic acid and other folates, vitamin K, niacin, and pantothenic acid). In some embodiments, the particles comprise at least one mineral (e.g., sodium, potassium, magnesium, calcium, phosphorus, chlorine, iron, zinc, manganese, flourine, copper, molybdenum, chromium, selenium, and iodine). In some particularly preferred embodiments, a dosage of a plurality of particles includes vitamins or minerals in the range of the recommended daily allowance (RDA) as specified by the United States Department of Agriculture. In still other embodiments, the particles comprise an amino acid supplement formula in which at least one amino acid is included (e.g., 1-carnitine or tryptophan).

## C. Uses of Krill Oil

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Previously, it was disclosed that omega-3 fatty acids have anti-inflammatory properties. See, e.g., Calder. Am. J. Clin. Nutr. 83 (2006) 1505S. In addition, in it was disclosed that a phospholipid emulsion derived from a marine and/or synthetic origin comprising polyunsaturated fatty acids have anti-inflammatory and/or immuno-suppressive effects. See, e.g., 5,434,183. An embodiment of this invention is a krill oil composition effective for reducing inflammation i.e. reducing the levels of TNF-α, IL-1 beta, IL-6, IL-10, TGF beta and fibrinogen in the blood.

Type 2 diabetes is a metabolic disorder characterized by impaired glycemic control (high blood glucose levels). In type 2 diabetes, it is the tissue wide insulin resistance that contributes to the development of the disease. Strategies reducing insulin resistance or improving tissue sensitivity to insulin are recognized as beneficial in preventing type 2 diabetes. In healthy humans, a 3-week supplementation with fish oil (1.1 g EPA/d and 0.7 g DHA/d) decreased the insulin response to an oral glucose load by 40%. Omega-3 PUFA dietary enrichment resulted in lower glucose oxidation, higher fat oxidation, and increased glycogen storage; the glycemic response was unchanged, however, which indicates an improved sensitivity to insulin. In another embodiment of this invention is a krill oil composition effective for reducing the insulin resistance.

Krill oil has not been disclosed as being effective in treating one of the most important life style problems of modern societies, i.e., excess weight gain and obesity. Excess adipose tissue mass (overweight and obesity) is associated with low grade inflammation in adipose tissue and in the whole body reflecting the inflammatory mediators "spilling over" from fat tissue. Travhurn et al., Br. J. Nutrition (2004), 92(3), 347-355. Inflammation appears to be an important link between obesity and metabolic syndrome/type-II diabetes as well as cardiovascular disease. Libby et al., J. Amer. Coll. Card. (2006), 48(9, Suppl. A), A33-A46. Thus, excess adipose tissue is an unhealthy condition. Weight reduction will improve the inflammatory condition, but persistent weight reduction is difficult to achieve. Omega-3 fatty acid supplementation may alleviate the inflammatory condition in adipose tissue and thus ideally complement the principal strategies of weight reduction i.e. low calorie diet and exercise. There are clinical studies in humans that demonstrate that omega-3 enhance the effect of very low calorie diet and exercise in reducing body fat mass. Kunesova et al., Physiological research / Academia Scientiarum Bohemoslovaca (2006), 55(1), 63-72. Although diet and exercise regime may fail to result in consistent decrease in weight in long term, the effect of omega-3 fatty acids alleviating the inflammatory condition in the adipose tissue may persist generating a condition that can be described as "healthy adipose tissue". Previously, it was shown that dietary omega-3 fatty acids can be used to reduce inflammation in adipose tissue without influencing level of obesity. Todoric et al., Diabetologia (2006), 49(9), 2109-2119. Reduction in adipose tissue inflammation was demonstrated by an increase in circulating levels of adiponectin. Adiponectin is an adipose tissue derived anti-inflammatory hormone. Results on the treatment of obese people with omega-3 fatty acids to alleviate circulating levels of inflammatory markers are inconclusive. Trebble et al., Br. J. Nutrition (2003), 90(2), 405-412. However, duration of these studies may not have been sufficient given the slow turnover of adipose tissue in humans. Itoh et al. found that 1.8 g/d of EPA increased adiponectin, a marker of adipose tissue derived inflammation, in a group of overweight subjects with metabolic syndrome. Itoh et al., Arteriosclerosis, Thrombosis, and Vascular Biology (2007), 27(9), 1918-1925.

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An embodiment of the invention is the use of krill oil to increase serum adiponectin levels. Adiponectin is a protein hormone that modulates a number of metabolic processes, including glucose regulation and fatty acid catabolism. Adiponectin is exclusively secreted from adipose tissue into the bloodstream and is very abundant in plasma relative to many hormones.

Levels of the hormone are inversely correlated with body mass index (BMI). The hormone plays a role in alleviating the metabolic dysregulation that may result in type 2 diabetes, obesity, atherosclerosis and non-alcoholic fatty liver disease (NAFLD). Díez et al., Eur. J. Endocrinol. 148 (3): 293-300; Ukkola et al., J. Mol. Med. 80 (11): 696-702.

Another embodiment of the invention is to use krill oil in an overweight and obese subjects for alleviating diet induced adipose tissue dysfunction and diet induced changes in the lipid metabolism.

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In further embodiments, krill oil is effective in reducing risk factors of type 2 diabetes such as hyperinsulinemia and insulin resistance and cardiovascular disease risk factors in overweight subjects. In addition this invention discloses that krill oil is effective in preventing accumulation of fat in muscles and in the liver (liver steatosis).

It is well known in the art that the obese Zucker rat is a useful rat model to study metabolic Syndrome X and non-insulin dependent diabetes mellitus, including glucose tolerance, insulin resistance and hyperinsulinaemia. It has also been shown previously that astaxanthin is a powerful antioxidant, useful for prevention of oxidative stress in vivo and in Zucker rats using vitamin E. See, e.g., Aoi et al., (2003). Antioxidants & Redox Signaling. 5(1):139-44; Laight et al., Eur. J. Pharmacol. 377 (1999) 89.

In yet another embodiment of the invention is a krill oil composition effective of improving the blood lipid profile by increasing the HDL cholesterol levels, decreasing the LDL cholesterol and triglyceride levels. Hence the novel krill oil composition is effective for treating metabolic syndrome. Metabolic syndrome is defined as the coexistence of 3 or more components selected from the group: abdominal obesity, high serum triglyceride levels, low HDL levels, elevated blood pressure and high fasting plasma glucose levels.

In another embodiment of the invention, the krill oil compositions are found to be effective and safe for the treatment of metabolic syndrome in humans.

In still other embodiments, the krill oil compositions of the present invention find use in increasing or inducing diuresis. In some embodiments, the krill oil compositions of the present invention find use in decreasing protein catabolism and increasing the muscle mass of a subject.

In some embodiments, the kill oil composition of the present invention find use in the treatment of fatty heart disease and non-alcoholic fatty acid liver disease. Thus, the krill oil

compositions are useful for decreasing the lipid content of the heart and/or liver and/or muscle of a subject.

In yet another embodiment of the invention is a method to increase the transfer of DHA to the brain.

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### **EXAMPLE 1**

Antarctic krill (*Euphausia superba*) was captured and brought on board alive, before it was processed into krill meal, an oil (asta oil) and stickwater. The composition and properties of the krill meal was monitored during the processing and compared to a commercial competitor (Table 1 and 2). Furthermore, the amino acid composition of the krill meal and stickwater was determined (Table 3), showing that krill meal is a suitable feed source for to be used in aquaculture due to the presences of all the essential amino acids teleost fish require. During the krill meal processing a neutral oil (asta oil) is recovered, the chemical composition of the asta oil is shown in Tables 4 and 5.

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Table 1. Composition of products from the processing line

	Round frozen krill	After decanter	After drier	Konstruktor Koshkin (Ukranian vessel)
Protein	13,5 g/100 g	20,9 g/100 g	58,5 g/100 g	60,2 g/100 g
Moisture	76,3 g/100 g	65,6 g/100 g	9,1 g/100 g	9,6 g/100 g
Lipid (Folch)	8,6 g/100 g	10 g/100 g	21,8 g/100 g	21,4 g/100 g
Free fatty acids	29,8 g/100 g	25,3 g/100 g	24,8 g/100 g	23,3 g/100 g
Total	53,3 mg/kg	81,3 mg/kg	145 mg/kg	126 mg/kg
astaxanthin				

**Table 2.** Lipid class composition in products from the processing line

Crude protein	Round frozen	After decanter	After drier	Konstruktor
	krill (g/100 g)	(g/100 g)	(g/100 g)	Koshkin
				(Ukranian
				vessel) (g/100
				g)
Wax ester/cholesterol	2,5	3,0	1,9	3,3
ester				
Triglycerides/pigments	30,2	33,7	29,3	32,2
Free fatty acids	15,1	2,5	9,0	5,9
Monoglycerides	3,9	Nd	1,3	Nd
PE	6,6	10,4	7,9	6,3

PS	1,2	1,6	1,4	2,7
PI	1,9	2,0	2,1	3,5
PC	28	35,9	32,0	32,1
Sphingomyeline/lyso PC	2,0	0,5	3,0	3,0

Nd= not detected

Table 3. Amino acids in krill meal and stick water

Amino acid	Total in meal	Free in meal	Free in stickwater (g/100
	(g/100 g	(g/100g	g protein)
	protein)	protein)	
Aspartic acid	10,5	0,02	0,22
Glutamic acid	13,5	0,007	0,51
Hydroxiproline	<0,5	<0,001	<0,05
Serine	4,2	0,02	0,13
Glycine	4,4	0,18	3,28
Histidine	2,1	<0,01	<0,05
Arginine	6,7	0,56	4,86
Threonine	4,1	<0,01	0,22
Alanine	5,4	0,08	0,87
Proline	3,8	0,53	2,32
Tyrosine	4,0	0,01	0,2
Valine	5,0	0,02	0,13
Methionine	2,9	<0,01	0,12
Isoleucine	5,0	0,02	0,1
Leucine	7,8	0,14	0,19
Phenylalanine	4,4	0,01	0,1
Lysine	7,8	0,02	0,27
Cysteine/Cystine	1,4	<0,01	<0,05
Thryptophan	1,1	<0,02	<0,05
Creatinine		<0,01	<0,05
Asparagine		<0,01	0,05

Glutamine	<0,01	<0,05
3-aminopropanoic acid	0,5	8,99
Taurine	0,5	8,52
4-aminobutanoic acid	<0,01	<0,05
Citrulline	0,04	0,14
Carnosine	<0,01	<0,05
Anserine	<0,01	<0,05
Ornithine	0,02	1,04

<sup>3-</sup>aminopropanoic acid is also known as  $\beta$ -alanine

**Table 4.** Composition and quality parameters of asta oil.

Moisture	0,14 g/100 g
Insoluble impurities	0,02 g/100 g
Unsaponifable matter	1,5 g/100 g
Nitrogen	0,5 g/100 g
Free fatty acids	0,3 g/100 g
Peroxide value	<2 meq peroxide/kg oil
Ansidine value	<1
Phosphorous	23 mg/kg
Phopspholipids	575 mg/kg
Astaxanthin	1245 mg/kg

**Table 5.** Fatty acid composition of the asta oil

Fatty Acid	Asta oil
File	
C4:0	0,00
C6:0	0,00
C8:0	0,00
C10:0	0,00
C12:0	0,00
C14:0	17,5
C14:1	0,00
C15:0	0,00
C16:0	19,3
C16:1	9,7
C18:0	1,2
C18:1	22,6
C18:2N6	1,4
C18:3N6	0,1

<sup>4-</sup>aminobutanoic acid is alos known as  $\gamma$ -aminobutyric acid or GABA

C18:3N3	0,7
C18:4N3	3,0
C20:0	0,1
C20:1	1,3
C20:2N6	<0,1
C20:3N6	0,1
C20:4N6	0,1
C20:3N3	<0,1
C20:4N3	0,2
C20:5N3 (EPA)	5,6
C22:0	0,1
C22:1	0,3
C22:2N6	0,0
C22:4N6	<0,1
C22:5N6	0,00
C22:5N3	0,2
C22:6N3 (DHA)	2,00
C24:1	0,03
Total	88,4
Saturated	38,0
Monounsaturated	33,9
Polyunsaturated	16,4
Total	88,4
Omega-3	11,9
Omega-6	1,6

#### **EXAMPLE 2**

The krill meal obtained in example 1 was then ethanol extracted according to the method disclosed in JP02215351. The results showed that around 22% fat from the meal could be extracted, somewhat lower than was extracted using Folch (25%). Table 6 shows the fatty acid composition of the krill meal and the krill oil extracted from the meal using ethanol. Table 7 shows the composition and properties of the krill meal and products before and after extraction, whereas table 8 shows the lipid composition.

**Table 6.** Fatty acid distribution in krill meal (g/100 g lipid) and the ethanol extracted krill oil.

Fatty Acid	Krill meal	EtOH KO
	IXI III III Cai	
File	0.00	
C4:0	0,00	
C6:0	0,00	
C8:0	0,00	
C10:0	0,00	
C12:0	0,00	
C14:0	7,8	6,4
C14:1	0,00	
C15:0	0,00	14.7
C16:0	15,8	14,7
C16:1	5,1	4,2
C18:0	0,9	0,7
C18:1	13,4	11,8
C18:2N6	1,1	1,2
C18:3N6	0,1	0,1
C18:3N3	0,4	0,4
C18:4N3	1,1	0,1
C20:0	0,1	0,1
C20:1	0,8	0,6
C20:2N6	<0,1	<0,1
C20:3N6 C20:4N6	0,1	<0,1 0,2
C20:4N0 C20:3N3	0,2 <0,1	<0,1
C20:4N3	0,1	0,2
C20.4N3 C20:5N3 (EPA)	10,5	10,4
C20.5N3 (E1 A) C22:0	<0,1	<0,1
C22:1	0,5	0,4
C22:2N6	<0,1	<0,1
C22:4N6	<0,1	, i
C22:5N6	0,00	
C22:5N3	0,2	
C22:6N3 (DHA)	5,4	4,8
C24:1	0,03	1,0
C24.1	0,03	
Saturated	24,6	21,9
Monounsaturated	19,9	17,0
Polyunsaturated	21,0	19,4
	,	,
Total	65,5	58,2

Omega-3	18,2	17,0
Omega-6	1,3	

**Table 7.** Composition and properties of the krill meal and products after extraction

	Krill meal	Delipidated krill	EtOH extracted krill
		meal	oil
Crude protein	586 g/kg	735 g/kg	
Fat (Folch)	250 g/kg	30 g/kg	
Moisture/ethanol	71 g/kg	134 g/kg	85 g/kg
Astaxanthin esters	144 mg/kg	10 mg/kg	117 mg/kg
Diesters	110 mg/kg	8,5 mg/kg	117 mg/kg
Monoesters	33 mg/kg	1,8 mg/kg	37 mg/kg
Biological digestable	854 g/kg protein	870 g/kg protein	
protein			
Flow number	4,8	1,9	
NH3	9 mg N/100 g	0	3 mg N/100 g
TMA	2 mg N/100 g	0	70 mg N/100 g
TMAO	125 mg N/100 g	0	456 mg N/100 g

# 5 Table 8. Lipid class distribution

Table 6. Lipid class			
	Krill meal	Delipidated krill	EtOH extracted KO
		meal	
Cholesterol ester	3,5		
TG	32,7	37,4	31,1
FFA	7,8	14,1	16,0
Cholesterol	9,1	8,0	12,6
DG	1,1		3,3
MG	3,7		
Sphingolipid			2,8
PE	6,5	2,5	2,7
Cardiolipin		4,2	
PI	1,1	11,0	
PS	1,4		
PC	28,6	20,2	25,3
LPC	2,9	2,6	6,2
Total polar lipids	40,6	40,5	36,9

Total neutral lipids	54.2	59.5	63.1
1 0 0001 11 0 0 0 1 1 1 p 1 0 0	· ·,-	,-	1 00,1

#### **EXAMPLE 3**

The krill meal obtained in example 1 was then subjected to a supercritical fluid extraction method in two stages. During stage 1, 12.1% fat (neutral krill oil) was removed using neat CO<sub>2</sub> only at 300 bars, 60° C and for 30 minutes. In stage 2, the pressure was increased to 400 bar and 20% ethanol was added (v/v) for 90 minutes. This resulted in further extraction of 9% polar fat which hereafter is called polar krill oil. The total fatty acid composition of the polar krill oil, the neutral krill oil and a commercial product obtained from Neptune Biotech (Laval, Quebec, Canada) are listed in Table 9. In addition the fatty acid composition for the phospholipids (Table 10), the neutral lipids (Table 11), the free fatty acids, diglycerides (Table 12), triglycerides, lysophosphatidylcholine (LPC) (Table 13), phosphatidylcholine (PC), phosphatidylethanolamine (PE) (Table 14), phosphatidylinositol (PI) and phosphatidylserine (PS) (Table 15) are shown. Table 16 shows the level of astaxanthin and cholesterol for the different fractions.

**Table 9.** Total fatty acids compositions of the krill oil products (% (w/w))

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	Tot	Total Fatty Acids						
Fatty Acid	Neutral KO	Polar KO	NKO					
File								
C4:0	0,00	0,00	0,00					
C6:0	0,00	0,00	0,00					
C8:0	0,00	0,00	0,00					
C10:0	0,00	0,00	0,00					
C12:0	0,47	0,04	0,24					
C14:0	22,08	3,28	12,48					
C14:1	0,33	0,01	0,17					
C15:0	0,58	0,36	0,52					
C16:0	27,03	29,25	23,25					
C16:1	0,07	0,01	8,44					
C18:0	1,72	1,03	1,42					
C18:1	30,29	13,57	18,92					
C18:2N6	2,10	1,96	1,71					
C18:3N6	0,30	0,21	0,00					
C18:3N3	0,69	1,02	1,32					
C18:4N3	0,05	1,81	3,50					
C20:0	0,06	0,00	0,05					
C20:1	1,87	0,80	1,16					
C20:2N6	0,05	0,05	0,05					

C20:3N6	0,22	0,73	0,04
C20:4N6	0,00	0,00	0,49
C20:3N3	0,09	0,09	0,06
C20:4N3	0,24	0,51	0,33
C20:5N3 (EPA)	7,33	29,88	16,27
C22:0	0,01	0,06	0,05
C22:1	0,64	1,78	0,82
C22:2N6	0,00	0,00	0,00
C22:4N6	0,00	0,00	0,07
C22:5N6	0,00	0,03	0,00
C22:5N3	0,21	0,67	0,36
C22:6N3 (DHA)	3,51	12,61	8,17
C24:0	0,05	0,00	0,01
C24:1	0,03	0,25	0,11
Total	100,00	100,00	100,00
Saturated	52,00	34,01	38,01
Monounsaturated	33,22	16,43	29,61
Polyunsaturated	14,77	49,56	32,37
Total	100,00	100,00	100,00
Omega-3 Omega-6	12,11 2,67	46,58 2,98	30,02 2,35

**Table 10.** Fatty acid composition of the phospholipid fraction ((w/w)).

	Total Phospholipid						
Fatty Acid	Neutral Polar Neptun KO KO KO						
File							
C4:0	0,00	0,00	0,00				
C6:0	0,00	0,00	0,00				
C8:0	0,00	0,00	0,00				
C10:0	0,00	0,00	0,00				
C12:0	0,00	0,00	0,00				
C14:0	0,01	0,00	0,00				
C14:1	0,42	0,01	0,01				
C15:0	2,52	0,00	0,00				
C16:0	4,73	35,78	32,81				
C16:1	0,19	0,17	0,19				
C18:0	6,31	1,18	1,55				
C18:1	38,40	15,58	13,54				
C18:2N6	4,18	2,16	1,90				
C18:3N6	0,18	0,22	0,19				

C18:3N3	1,02	1,05	1,48
C18:4N3	3,08	1,62	2,15
C20:0	0,27	0,00	0,07
C20:1	2,55	1,02	0,78
C20:2N6	0,19	0,06	0,06
C20:3N6	0,00	0,14	0,10
C20:4N6	0,57	0,62	0,64
C20:3N3	0,43	0,08	0,09
C20:4N3	0,17	0,45	0,42
C20:5N3 (EPA)	20,58	25,53	26,47
C22:0	0,14	0,06	0,00
C22:1	0,00	2,09	1,94
C22:2N6	0,25	0,71	0,85
C22:4N6	0,44	0,00	0,03
C22:5N6	0,11	0,00	0,00
C22:5N3	0,00	0,60	0,63
C22:6N3 (DHA)	10,93	10,30	13,34
C24:0	1,77	0,30	0,37
C24:1	0,59	0,28	0,38
Total	100,00	100,00	100,00
Saturated	15,74	37,32	34,81
Monounsaturated	42,14	19,15	16,84
Polyunsaturated	42,12	43,53	48,34
Total	100,00	100,00	100,00
Omega-3 Omega-6	36,22 5,91	39,62 3,90	44,56 3,78
Omega-o	],71	3,50	3,70

Table 11. Fatty acid composition of the total neutral lipid fraction (% (w/w)).

	Total neutral lipid						
Fatty Acid	Neutral KO	Neptune KO					
File							
C4:0	0,00	0,00	0,00				
C6:0	0,00	0,00	0,00				
C8:0	0,00	0,00	0,00				
C10:0	0,00	0,00	0,00				
C12:0	0,00	0,00	0,00				
C14:0	20,35	11,31	18,44				
C14:1	0,30	0,29	0,25				
C15:0	0,53	1,53	0,62				

C16:0	23,79	0,49	24,11
C16:1	12,42	5,22	11,86
C18:0	1,54	3,27	1,67
C18:1	26,81	33,09	23,82
C18:2N6	1,68	2,37	1,79
C18:3N6	0,20	0,23	0,25
C18:3N3	0,59	0,62	0,03
C18:4N3	0,03	1,27	0,05
C20:0	0,07	0,00	0,06
C20:1	1,63	1,41	1,39
C20:2N6	0,04	0,00	0,05
C20:3N6	0,18	0,94	0,01
C20:4N6	0,00	0,00	0,00
C20:3N3	0,09	0,00	0,01
C20:4N3	0,18	0,41	0,23
C20:5N3 (EPA)	5,88	19,26	9,68
C22:0	0,02	0,00	0,03
C22:1	0,56	0,60	0,53
C22:2N6	0,00	0,00	0,00
C22:4N6	0,00	0,00	0,04
C22:5N6	0,01	0,00	0,00
C22:5N3	0,17	0,27	0,22
C22:6N3 (DHA)	2,74	17,22	4,64
C24:0	0,15	0,00	0,17
C24:1	0,03	0,21	0,06
Total	100,00	100,00	100,00
Saturated	46,45	16,60	45,10
Monounsaturated	41,75	40,82	37,91
Polyunsaturated	11,80	42,59	16,99
<i>y</i>		,	,
Total	100,00	100,00	100,00
Omega-3 Omega-6	9,68 2,11	39,05 3,54	14,86 2,14
Omega-0	<b>∠</b> ,11	3,34	∠,14

Table 12. Fatty acid composition of the diglyceride and free fatty acids (% (w/w)).

		Diglycerides			Free fatty acids		
Fatty Acid	Neutral KO	Polar KO	Neptune KO	Neutral KO	Polar KO	Neptune KO	
File							
C4:0	0,00	0,00	0,00	0,00	0,00	0,00	
C6:0	0,00	0,00	0,00	0,00	0,00	0,00	
C8:0	0,00	0,00	0,00	0,00	0,00	0,00	

C10:0	0,00	0,00	0,00	0,00	0,00	0,00
C12:0	0,00	0,00	0,00	0,00	0,00	0,00
C14:0	13,85	14,35	12,22	5,86	7,19	5,45
C14:1	0,18	0,00	0,17	0,05	0,00	0,08
C15:0	0,49	1,08	0,66	0,46	1,60	0,45
C16:0	23,68	35,24	25,81	28,30	29,37	21,12
C16:1	9,49	6,80	0,09	3,27	3,08	4,91
C18:0	1,56	3,63	1,89	1,13	2,43	0,99
C18:1	23,67	19,85	23,82	14,50	14,77	17,41
C18:2N6	1,79	0,21	1,90	1,69	0,97	1,86
C18:3N6	0,17	0,00	0,01	0,14	0,00	0,22
C18:3N3	0,69	0,00	1,19	0,85	0,00	1,34
C18:4N3	1,92	0,00	2,75	1,30	0,00	2,72
C20:0	0,00	0,00	0,00	0,00	0,00	0,00
C20:1	1,09	0,00	1,01	0,48	0,00	0,57
C20:2N6	0,00	0,00	0,00	0,00	0,00	0,00
C20:3N6	0,13	0,00	0,00	0,08	0,00	0,05
C20:4N6	0,45	0,00	0,64	0,78	0,00	1,43
C20:3N3	0,00	0,00	0,00	0,00	0,00	0,00
C20:4N3	0,35	0,00	0,43	0,39	0,00	0,43
C20:5N3 (EPA)	14,03	9,80	18,00	24,33	23,57	25,36
C22:0	0,18	0,00	0,10	0,00	0,00	0,05
C22:1	0,41	0,00	0,57	0,80	0,69	0,37
C22:2N6	0,28	0,00	0,50	0,46	0,00	0,54
C22:4N6	0,00	0,00	0,00	0,00	0,00	0,00
C22:5N6	0,00	0,00	0,00	0,00	0,00	0,00
C22:5N3	0,20	0,00	0,27	0,34	0,00	0,32
C22:6N3 (DHA)	4,74	9,04	7,53	14,31	16,33	13,95
C24:0	0,64	0,00	0,42	0,49	0,00	0,39
C24:1	0,00	0,00	0,00	0,00	0,00	0,00
Total	100,00	100,00	100,00	100,00	100,00	100,00
Saturated	40,40	54,30	41,10	36,24	40,59	28,45
Monounsaturated	34,84	26,64	25,66	19,09	18,54	23,34
Polyunsaturated	24,77	19,06	33,24	44,67	40,87	48,22
Total	100,00	100,00	100,00	100,00	100,00	100,00
Omega-3	21,95	18,85	30,18	41,51	39,90	44,13
Omega-6	2,82	0,21	3,05	3,15	0,97	4,09

**Table 13.** Fatty acid composition of the triglyceride and lyso-phophatidylcholine fractions (% (w/w)).

	Triglycerides		Lyso PC			
Fatty Acid	Neutral KO	Polar KO	Neptune KO	Neutral KO	Polar KO	Neptune KO
File						
C4:0	0,00	0,00	0,00	0,00	0,00	0,00
C6:0	0,00	0,00	0,00	0,00	0,00	0,00
C8:0	0,00	0,00	0,00	0,00	0,00	0,00
C10:0	0,00	0,00	0,00	0,00	0,00	0,00
C12:0	0,00	0,00	0,00	0,00	0,00	0,00
C14:0	23,06	26,65	25,13	19,38	4,27	2,87
C14:1	0,36	0,93	0,36	0,00	0,08	0,00
C15:0	0,56	2,64	0,78	0,00	0,52	0,45
C16:0	23,17	4,93	27,80	41,00	44,14	30,56
C16:1	13,68	11,58	0,04	0,00	1,84	2,24
C18:0	1,52	3,12	1,99	0,76	1,59	1,32
C18:1	27,83	34,39	27,92	6,65	14,24	11,29
C18:2N6	1,64	2,05	1,92	0,00	1,75	2,07
C18:3N6	0,20	0,00	0,30	0,00	0,00	0,06
C18:3N3	0,51	0,00	0,00	7,95	0,67	1,75
C18:4N3	1,99	0,00	4,83	0,00	1,11	2,46
C20:0	0,06	0,00	0,08	0,00	0,00	0,00
C20:1	1,67	0,00	1,76	0,00	0,52	0,00
C20:2N6	0,04	0,00	0,05	0,00	0,00	0,00
C20:3N6	0,05	0,00	0,01	0,00	0,00	0,54
C20:4N6	0,00	0,00	0,00	0,00	0,40	0,00
C20:3N3	0,05	0,00	0,07	0,00	0,00	0,00
C20:4N3	0,11	0,00	0,17	0,00	0,31	0,55
C20:5N3 (EPA)	2,10	7,97	4,44	0,00	18,59	28,48
C22:0	0,02	0,00	0,04	0,00	0,00	0,00
C22:1	0,37	0,00	0,42	0,00	1,46	0,91
C22:2N6	0,00	0,00	0,00	0,00	0,00	0,00
C22:4N6	0,01	0,00	0,01	0,00	0,00	0,00
C22:5N6	0,00	0,00	0,01	0,00	0,00	0,00
C22:5N3	0,10	0,00	0,16	0,00	0,41	0,62
C22:6N3 (DHA)	0,67	3,97	1,42	24,26	7,79	13,82
C24:0	0,26	1,78	0,26	0,00	0,32	0,00
C24:1	0,00	0,00	0,03	0,00	0,00	0,00
Total	100,00	100,00	100,00	100,00	100,00	100,00
Saturated	48,64	39,12	56,08	61,14	50,83	35,21
Monounsaturated	43,90	46,89	30,52	6,65	18,14	14,44
Polyunsaturated	7,45	13,99	13,41	32,20	31,02	50,35
Total	100,00	100,00	100,00	100,00	100,00	100,00

Omega-3	5,51	11,94	11,11	32,20	28,87	47,69
Omega-6	1,94	2,05	2,30	0,00	2,15	2,66

**Table 14.** Fatty acid composition of the phosphatidylcholine and the phosphatidylserine fractions (% (w/w)).

		PC			PS	
Fatty Acid	Neutral KO	Polar KO	Neptune KO	Neutral KO	Polar KO	Neptune KO
File						
C4:0	0,00	0,00	0,00	0,00	0,00	0,00
C6:0	0,00	0,00	0,00	0,00	0,00	0,00
C8:0	0,00	0,00	0,00	0,00	0,00	0,00
C10:0	0,00	0,00	0,00	0,00	0,00	0,00
C12:0	0,00	0,00	0,00	0,00	0,00	0,00
C14:0	0,75	3,29	2,77	7,60	9,52	2,31
C14:1	2,07	0,04	0,02	0,00	0,00	0,00
C15:0	1,34	0,00	0,00	3,83	0,00	0,00
C16:0	16,65	31,92	29,83	30,44	43,61	19,49
C16:1	0,96	0,01	0,17	9,96	3,47	2,79
C18:0	1,33	1,06	1,33	2,08	3,34	2,24
C18:1	34,34	13,55	11,16	0,00	7,37	11,87
C18:2N6	10,55	2,27	1,90	0,00	0,00	0,00
C18:3N6	1,44	0,25	0,20	0,00	0,00	0,00
C18:3N3	2,49	1,19	1,54	0,00	0,00	0,00
C18:4N3	2,38	1,92	2,41	0,00	0,00	0,00
C20:0	2,79	0,03	0,05	0,00	0,00	0,00
C20:1	2,42	0,82	0,74	0,00	0,00	0,00
C20:2N6	0,56	0,05	0,06	0,00	0,00	0,00
C20:3N6	0,67	0,13	0,09	0,00	0,00	0,00
C20:4N6	1,85	0,61	0,56	0,00	0,00	0,00
C20:3N3	3,94	0,07	0,06	0,00	0,00	0,33
C20:4N3	4,32	0,50	0,46	0,00	0,00	0,00
C20:5N3 (EPA)	1,08	29,85	30,09	25,84	15,81	16,35
C22:0	0,00	0,05	0,02	0,00	0,00	0,00
C22:1	2,77	0,00	1,87	0,00	0,00	0,00
C22:2N6	0,00	0,81	0,97	0,00	0,00	0,00
C22:4N6	0,00	0,01	0,02	0,00	0,00	0,00
C22:5N6	1,49	0,01	0,00	0,00	0,00	0,00
C22:5N3	1,48	0,67	0,68	0,00	0,00	0,00
C22:6N3 (DHA)	0,00	10,53	12,49	20,25	16,89	44,63
C24:0	2,34	0,10	0,18	0,00	0,00	0,00
C24:1	0,00	0,25	0,34	0,00	0,00	0,00
Total	100,00	100,00	100,00	100,00	100,00	100,00

Saturated	25,19	36,46	34,18	43,95	56,47	24,04
Monounsaturated	42,56	14,67	14,29	9,96	10,84	14,65
Polyunsaturated	32,25	48,87	51,53	46,09	32,69	61,31
Total	100,00	100,00	100,00	100,00	100,00	100,00
Omega-3	15,69	44,73	47,73	46,09	32,69	61,31
Omega-6	16,56	4,13	3,81	0,00	0,00	0,00

**Table 15.** Fatty acid composition of the phosphatidylinositol and phophatidylethanolamine fractions ( $\frac{6}{6}$  (w/w)).

		PΙ			PE	
Fatty Acid	Neutral KO	Polar KO	Neptune KO	Neutral KO	Polar KO	Neptune KO
File						
C4:0	0,00	0,00	0,00	0,00	0,00	0,00
C6:0	0,00	0,00	0,00	0,00	0,00	0,00
C8:0	0,00	0,00	0,00	0,00	0,00	0,00
C10:0	0,00	0,00	0,00	0,00	0,00	0,00
C12:0	0,00	0,00	0,00	0,00	0,00	0,00
C14:0	11,15	5,82	5,72	14,42	4,60	0,83
C14:1	3,03	0,66	0,00	0,00	0,00	0,10
C15:0	5,86	1,95	3,18	0,00	1,30	0,23
C16:0	37,02	30,66	31,39	35,91	31,21	18,38
C16:1	18,05	2,24	1,16	0,00	1,51	0,75
C18:0	6,72	2,83	5,56	12,72	16,70	1,84
C18:1	18,15	24,77	14,23	36,96	19,91	18,45
C18:2N6	0,00	2,67	0,00	0,00	2,62	0,85
C18:3N6	0,00	0,00	0,00	0,00	0,00	0,00
C18:3N3	0,00	0,00	0,00	0,00	0,00	0,33
C18:4N3	0,00	0,00	0,00	0,00	0,00	0,00
C20:0	0,00	0,00	0,00	0,00	0,00	0,00
C20:1	0,00	0,00	0,00	0,00	0,00	0,00
C20:2N6	0,00	0,00	0,00	0,00	0,00	0,00
C20:3N6	0,00	0,00	0,00	0,00	0,00	1,15
C20:4N6	0,00	0,00	0,00	0,00	0,00	0,00
C20:3N3	0,00	0,00	0,00	0,00	0,00	0,00
C20:4N3	0,00	0,00	0,00	0,00	0,00	0,00
C20:5N3 (EPA)	0,00	17,60	20,45	0,00	10,76	21,26
C22:0	0,00	0,00	0,00	0,00	0,00	0,00
C22:1	0,00	0,00	0,00	0,00	0,00	0,00

C22:2N6	0,00	0,00	0,00	0,00	0,00	0,00
C22:4N6	0,00	0,00	0,00	0,00	0,00	0,00
C22:5N6	0,00	0,00	0,00	0,00	0,00	0,00
C22:5N3	0,00	0,00	0,00	0,00	0,00	0,67
C22:6N3 (DHA)	0,00	10,79	18,32	0,00	11,39	35,16
C24:0	0,00	0,00	0,00	0,00	0,00	0,00
C24:1	0,00	0,00	0,00	0,00	0,00	0,00
Total	100,00	100,00	100,00	100,00	100,00	100,00
Saturated	60,76	41,26	45,84	63,04	53,81	21,28
Monounsaturated	39,24	27,67	15,39	36,96	21,42	19,30
Polyunsaturated	0,00	31,07	38,77	0,00	24,77	59,42
Total	100,00	100,00	100,00	100,00	100,00	100,00
Omega-3 Omega-6	0,00 0,00	28,40 2,67	38,77 0,00	0,00 0,00	22,15 2,62	57,43 1,99

**Table 16.** Compositional data for the novel krill oil composition obtained and NKO krill oil.

Compounds	Neptune KO	Ethanol	Polar KO	Neutral KO
		extracted KO		
Astaxanthin esters	472 mg/kg	117 mg/kg	580 mg/kg	98 mg/kg
Astaxanthin free	11 mg/kg	< 1 mg/kg	<1 mg/kg	<1 mg/kg
Total cholesterol	1 g/100g	12 g/100g	< 0,5 g/100g	5,7 g/100g

#### **EXAMPLE 4**

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Neutral lipids were extracted from krill meal (138 kg) using SFE with neat CO<sub>2</sub> (solvent ratio 25 kg/kg) at 500 bar and 75 °C. The neutral lipids were fractionated at 200 bar (75 °C) and at 60 bar (35 °C) at separator S1 and S2, respectively. The extract obtained in S1 (19,6 kg) were characterized and the results can be found in Tables 17A-C. The extract in table S2 (0,4 kg) were rich in water and were not further used. Next, the polar lipids were extracted using CO<sub>2</sub> at 500 bar, 20% ethanol and at a temperature of 75 °C. Using a solvent ratio of 32 (kg/kg) and collecting an extract of 18,2 kg using a separator at 60 bars and 35°C. The polar lipids were collected and analyzed (Tables 18A-C). Next, the polar lipids were mixed in a 50/50 ratio with the neutral

lipids collected from S1 before finally the ethanol was removed carefully by evaporation. The product obtained was red and transparent. If the ethanol is removed before the mixing if the fractions a transparent product is not obtained. The composition of the 50/50 red and transparent product can be found in Tables 19A-C.

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Table 17A Fatty acid composition of the extract collected in S1

Fatty acid	Unit	Amount
14:0	g/100g	18,4
16:0	g/100g	22,2
18:0	g/100g	1,5
16:1 n-7	g/100g	10,9
18:1 (n-9) + (n-7) + (n-5)	g/100g	25,6
20:1 (n-9) + (n-7)	g/100g	1,8
22:1 (n-11) + (n-9) + (n-7)	g/100g	0,5
16:2 (n-4)	g/100g	1,3
16:4 (n-1)	g/100g	1,2
18:2 n-6	g/100g	1,3
18:3 n-3	g/100g	0,8
18:4 n-3	g/100g	2,9
20:5 n-3	g/100g	4,1
22:6 n-4	g/100g	1,7

Table 17B. Lipid class composition of the extract collected in S1

Lipid	Unit	Amount
Triacylglycerol	g/100g	84
Diacylglycerol	g/100g	0,7
Free fatty acids	g/100g	1,5
Cholesterol	g/100g	2,7
Cholesterol esters	g/100g	0,9

10 **Table 17C.** Miscellaneous analysis of the extract in S1.

Compound	Unit	Amount
Free astaxanthin	mg/kg	4,3
Astaxanthin esters	mg/kg	462
Trimethylamin	mg N/100 g	<1
Trimethylamineoxide	mg N/100 g	2

Table 18A Fatty acid composition of the extract collected after CO<sub>2</sub> and 20% ethanol in S1.

Fatty acid	Unit	Amount
14:0	g/100g	1,3
16:0	g/100g	13,8
18:0	g/100g	0,6
16:1 n-7	g/100g	0,9
18:1 (n-9) + (n-7) + (n-5)	g/100g	6,5
20:1 (n-9) + (n-7)	g/100g	0,6
22:1 (n-11) + (n-9) + (n-7)	g/100g	0,1
16:2 (n-4)	g/100g	<0,1
16:4 (n-1)	g/100g	<0,1
18:2 n-6	g/100g	0,8
18:3 n-3	g/100g	0,6
18:4 n-3	g/100g	1,0
20:5 n-3	g/100g	14,7
22:6 n-4	g/100g	6,5

Table 18B. Lipid class composition of the extract collected after CO<sub>2</sub> and 20% ethanol in S1.

Lipid	Unit	Amount
Triacylglycerol	g/100g	<0,5
Cholesterol	g/100g	<0,5
Phophatidylethanolamine	g/100g	1,6
Phosphatidylcholine	g/100g	67
Lyso-phophatidylcholine	g/100g	4,4

**Table 18C.** Miscellaneous analysis of the extract in S1.

Compound	Unit	Amount
Trimethylamin	mg N/100 g	422
Trimethylamineoxide	mg N/100 g	239

Table 19A Fatty acid composition of the final blended product obtained in Example 4 in S1.

Fatty acid	Unit	Amount
14:0	g/100g	9,7
16:0	g/100g	18,5
18:0	g/100g	1,0
16:1 n-7	g/100g	5,8
18:1 (n-9) + (n-7) + (n-5)	g/100g	16,0
20:1 (n-9) + (n-7)	g/100g	1,2
22:1 (n-11) + (n-9) + (n-7)	g/100g	1,0
16:2 (n-4)	g/100g	0,3
16:4 (n-1)	g/100g	<0,1
18:2 n-6	g/100g	1,0
18:3 n-3	g/100g	0,8
18:4 n-3	g/100g	2,1
20:5 n-3	g/100g	10,7
22:6 n-4	g/100g	4,7

# 5 **Table 19B.** Lipid class composition of the final blended product obtained in Example 4.

Lipid	Unit	Amount
Triacylglycerol	g/100g	53
Diacylglycerol	g/100g	1,3
Free fatty acids	g/100g	0,5
Cholesterol	g/100g	0,6
Cholesterol esters	g/100g	<0,5
Phophatidylethanolamine	g/100g	<1

Phosphatidylcholine	g/100g	42
Lyso-phophatidylcholine	g/100g	5,9

**Table 19C.** Miscellaneous analysis of the final blended product obtained in example 4.

Compound	Unit	Amount
Free astaxanthin	mg/kg	1,1
Astaxanthin esters	mg/kg	151
Trimethylamin	mg N/100 g	109
Trimethylamineoxide	mg N/100 g	80

#### **EXAMPLE 5**

The asta oil obtained in example 1 was blended with the polar lipids obtained in example 4 in a ratio of 46:54 (v/v). Next the ethanol was removed by evaporation and a dark red and transparent product was obtained. The product was analyzed and the results can be found in Tables 20A-C. Furthermore, the product was encapsulated into soft gels successfully. During the encapsulation it was observed that any further increase in phospholipids and thereby viscosity will make it very difficult to encapsulate the final product.

**Table 20A** Fatty acid composition of the final blended product obtained in Example 5.

Fatty acid	Unit	Amount
14:0	g/100g	8,2
16:0	g/100g	17,7
18:0	g/100g	1,0
16:1 n-7	g/100g	4,9
18:1 (n-9) + (n-7) + (n-5)	g/100g	14,9
20:1 (n-9) + (n-7)	g/100g	1,1
22:1 (n-11) + (n-9) + (n-7)	g/100g	1,0
16:2 (n-4)	g/100g	0,4
16:4 (n-1)	g/100g	<0,1
18:2 n-6	g/100g	1,2

18:3 n-3	g/100g	0,8
18:4 n-3	g/100g	1,8
20:5 n-3	g/100g	10,6
22:6 n-4	g/100g	4,8

**Table 20B.** Lipid class composition of the final blended product obtained in Example 5.

Lipid	Unit	Amount
Triacylglycerol	g/100g	41
Diacylglycerol	g/100g	0,8
Free fatty acids	g/100g	1,2
Cholesterol	g/100g	0,4
Cholesterol esters	g/100g	0,3
Phophatidylethanolamine	g/100g	0,6
Phosphatidylcholine	g/100g	51
Lyso-phophatidylcholine	g/100g	<0,5
Total polar lipids	g/100g	52,4
Total neutral lipids	g/100g	43,6

Table 20C. Miscellaneous analysis of the final blended product obtained in Example 5

Compound	Unit	Amount
Free astaxanthin	mg/kg	12
Astaxanthin esters	mg/kg	1302
Trimethylamin	mg N/100 g	193
Trimethylamineoxide	mg N/100 g	1,7

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## **EXAMPLE 6**

Fresh krill was pumped from the harvesting trawl directly into an indirect steam cooker, and heated to 90°C. Water and a small amount of oil were removed in a screw press before

ethoxyquin (antioxidant) was added and the denatured meal was dried under vacuum at a temperature not exceeding 80C. After 19 months storage in room temperature, a sample of the denatured meal was extracted in two steps with supercritical CO<sub>2</sub> in laboratory scale at a flow rate of 2ml/min at 100C and a pressure of 7500 psi. In the second step 20% ethanol was added to the CO<sub>2</sub>. The two fractions collected were combined and analyzed by HPLC using ELS detection. The phosphatidylcholine was measured to 42.22% whereas the partly decomposed phosphatidylcholine was 1.68%. This data strongly contrasts the data obtained by analysis of a krill oil sample in the marketplace that showed a content of 9.05% of phosphatidylcholine and 4.60% of partly decomposed phosphatidylcholine.

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#### **EXAMPLE 7**

Krill lipids were extracted from krill meal (a food grade powder) using supercritical fluid extraction with co-solvent. Initially, 300 bar pressure, 333°K and 5% ethanol (ethanol:CO<sub>2</sub>, w/w) were utilized for 60 minutes in order to remove neutral lipids and astaxanthin from the krill meal. Next, the ethanol content was increased to 23% and the extraction was maintained for 3 hours and 40 minutes. The extract was then evaporated using a falling film evaporator and the resulting krill oil was finally filtered. The product obtained was then analyzed and the results can be found in Table 21.

**Table 21.** Analysis of the krill oil obtained using supercritical fluid extraction.

Parameter	Value
Ethanol	1.11% w/w
Water Content	2.98 % w/w
C20:5 n-3 (EPA)	19.9
C22:6 n-3 (DHA)	11.3
Total Omega 3	35.7
Total Omega 6	3.0
Total Phospholipids	50.55 wt%
Ratio Omega 3-PL/Total Omega 3	77.6 % w/w
Ratio EPA- PL/Total EPA	84.4 %w/w
Ratio DHA-PL/Total DHA	74.7 %w/w
Triglycerides	25.9 g/100g
Astaxanthin	2091 mg/kg
Peroxide Value	<0.1

#### **EXAMPLE 8**

Krill oil was prepared according to the method described in example 7 extracting from the same krill meal. The oil was subjected to <sup>31</sup>P NMR analysis for the identification and quantification of the various forms of phospholipids. The analysis was performed according to the following methods: Samples (20 – 40 mg) were weighed into 1.5 ml centrifuge tubes. Next, NMR detergent (750 µl -10% Na cholate, 1% EDTA, pH 7.0 in H<sub>2</sub>O+D<sub>2</sub>O, 0.3 g L-1 PMG internal standard) was added. Next, the tube was placed in a oven at 60°C and periodically shaken/sonicated until completely dispersed. The solution was then transferred to a 5 ml NMR tube for analysis. Phosphorus NMR spectra were recorded on the two-channel Bruker Avance300 with the following instrument settings: spectrometer frequency 121.498MHz, sweep width 24,271 Hz, 64,000 data points, 30 degree excitation pulse, 576 transients were normally taken, each with an 8 second delay time and f.i.d. acquisition time of 1.35 sec. Spectra were processed with a standard exponential weighting function with 0.2 Hz line broadening before Fourier transformation.

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Peaks were identified using known chemical shifts. Deacylation of samples with monomethylamine was also used on two samples for confirmation of peak identity and to achieve better peak resolution. Example spectra are presented in Figure 1. Peak area integration gave relative molar amounts of each lipid class. Weight percent values were calculated using molecular masses calculated from a krill sample fatty acid profile (average chain length = 18.6). Total PL levels were calculated from the PMG internal standard peak. The quantification of the phospholipids are shown in table 25 for both the raw material, the final product and for a commercially available krill oil (Neptune Krill Oil). The main polar ether lipids of the krill meal are alkylacylphosphatidylcholine (AAPC) at 7-9 % of total polar lipids, lysoalkylacylphosphatidylcholine (LAAPC) at 1 % of total polar lipids (TPL) and alkylacylphosphatidyl-ethanolamine (AAPE) at < 1 % of TPL.

 Table 22:
 Phospholipid profiles

	Type B krill		
	<u>powder</u>	<u>NKO</u>	Krill Oil obtained in Example 7
PC	66.0	68.6	75.3
AAPC	12.0	7.0	13.0
PI			
1LPC	1.2	1.3	0.4
PS			
2LPC	7.4	13.8	2.9
LAAPC	2.2	1.2	0.9
PE	6.0	3.4	3.4
AAPE			1.5
SM			
GPC		1.3	
DHSM			
NAPE		3.4	
CL	5.3		2.1
LPE			0.5
LCL			
% PL in			
powder or			
lipid sample	8.3	30.0	47.9

Analysis has been carried out on the fatty acid and ether/alcohol profiles of the AAPC. The following results are presented in Table 23.

 Table 23. Fatty acid profile of the alkylacylphosphatidylcholine.

AAPC fatty acid composition	AAPC alcohol composition	
	alcohol	%
20:5(n-3) –		
46.9%;	16:0	47.6
22:6(n-3) –		
36.1%;	18:1	17.8
18:1(n-9) - 4.6%	16:1	14.1
22:5(n-3) - 2.6%	14:0	10
20:4(n-6)-1.9%	18:0	8.6

21:5(n-3) - 1.5%	18:2	5.1
18:2(n-6)-0.9%	17:0	4.4
16:1(n-9)-0.8%	15:0-i	2.1
16:0-0.7%	15:0	1.7
phytanic – 0.6%	20:1	1.4
18:3(n-3)-0.5%	15:0-a	1.3
18:4(n-3) - 0.4%	18:0-i	0.4
18:1(n-7)-0.4%		
24:1-0.4%		
14:0-0.3%		

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The rest of alcohols (i17:0, etc.), were less than 0.3% each. Only part of 20:1 was confirmed by GC-MS. Alcohol moieties composition of Krill AAPC was determined (identification was performed in the form of 1-alkyl-2,3-diTMS glycerols on GC-MS, % of total fatty alcohols were obtained by GC with FID). Ten other fatty acids were all below 0.3 % by mass.

#### **EXAMPLE 9**

The purpose of this experiment was to investigate the effect of different omega-3 fatty acid sources on metabolic parameters in the Zucker rat. The Zucker rat is a widely used model of obesity and insulin resistance. Obesity is due to a mutation in the leptin receptor which impairs the regulation of intake. Omega-3 sources compared in this study were fish oil (FO) and two types of krill oil. The krill oil were either from a commercial supplier (Neptune Krill oil) or prepared according to example 7 (Superba<sup>TM</sup>). Four groups of rats (n = 6 per group) were fed ad lib either a control diet (CTRL) or a diet supplemented with a source of omega-3 fatty acids (FO, NKO, Superba). All diets supplied same amount of dietary fatty acids, oleic acid, linoleic acid and linolenic acid. Omega-3 diets (FO, NKO and Superba<sup>TM</sup>) were additionally balanced for EPA and DHA content. The Zucker rats were 4 wk old at the start of the study with average initial weight of 250 g. At this stage the Zucker rats can be characterized as being pre-diabetic. Rats were fed the test diets for 4 wk after which they were sacrificed and blood and tissue samples were collected. Data presented in the following figures are means  $\pm$  SE. This example shows that supplementation of the Zucker rat with krill oil prepared as in example 7 results in an improvement of metabolic parameters characteristic of the obesity induced type two diabetic condition. The effect induced by the novel krill oil is often more pronounced than the effect of

FO an in several cases greater than the effect induced by NKO. Specifically, the effects of the two types of krill oil differentiated with respect to the reduction of blood LDL cholesterol levels as well as lipid accumulation in the liver and muscle (Figure 2-9). Furthermore, the efficacy of transfer of DHA from the diet to the brain tissue was greatest with the krill oil prepared as in example 7 (Figure 10).

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#### **EXAMPLE 11**

This example describes the effect of the supplementation of human diets with krill oil, fish oil (positive control), or a negative control oil (no omega-3 fatty acids) on blood urea nitrogen (BUN).

BUN measures the amount of nitrogen in the blood that comes from urea. BUN is used as a measure of renal function. Serum creatinine is, however, considered to be a more specific measure of renal function. In this study, krill oil decreased BUN by 11.8% while creatinine levels were unchanged. Thus, it is likely that the decrease in BUN is due to some other effect than improved renal function. BUN decreases if krill oil induced diuresis i.e. excretion of urine (diuretic effect).

BUN also decreases if body protein catabolism is reduced. Protein catabolism is a normal feature of body protein turnover. Many tissues express high protein turnover rates. For example the gastrointestinal system expresses high rates of protein turnover. In growing animals a reduction in GI protein catabolism improves weight gain. Mice supplemented with krill oil grew at a faster rate than mice supplemented with fish oil or control diet (Figure 11).

**Table 24.** The effect on blood urea nitrogen in humans for the different treatment groups.

		Control	Krill Oil	Menhaden oil	p
30		n = 23	n = 24	n = 25	
	BUN, mg/dL				
	Baseline	11.5 (7.8, 13.8)	11.5 (9.5, 13.5)	11.5 (9.5, 14.0)	0.523
	$\Delta$ from baseline, %	11.0 (-14.3, 26.1)	-11.8 (-20.0, 1.5)	9.1 (-9.1, 35.7)	0.014r

Creatinine, mg/dL

Baseline	0.9 (0.7, 0.9)	0.9 (0.7, 0.9)	0.8 (0.8, 1.0)	0.952r (r)
$\Delta$ from baseline, %	0.0 (-9.6, 2.9)	0.0 (-2.0, 5.9)	0.0 (-5.9, 6.7)	0.416

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#### **EXAMPLE 12**

The purpose of this experiment was to investigate the effect of dietary krill oil on metabolic parameters in high-fat fed mice and to compare the effect of dietary krill oil with that of fish oil containing the same amount of omega-3 fatty acids. Four groups of C57BL/6 mice (n = 10 per group) were fed 1) chow (N), 2) high fat diet comprising 21% butter fat and 0.15% cholesterol (HF), 3) high fat diet + krill oil (HFKO) or 4) high fat diet + fish oil (HFFO). Treatment 3 contained 2.25% (w/w) krill oil as prepared in example 5 (except that the astaxanthin content was 500 ppm) which were equivalent to 0.36% omega-3 fatty acids. Treatment 4 also contained 0.36% omega-3 fatty acids obtained from regular 18-12 fish oil. The diets were fed to the mice for 7 weeks with free access to drinking water. Data represented in this example means  $\pm$  SE. Columns not sharing a common letter are significantly different (P < 0.05) by ANOVA followed by Tukey's multiple comparison test. N = normal chow diet (n = 10); HF = high-fat diet (n = 10); HFFO = high-fat diet supplemented with fish oil (n = 9); HFKO = high-fat diet supplemented with krill oil (n = 8). The data are presented in Figures 18-25.

This example shows that supplementation of high-fat fed mice with krill oil results in an amelioration of diet-induced hyperinsulinemia, insulin resistance, increase in muscle lipid content (measured as a change in muscle mass), serum adiponectin reduction and hepatic steatosis. These potentially beneficial atheroprotective effects were similar or greater than those achieved with a supplement containing a comparable level of omega-3 fatty acids (Figure 12-19).

#### **CLAIMS**

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- 1. A method of production of krill oil comprising:
  - a) providing krill;
- b) treating said krill to denature lipases and phospholipases in said krill to provide a denatured krill product;
  - c) storing said denatured krill product for a period of from 1 to 24 months;
- d) extracting oil from said denatured krill product with a polar solvent to provide a krill oil with from about 3% to about 15% ether phospholipids w/w of said krill oil astaxanthin esters in amount of greater than about 100 mg/kg of said krill oil.
  - 2. The method of claim 1, wherein said steps a and b are performed on a ship.
- 15 3. The method of claim 1, wherein said treating comprises heating.
  - 4. The method of claim 1, wherein said denatured krill product is a krill meal.
  - 5. The method of claim 1, wherein said krill is freshly harvested.

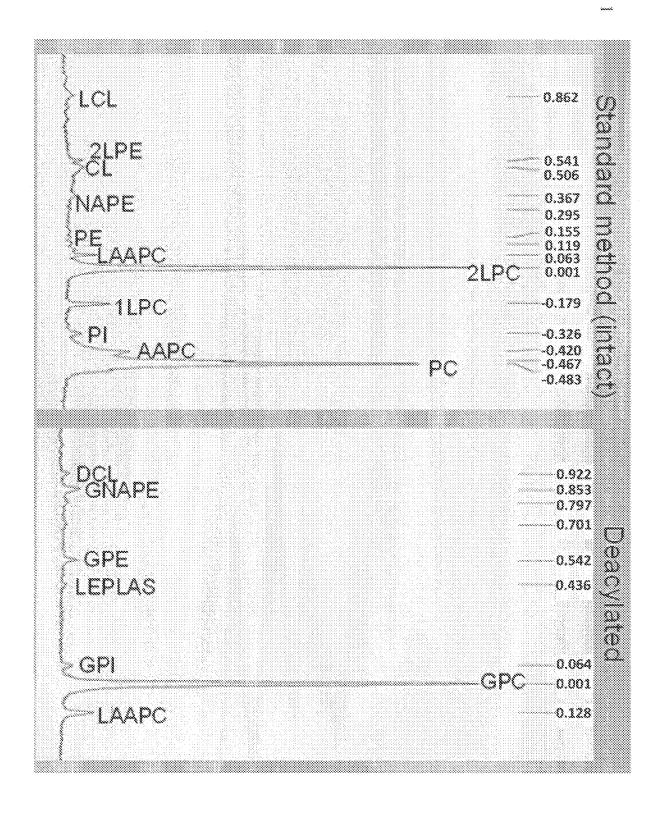
- 6. The method of claim 1, further comprising encapsulating said krill oil.
- 7. The method of claim 1, wherein said krill is Antarctic krill.
- 25 8. The method of claim 1, wherein said Antarctic krill is Euphausia superba.
  - 9. The method of claim 1, wherein said krill oil contains astaxanthin esters in an amount of greater than about 200 mg/kg of said krill oil.
- 30 10. The method of claim 1, wherein said krill oil comprises at least 30% total phospholipids w/w of said krill oil.

- 11. The method of claim 1, wherein said krill oil comprises at least 30% phosphatidylcholine w/w of said krill oil.
- 5 12. A method of production of krill oil comprising:
  - a) obtaining a denatured krill product produced by treating freshly harvested krill krill to denature lipases and phospholipases in said krill and that has been stored from 1 to 24 months; and
- b) extracting oil from said denatured krill product with a polar solvent to provide a krill oil with from about 3% to about 15% ether phospholipids w/w of said krill oil astaxanthin esters in amount of greater than about 100 mg/kg of said krill oil.
  - 13. The method of claim 1, wherein said treating comprises heating.
- 15 14. The method of claim 1, wherein said denatured krill product is a krill meal.
  - 15. The method of claim 1, wherein said krill is freshly harvested.
  - 16. The method of claim 1, further comprising encapsulating said krill oil.
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  - 17. The method of claim 1, wherein said krill is Antarctic krill.
  - 18. The method of claim 10, wherein said Antarctic krill is Euphausia superba.
- The method of claim 1, wherein said krill oil contains astaxanthin esters in an amount of greater than about 200 mg/kg of said krill oil.
  - 20. The method of claim 1, wherein said krill oil comprises at least 30% total phospholipids w/w of said krill oil.

## **ABSTRACT**

This invention discloses new krill oil compositions characterized by having high amounts of phospholipids, astaxanthin esters and/or omega-3 contents. The krill oils are obtained from krill meal using supercritical fluid extraction in a two stage process. Stage 1 removes the neutral lipid by extracting with neat supercritical CO<sub>2</sub> or CO<sub>2</sub> plus approximately 5% of a co-solvent. Stage 2 extracts the actual krill oils by using supercritical CO<sub>2</sub> in combination with approximately 20% ethanol. The krill oil materials obatined are compared with commercially available krill oil and found to be more bioeffective in a number of areas such as anti-inflammation, anti-oxidant effects, improving insulin resistances and improving blood lipid profile.

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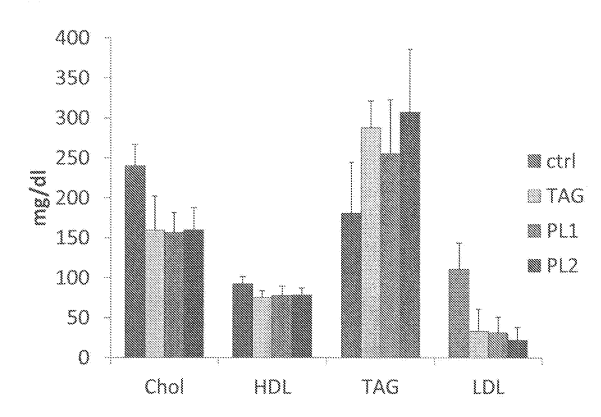


FIGURE 3

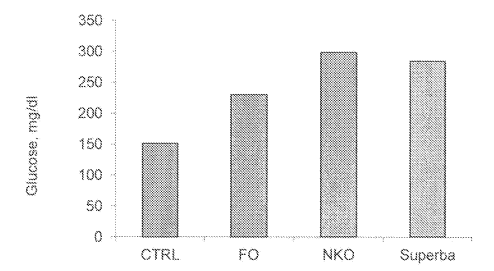


FIGURE 4

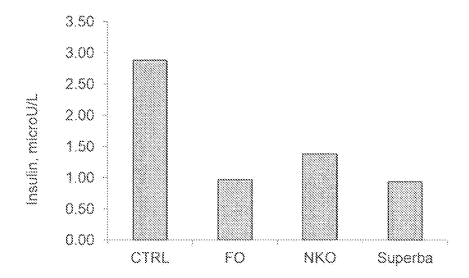


FIGURE 5

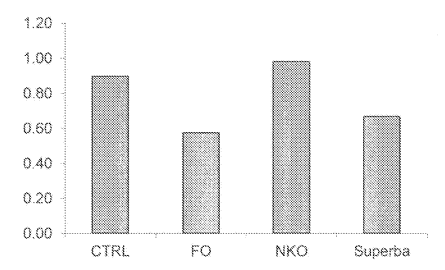


FIGURE 6

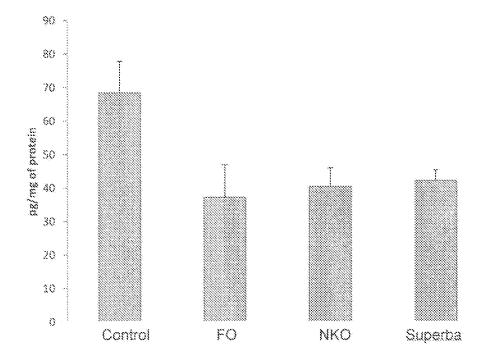


FIGURE 7

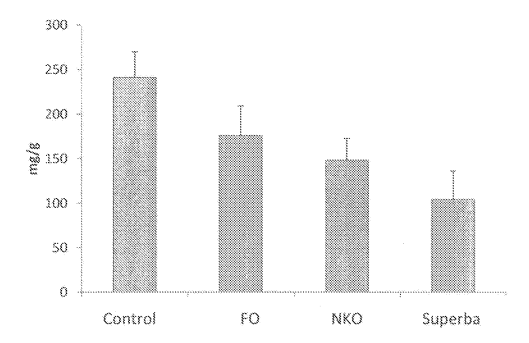
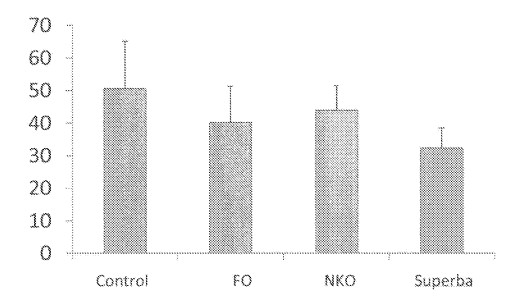
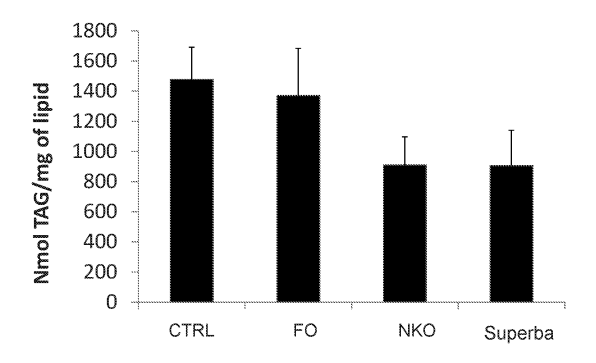
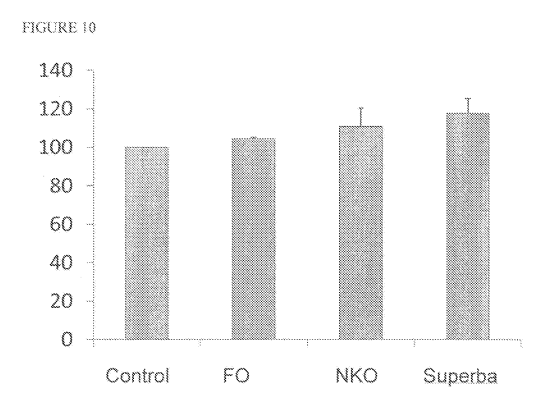


FIGURE 8



# FIGURE 9





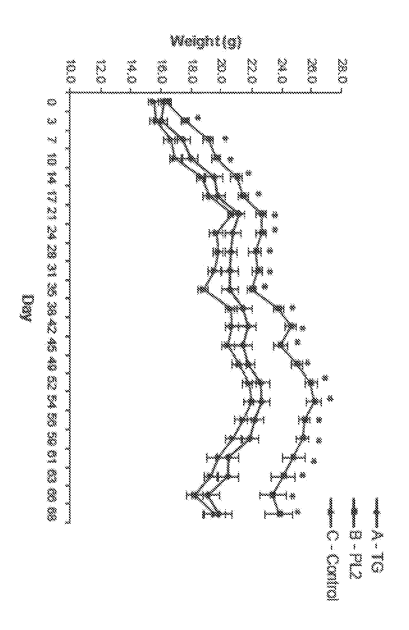


FIGURE 12

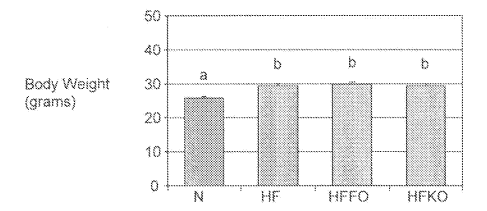


FIGURE 13

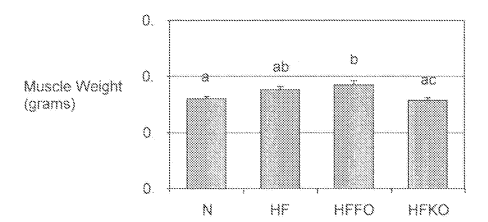


FIGURE 14

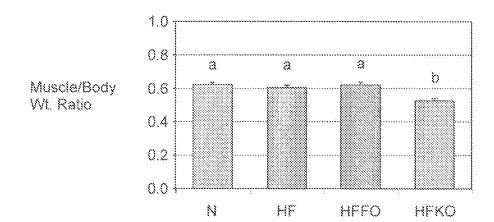


FIGURE 15

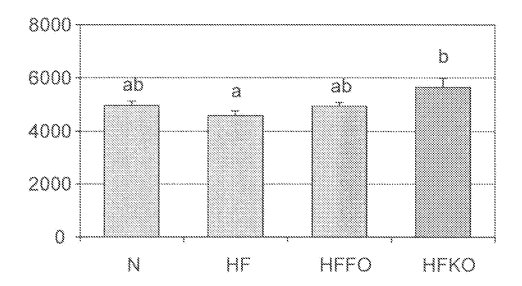


FIGURE 16

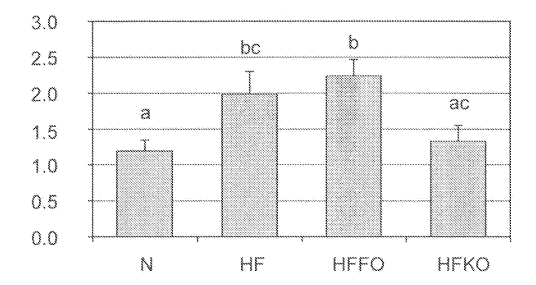


FIGURE 17

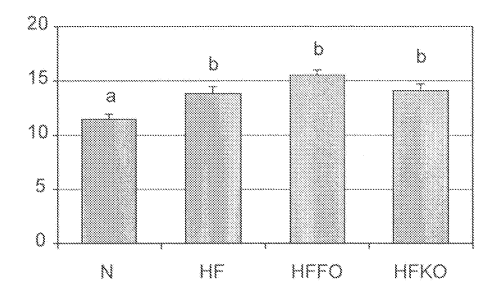


FIGURE 18

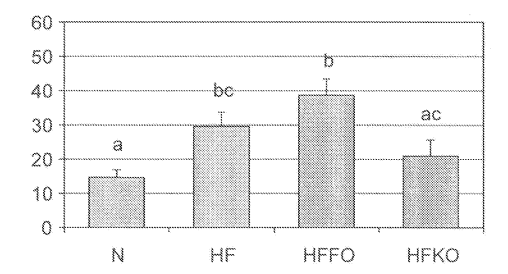
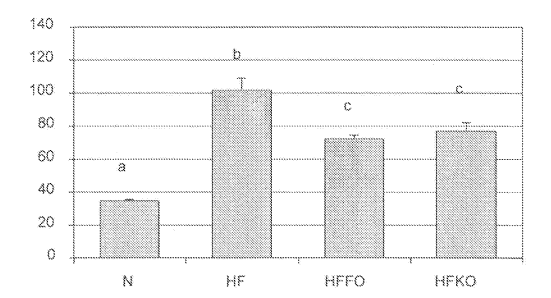


FIGURE 19



#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: Inge Bruheim et al. Group No.: TBD Serial No.: / , Examiner: TBD

Filed: Herewith

Entitled: BIOEFFECTIVE KRILL OIL COMPOSITIONS

#### 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).

Copies of any patent, publication, pending U.S. application or other information listed in the attached IDS Form PTO-SB08, not provided herewith, have been previously cited in parent U.S. patent application numbers 14/020,162 filed on September 6, 2013, and 12/057,775 filed on March 28, 2008. The documents can be found in the image file wrappers of the parent applications. In compliance with 37 C.F.R. § 1.98(d), Applicants have not included copies of these documents.

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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-12/CON.

Respectfully submitted,

Dated: June 13, 2016 /J. Mitchell Jones/

J. Mitchell Jones Registration No. 44,174

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	Filing Date		2016-06-13	
INFORMATION DISCLOSURE	First Named Inventor Inge Bi		Bruheim	
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit			
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	Attorney Docket Number		AKBM-14409/US-12/CON	

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Application Number		
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First Named Inventor Inge I		Bruheim
Art Unit		
Examiner Name		
Attorney Docket Number		AKBM-14409/US-12/CON

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Filing Date		2016-06-13
First Named Inventor Inge E		Bruheim
Art Unit		
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Application Number		
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First Named Inventor	Inge E	Bruheim
Art Unit		
Examiner Name		
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Art Unit		
Examiner Name		
Attorney Docket Number		AKBM-14409/US-12/CON

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43	Certificate of translation of Ex. 1074: Japanese Patent No. 60-153779, entitled "Nutritional Supplement"
44	Certificate of translation of Ex. 1076: Japanese Patent Publication No. H08-231391, entitled "Medicine for Improvement of Dementia Symptoms"
45	Certification of translation of Ex. 1070: Japanese Unexamined Patent Application Publication No. 02-215351

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	46	Certified translation of Ex. 1070: Japanese Unexamined Patent Application Publication No. 02-215351, titled Krill Phospholipids Fractioning Method ("Maruyama,"); Certificate of Translation provided as Ex. 1071.								
	47	Certified translation of Ex. 1072: Fisheries Agency, General Report on Research and Development of Techniques in Processing and Utilization of Marine Products, Chapter 6, Development of technology for recovery of valuable substances (astaxanthin) from krill, by Takao Fujita, pp. 273-307 (March 1985) ("Fujita"); Certificate of Translation provided as Ex. 1073.								
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Signature	/J. Mitchell Jones/	Date (YYYY-MM-DD)	2016-06-13
Name/Print	J. Mitchell Jones	Registration Number	44174

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	1	60-153779	JP			1985-08-13	Honen Seiyu Co. Lí al.	td. Et			
	2	H08-231391	JP			1996-08-09	Kanagawa Kagaku Kenkyuujo Co., Ltd.	. Et al.			

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	1	Declaration of Bjorn Ole Haugsgjerd submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("Haugsgjerd '348 Decl.")						
	2	Declaration of Dr. Albert Lee in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Lee"	")					
	3	Declaration of Dr. Albert Lee in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Lee'	")					
	4	Declaration of Dr. Chong Lee submitted during inter partes reexamination of parent patent U.S. ( Reexam Decl.")	8,030,348	("Yeboah				
	5	Declaration of Dr. Earl White submitted during prosecution of parent patent U.S. 8,030,348 ("20	11 White D	ecl.")				
	6	Declaration of Dr. Ivar Storrø in support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Sto	ггø")					
	7	Declaration of Dr. Ivar Storrø in support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Sto	ггø")					
	8	Declaration of Dr. Jacek Jaczynski from inter partes reexamination of the parent patent U.S. 8,0: Reexam. Decl.")	30,348 ("Ja	aczynski				
	9	Declaration of Dr. Jaczynski submitted during prosecution of parent patent U.S. 8,278,351 (Jacz	ynski '351	Decl.")				

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10	Declaration of Dr. Jeff Moore in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Moore")	
11	Declaration of Dr. Jeff Moore in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Moore")	
12	Declaration of Dr. Richard van Breemen in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Van Breemen")	
13	Declaration of Dr. Richard van Breemen in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Van Breemen")	
14	Declaration of Dr. Shahidi submitted during inter partes reexamination of parent patent U.S. 8,030,348 (Shahidi Reexam. Decl.")	
15	Declaration of Dr. Shahidi submitted during prosecution of parent patent U.S. 8,278,351 (Shahidi '351 Decl.")	
16	Declaration of Dr. Suzanne Budge in Support of Inter Partes Review of U.S. Pat. No. 8,278,351 ("Budge")	
17	Declaration of Dr. Suzanne Budge in Support of Inter Partes Review of U.S. Pat. No. 8,383,675 ("Budge")	
18	Declaration of Dr. Thomas Brenna in support of Inter Partes Review of U.S. Pat. No. 8,278,351	
19	Declaration of Dr. Thomas Brenna in support of Inter Partes Review of U.S. Pat. No. 8,383,675	
20	Declaration of Dr. Thomas Gundersen submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("Gundersen Decl.")	

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	21	Declaration of Dr. Tina Sampalis submitted during inter partes reexamination of parent patent U.S. 8,030,348 (Sampalis")	
	22	Declaration of Dr. Van Breemen submitted during Ex parte Reexamination of the '351 patent (Van Breemen '351 Reexam. Decl."	
	23	Declaration of Dr. Van Breemen submitted during Inter partes Reexamination of the '348 patent (Van Breemen '348 Reexam Decl."	
	24	Declaration of Dr. Yeboah submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("Yeboah Reexam Decl.")	
	25	Declaration of Dr. Yeboah submitted during prosecution of parent patent U.S. 8,278,351 ("Yeboah '351 Decl.")	
	26	Eichberg, "Lecithin – It Manufacture and Use in the Fat and Oil Industry," Oils and Soap 51-54, 1939 ("Eichberg")	
	27	Expert Witness Report of Dr. Theodore Welch submitted in relation to ITC Investigation No. 337-TA-877 ("Welch")	
	28	Farkas, Composition and Physical State of Phospholipids in Calanoid Copepods from India and Norway, LIPIDS, Vol. 23, No. 6 (1988)	
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	30		
	31	Folch, et al., A Simple Method for the Isolation and Purification of Total Lipids from Animal Tissues. J. Biol. Chem., 226, 497-509 (1957)	

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32	Grant of Request for Ex parte Reexamination of the '351 patent	
33	Grit et al., Hydrolysis of phosphatidylcholine in aqueous liposome dispersions, Int. J. Pharmaceutics 50:1-6 (1989)	
34	HENDERSON et al., Lipid Composition of the Pineal Organ from Rainbow Trout (Oncorhynchus mykiss), Lipids, Vol. 29, No. 5, pp. 311-317 (1994) ("Henderson ")	
35	HERMAN and GROVES, The Influence of Free Fatty Acid Formation on the pH of Phospholipid-Stabilized Triglyceride Emulsions, Pharmaceutical Research 10(5):774-776 (1993)	
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38	LE GRANDOIS et al., Investigation of Natural Phosphatidylholine Sources: Separation and Identification by Liquid Chromatography -Electrospray Ionization-Tandem Mass Spectrometry (LC-ESI-MS2) of Molecular Species, J. Agric. Food Chem., 57, 6014-20 (2009) ("Le Grandois")	
39	LIN et al., Effect of Dietary N-3 Fatty Acids Upon the PhospholipidMolecular Species of the Monkey Retina, Invest Ophthalmol Vis Sci. 1994;35:794-803	
40	MEDINA et al., C Nuclear Magnetic Resonance Monitoring of Free Fatty Acid Release After Fish Thermal Processing, J. Amer. Oil Chem. Soc. 71(5):479-82 (1994)	
41	October 24, 2012 Office Action, '675 patent	
42	Office Action dated January 5, 2012, '351 patent	

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4	3	Provisional Application No. 60/307,842 (Priority document for the '351 patent), available in PAIR					
4		Supplemental Declaration of Bjorn Ole Haugsgjerd submitted during inter partes reexamination of parent patent U.S. 3,030,348 ("Haugsgjerd '348 Supp. Decl.")					
4		Supplemental Declaration of Dr. Earl White submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("White Supp. Reexam. Decl.")					
4		Supplemental Declaration of Dr. Earl White submitted during prosecution of parent patent U.S. 8,278,351 ("White Supp. Decl.")					
4		Supplemental Declaration of Dr. Thomas Gundersen submitted during inter partes reexamination of parent patent U.S. 8,030,348 ("Gundersen Supp. Decl.")					
4	.8	Suzuki, T. and Shibata, N., "The utilization of Antarctic krill for human food," Food Rev. Int'l, 6:1, 119-147 (1990) ("Suzuki")					
4	.9	Takahashi et al., Compositional Changes in Molecular Species of Fish Muscle Phosphatidylcholine During Storage, Bull. Fac. Fish. Hokkaido Univ. 37(1), 80-84 1986.					
5	50	Takahashi et al., Molecular Species of Fish Muscle Lecithin, Bulletin of the Japanese Society of Scientific Fisheries 48 (12), 1803-1814 (1982)					
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Name/Print	J. Mitchell Jones	Registration Number	44174

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- 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.
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	Application Number			
	Filing Date		2016-06-13	
INFORMATION DISCLOSURE	First Named Inventor Inge Br		Bruheim	
STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)	Art Unit			
(Notion Submission under or of K 1.00)	Examiner Name			
	Attorney Docket Number	er	AKBM-14409/US-12/CON	

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	2	9072752		2015-07-07	AKER BIOMARINE ANTARCTIC AS			
	3	9034388		2015-05-19	Inge Bruheim et al.			
	4	9028877		2015-05-12	AKER BIOMARINE ANTARCTIC AS			
	5	9078905		2015-07-14	AKER BIOMARINE ANTARCTIC AS			
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	1	20140274968		2014-09-18	AKER BIOMARINE ANTARCTIC AS	
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	8	20100226977		2010-09-09	Snorre Tilseth et al.	
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First Named Inventor	Inge E	Bruheim
Art Unit		
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	11 20140		20140107072	2014-04-17		-17	Snorre Tilseth et al.					
	12 20090061067		2009-03-05		Snorre Tilseth et al.							
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	1		Takahashi et al., Prediction of Relative Retention Value of the Individual Molecular Species of Diacyl Glycerolipid on High Performance Liquid Chromatography, Bull. Fac. Fish. Hokkaido Univ. 38(4), 398-404. 1987									
	2	Tanaka, Biosynthesis of 1,2-dieicosapentaenoyl-sn-glycero-3-phosphocholine in Caenorhabditis elegans, Eur. J. Biochem. 263, 189±194 (1999)										
	3		her, Chapter 6, Glyce Mommsen (eds.)(19		nolipid m	etabolis	m, Biochemistry	and molecular biolog	gy of fis	hes, vol. 4,	Hochachka	

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4	Watanabe et al., Effective Components in Cuttlefish Meal and Raw Krill for Improvement of Quality of Red Seabream Pagrus major Eggs, Nippon Suisan Gakkaishi 57(4):681-694 (1991)("Watanabe")	
5	WHO News and Activities, Bulletin of the World Health Organization, 73(4), pp. 547-51 (1995) ("WHO Bulletin")	
6	VALERI, D., et al., "Visocities of Fatty acids, triglycerides and their binary mixtures," JAOCS 74 (1997) pp. 1221-1226	
7	CRC 2013-2014, 94th ed., pp. 6-231-6-235	
8	EP Opposition filed February 13, 2014 by Olympic Seafood AS, EP Patent Application No. EP08718910l6	
9	BRZUSTOWICZ, Michael R., et al., "Controlling Membrane Cholesterol Content. A Role for Polyunsaturated (Docosahexaenoate) Phospholipids," Biochemistry (2002), 41, pp. 12509-12519	
10	JONG-HO LEE, "A Review: Antioxygenic and Peroxide-decomposing Activities of Antarctic Krill Lipids," J. Korean Soc. Food Mutr. 13(3) pp. 326-333 (1984)	
11	KI WOONG CHO, et al., "Lipid and Fatty Acid Composition of the Antarctic Krill Euphausia superba," Ocean Research 21(2): 109-116 (1999)	
12	HVATTUM, Erlend, et al., "Effect of soybean oil and fish oil on individual molecular species of Atlantic salmon", Journal of Chromatography B, 748 (2000) 137-149	
13	IGARASHI, Daisuke, et al., "Positional Distribution of DHA and EPA in Phosphatidylcholine and Phosphatidylethanolamine from Different Tissues of Squids," J. Oleo Sci. Vol. 50, No. 9 (2001)	
14	TOCHIZAWA, Kaoru, et al., "Effects of Phospholipds Containing Docosahexaenoic Acid on Differentiation and Growth of HL-60 Human Promyelocytic Leukemia Cells," J. Jpn. Oil Chem. Soc. Vol. 46, No. 4 (1997)	

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		ZEROUGA, Mustapha, et al., "Comparison of phosphatidylcholines containing one or two docosahexaenoic acyl chains on properties of phospholipid monolayers and bilayers," Biochimica et Biophysica Acta 1236 (1995) 266-272								
	16	EUNG-HO LEE, et al., "Studies on the Processing of Krill Sauce," J. Korean Soc. Food Nutr. 13(1) 97-106 (1984)								
		HYUN-KU KIM, et al., "Effects of Cooking and Drying Methods on the Polar Lipds Composition of Shrimp," Korean J. Food Sci. Technol. Vol. 21, No. 1, pp. 25-30 (1989)								
		SHON, Mi-Yae, et al., "Effects of Krill and Cadmium on Lipid Composition of Plasma in Cholesterol-Fed Rats," J. Korean Soc. Food Nutr. 23(1), 38-43 (1994)								
	19	Summons Materials downloaded from ESPACE on December 16, 2014 for EP Patent Application No. 08 718 910.6								
		YANASE, M., "Innovations on the russian method for separating heat coagulated protein from antarctic krill, through autolysis," Bulletin of Tokai Regional Fisheries Research Laboratory, 1974, No. 78, p. 79-84								
		KOLAKOWSKI and GAJOWIECKI, "Optimization of autoproteolysis to obtain and edible product 'precipitate' from Antarctic krill," Seafood Science and Technology, pp. 331-336								
	22	EP Opposition filed May 8, 2015 by Olympic Seafood AS, EP Patent No. 2144618, 150 pages								
		ALLAHPICHAY et al., "Extraction of Growth Promoting Fractions from Non-muscle Krill Meal of Euphausia superba and its Effect on Fish Growth," Bulletin of the Japanese Society of Scientific Fisheries, 1984, 50(5): 821-826								
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Filing Date		2016-06-13		
First Named Inventor	Inge E	Bruheim		
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Examiner Name				
Attorney Docket Number		AKBM-14409/US-12/CON		

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Application Number			
Filing Date		2016-06-13	
First Named Inventor	Inge E	Bruheim	
Art Unit			
Examiner Name			
Attorney Docket Number		AKBM-14409/US-12/CON	

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

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Name/Print	J. Mitchell Jones	Registration Number	44174

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Electronic Patent Application Fee Transmittal				
Application Number:				
Filing Date:				
Title of Invention:	BIOEFFECTIVE KRILL O	IL COMPOSITIO	NS	
First Named Inventor/Applicant Name:	Inge Bruheim			
Filer:	John Mitchell Jones/M	allory Checkett		
Attorney Docket Number:	AKBM-14409/US-12/C	NC		
Filed as Large Entity				
Filing Fees for Track I Prioritized Examination - Nonpr	ovisional Applicatio	n under 35 U	SC 111(a)	
Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Basic Filing:				
Utility application filing	1011	1	280	280
Utility Search Fee	1111	1	600	600
Utility Examination Fee	1311	1	720	720
Request for Prioritized Examination	1817	1	4000	4000
Pages:				
Claims:				
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Petition:				
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Electronic Acknowledgement Receipt				
EFS ID:	26042498			
Application Number:	15180431			
International Application Number:				
Confirmation Number:	2763			
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS			
First Named Inventor/Applicant Name:	Inge Bruheim			
Customer Number:	72960			
Filer:	John Mitchell Jones/Mallory Checkett			
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Receipt Date:	13-JUN-2016			
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Application Type:	Utility under 35 USC 111(a)			

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Warnings:				'	
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6	Transmittal Letter	14409US12CON_IDS_Letter.pdf	88451	no	1
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to tre the a OR	ansact all busin attached transn reby appoint Pr	e Patent Practitioner(s) associated with the foliass in the United States Patent and Trademanittal letter (form PTO/AIA/82A) or identified a ractitioner(s) named in the attached list (form in the attached list (form in the attached list (form in the attached list).	ark Office connected the bove: 72960 PTO/AIA/82C) as my/ou	rewith for the application referenced in
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Inve	entor or Joint In	ventor (title not required below)		3
		ve of a Deceased or Legally Incapacitated Inv	ventor (title not required	below)
	•	to Whom the Inventor is Under an Obligation		
		wise Shows Sufficient Proprietary Interest (e.g ncurrently being filed with this document) (pro		
		SIGNATURE of Applic		
The under	signed (whose t	itie (& supplied balow) is authorized to act on bel	half of the applicant (e.g.,	where the applicant is a juristic entity).
Signature		(/////////////	Date (Options	<b>()</b>
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		orm must be signed by the applicant in accordar than one applicant, use multiple forms.	nce with 37 CFR 1.33. Se	e 37 CFR 1.4 for signature requirements
7 Total of	<del></del>	orne are submitted		

Total of 1 forms are submitted.

This collection of information is required by 37 CFR 1,131, 1,32, and 1,33. This information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to piccess) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1,13 and 1,14. This collection is estimated to to take 3 minutes to complete, including gathering, preparing, and submitting the completed application than to the USPTO. There will vary despending 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 Officer, U.S. Department of Generative, P.O. Sox 1458, Alexandria, VA 22313-1450, DO NOT SENO FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Sox 1458, Alexandria, VA 22313-1450.

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

Electronic Acknowledgement Receipt			
EFS ID:	26047101		
Application Number:	15180431		
International Application Number:			
Confirmation Number:	2763		
Title of Invention:	BIOEFFECTIVE KRILL OIL COMPOSITIONS		
First Named Inventor/Applicant Name:	Inge Bruheim		
Customer Number:	72960		
Filer:	John Mitchell Jones/Mallory Checkett		
Filer Authorized By:	John Mitchell Jones		
Attorney Docket Number:	AKBM-14409/US-12/CON		
Receipt Date:	13-JUN-2016		
Filing Date:			
Time Stamp:	15:39:50		
Application Type:	Utility under 35 USC 111(a)		
Payment information:	1		

Submitted with Payment	no
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## File Listing:

Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1	Power of Attorney	14409US12CON_POA_Exec.pdf	304245	no	1
·	Tower of Automey	,	b3e2124e58e3b129c1e2f600c0df553ae1ad 7f1a		·

## Warnings:

Information: RIMFROST EXHIBIT 1111 Page 0592 This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

### 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.