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## Krill Oil Capsules

### DEFINITION

#### Change to read:

▲Krill Oil Capsules contain NLT 95.0% of the labeled amount of Krill Oil calculated through the content of eicosapentaenoic acid (EPA), docosahexaenoic acid (DHA), and total phospholipids.▲ (USP 1-May-2019)

### IDENTIFICATION

#### Change to read:

#### • A. FATTY ACID PROFILE

**Antioxidant solution, System suitability solution 1, and Chromatographic system:** Proceed as directed in [Fats and Fixed Oils \(401\)](#), [Procedures, Omega-3 Fatty Acids Determination and Profile](#).

**Standard solution:** Prepare as directed in [Fats and Fixed Oils \(401\)](#), [Procedures, Omega-3 Fatty Acids Determination and Profile, Test solution 1](#), except use 250 mg of [USP Krill Oil RS](#).

**Sample solution:** Using the portion of oil from NLT 10 Capsules, prepare as directed in [Fats and Fixed Oils \(401\)](#), [Procedures, Omega-3 Fatty Acids Determination and Profile, Test solution 1](#).

#### System suitability

**Samples:** *System suitability solution 1* and *Standard solution*

#### Suitability requirements

**Chromatogram similarity:** The chromatogram from the *Standard solution* is similar to the reference chromatogram provided with the lot of [USP Krill Oil RS](#) being used.

**Resolution:** NLT 1.0 between the methyl oleate and methyl *cis*-vaccinate peaks, *Standard solution*

**Theoretical area percentages:** Meet the requirements for *System suitability solution 1*

#### Analysis

**Sample:** *Standard solution* and *Sample solution*

Identify the retention times of the peaks corresponding to the relevant fatty acid methyl esters by comparing the chromatogram of the *Standard solution* to the reference chromatogram provided with the lot of [USP Krill Oil RS](#) used.

Calculate the area percentage for each fatty acid as methyl esters in the portion of oil taken from the Capsules:

$$\text{Result} = (r_A/r_B) \times 100$$

$r_A$  = peak area of each individual fatty acid from the *Sample solution*

$r_B$  = total area of all peaks, except the solvent and butylated hydroxytoluene peaks, from the *Sample solution*

**Acceptance criteria:** See [Table 1](#).

Table 1

Fatty Acid	Short-Hand Notation	Lower Limit (Area %)	Upper Limit (Area %)
<b>Saturated fatty acids</b>			
Myristic acid	14:0	▲5.0▲ (USP 1-May-2019)	13.0
Palmitic acid	16:0	17.0	24.6
Palmitic acid:myristic acid ratio	16:0/14:0	1.6	▲3.6▲ (USP 1-May-2019)
<b>Monounsaturated fatty acids</b>			
Palmitoleic acid	16:1 n-7	2.5	▲12.0▲ (USP 1-May-2019)
<i>cis</i> -Vaccenic acid	18:1 n-7	4.7	8.0
Oleic acid	18:1 n-9	▲6.0▲ (USP 1-May-2019)	14.5
Eicosenic acid	20:1 n-9	0.0	2.0

Fatty Acid	Short-Hand Notation	Lower Limit (Area %)	Upper Limit (Area %)
<b>Polyunsaturated fatty acids</b>			
Linoleic acid	18:2 n-6	0.0	3.0
Eicosapentaenoic acid	20:5 n-3	14.0	▲28.0▲ (USP 1-May-2019)
Docosapentaenoic acid	22:5 n-3	0.0	0.7
Docosahexaenoic acid	22:6 n-3	7.1	15.7

**Change to read:**• **B. PHOSPHOLIPID PROFILE**

**Solution A**, ▲ (USP 1-May-2019) **Internal standard**, ▲ **Sample stock solution**, ▲ (USP 1-May-2019) **Sample solution**, **Standard solution**, **Instrumental conditions**, **System suitability**, and **Analysis**: Proceed as directed in the test for *Content of Total Phospholipids in Strength*.

**Acceptance criteria**: ▲The *Sample solution* shows <sup>31</sup>P nuclear magnetic resonance (NMR) spectra similar to those obtained with [USP Krill Oil RS](#). The main signal in the <sup>31</sup>P NMR is due to phosphatidylcholine (PC); a signal due to phosphatidylcholine ether has an intensity of about 10% of that due to PC; and the second signal in intensity is due to 2-lysophosphatidylcholine (2-LPC). Minor signals due to phosphatidylethanolamine (PE), *N*-acylphosphatidylethanolamine (NAPE), lysophosphatidylethanolamine (LPE), and 1-lysophosphatidylcholine (1-LPC) among others are also observed.▲ (USP 1-May-2019)

**STRENGTH****Delete the following:**▲• **CONTENT OF KRILL OIL**

**Analysis**: Weigh NLT 10 Capsules in a tared weighing bottle, and carefully open the Capsules, without loss of shell material. Transfer the combined Capsule contents to a 100-mL beaker. Remove any adhering substance from the emptied Capsules by washing with several small portions of 2,2,4-trimethylpentane. Discard the washings, and allow the empty Capsules to dry in a current of dry air until the 2,2,4-trimethylpentane is completely evaporated. Weigh the empty Capsules in the original tared weighing bottle, and calculate the average net weight per Capsule.

**Acceptance criteria**: 95.0%–105.0%▲ (USP 1-May-2019)

**Change to read:**• **CONTENT OF EPA AND DHA**

(See [Fats and Fixed Oils \(401\)](#), [Procedures, Omega-3 Fatty Acids Determination and Profile](#)).

**Standard solution 1a**, **Standard solution 1b**, **Standard solution 2a**, **Standard solution 2b**, **System suitability solution 1**, and **Chromatographic system**: Proceed as directed in the chapter.

**Test solution 1**: ▲Weigh NLT 10 Capsules in a tared weighing bottle, and carefully open the Capsules without loss of shell material. Transfer the combined contents to a 100-mL beaker. Remove any adhering substance from the emptied Capsules by washing with several small portions of [acetone](#). Discard the washings, and allow the empty Capsules to dry in a current of air until the [acetone](#) is completely evaporated. Weigh the empty Capsules in the original tared weighing bottle, and calculate the average fill weight per Capsule. Take 250 mg of the combined Capsule contents and▲ (USP 1-May-2019) proceed as directed in [Fats and Fixed Oils \(401\)](#), [Procedures, Omega-3 Fatty Acids Determination and Profile, Test solution 1](#).

**Test solution 2**: ▲▲ (USP 1-May-2019) Prepare as directed in [Fats and Fixed Oils \(401\)](#), [Procedures, Omega-3 Fatty Acids Determination and Profile, Test solution 2](#)▲ using 250 mg of the combined Capsule contents.▲ (USP 1-May-2019)

**Analysis**: Proceed as directed in [Fats and Fixed Oils \(401\)](#), [Procedures, Omega-3 Fatty Acids Determination and Profile, Analysis \(for triglycerides\)](#).

▲Calculate the percentage of the labeled amounts of EPA and DHA in the portion of Capsules taken:

$$\text{Result} = A \times W_{Av} \times 100/L$$

A = content of EPA or DHA in the portion of Capsules taken (mg/mg)

$W_{Av}$  = average fill weight (mg/Capsule)

L = label claim of EPA or DHA (mg/Capsule)▲ (USP 1-May-2019)

**Acceptance criteria**: ▲NLT 95.0% of the labeled amounts of EPA and DHA▲ (USP 1-May-2019)

**Change to read:**• **CONTENT OF TOTAL PHOSPHOLIPIDS**

(See [Nuclear Magnetic Resonance Spectroscopy \(761\)](#), [Qualitative and Quantitative NMR Analysis](#).)

[NOTE—All deuterated solvents used in this method should be NLT 99.8 atom % D. ▲▲ (USP 1-May-2019)]

**Solution A**: 0.2 M ethylenediaminetetraacetic acid (EDTA) adjusted with a 1 M cesium carbonate solution to a pH of 7.2–7.5. ▲▲ (USP 1-May-2019) [NOTE—Use cesium carbonate of a sufficient grade for trace metals analysis.]

▲▲ (USP 1-May-2019)

**Internal standard**: Use a triphenyl phosphate nuclear magnetic resonance (NMR) reference standard with NLT 99% purity.

▲**Sample stock solution**: Transfer a number of Capsules (NLT 5) to a conical flask with stopper. Add 2 mL/Capsule of [water](#) and melt the gelatin shells at 50°–60°. Add 10 mL/Capsule of [chloroform](#) and [methanol \(2:1\)](#), insert the stopper and shake intensively for 20 min. Transfer to a separation funnel fitted with glass wool

**Sample solution:** Transfer an aliquot of the *Sample stock solution* equivalent to 300 mg of Krill Oil to a suitable sealable glass vial, and add about 25 mg of the *Internal standard* accurately weighed. Evaporate to dryness. Add 2 mL of deuterated chloroform (chloroform-d) containing 0.05% tetramethylsilane (TMS), 2 mL of deuterated methanol (methanol-d4), and 2 mL of *Solution A*. Seal the vial and vortex intensively for 30–60 s, shake for an additional 30 min on a shaking device, and centrifuge the contents of the vial. Pass the entire amount of the lower organic phase through a glass fiber filter, and collect the filtrate in the appropriate NMR tube.▲ (USP 1-May-2019)

**Standard solution:**▲ Transfer 300 mg of [USP Krill Oil RS](#) to a suitable sealable glass vial, and add about 25 mg of the *Internal standard* accurately weighed. Add 2 mL of methanol-d4, 2 mL of chloroform-d containing 0.05% TMS, and 2 mL of *Solution A*. Seal the vial and vortex intensively for 30–60 s, shake for an additional 30 min on a shaking device, and centrifuge the contents of the vial. Pass the entire amount of the lower organic phase through a glass fiber filter, and collect the filtrate in the appropriate NMR tube.▲ (USP 1-May-2019)

#### Instrumental conditions

**Magnetic field strength:**▲ NLT 7.05 Tesla (resonance frequencies of 121 MHz for  $^{31}\text{P}$  or 300 MHz for  $^1\text{H}$ )▲ (USP 1-May-2019)

**Probe:** Direct observe probe capable of tuning to the resonance frequency of  $^{31}\text{P}$  (dependent on the specific magnetic field strength used)▲ at a temperature of 35°▲ (USP 1-May-2019)

**Data collection:** Use the parameters specified in [Table 2](#). Use 90° pulses, and calibrate pulses before use according to the recommendations supplied by the instrument manufacturer.

Table 2

Parameter	$^{31}\text{P}$ NMR Quantitative Measurement	▲▲ (USP 1-May-2019)
Pulse program	$^1\text{H}$ -decoupled $^{31}\text{P}$ (inverse gated)	▲▲ (USP 1-May-2019)
Spectral width	50 ppm (25 to –25 ppm)	▲▲ (USP 1-May-2019)
Transmitter offset	Center of spectral width, 0 ppm	▲▲ (USP 1-May-2019)
Relaxation delay	5–15 s	▲▲ (USP 1-May-2019)
Acquisition time	1–6 s	▲▲ (USP 1-May-2019)
Size of data set	NLT 64k (32k with zero-filling)	▲▲ (USP 1-May-2019)

[NOTE—The acquisition time is dependent upon the dwell time and the number of data points collected. The number of scans acquired using a▲7.05 Tesla magnet▲ (USP 1-May-2019) must be NLT 512.]

**System suitability:**▲ The *Standard solution* shows the  $^{31}\text{P}$  NMR signal for triphenyl phosphate at –17.80 ppm, and the signal for phosphatidylcholine at –0.89 ppm. The signal-to-noise ratio for the phosphatidylcholine signal in the  $^{31}\text{P}$  spectrum of the *Sample solution* obtained in the *Analysis* is NLT 2000. Using the baseline as a reference, determine the height of the phosphatidylcholine ether peak and draw a line parallel to the baseline at 30% of that total peak height (intensity). The valley between the peaks of phosphatidylcholine ether and phosphatidylcholine is below the line drawn.▲ (USP 1-May-2019)

**Analysis:**▲ Obtain the quantitative  $^{31}\text{P}$  spectrum of the *Sample solution* and the *Standard solution* as directed in *Data collection*. Record the resulting spectra, and integrate the complete set of phospholipid peaks as identified by a comparison with the reference spectrum provided with [USP Krill Oil RS](#). The integration region for each signal must extend  $\pm 0.05$  ppm on either side of the  $^{31}\text{P}$  signal.▲ (USP 1-May-2019)

**Calculations:** Use the following equations and molecular weights listed in [Table 3](#) to▲ calculate the content of the phospholipid of interest in the Capsules taken:▲

(USP 1-May-2019)

$$mmol_{IS} = (W_{IS} \times C_{IS}) / (MW_{IS} \times 100)$$

$mmol_{IS}$  = millimoles of the *Internal standard* in the *Sample solution* (mmol)

$W_{IS}$  = weight of the *Internal standard* added to the *Sample solution* (mg)

$C_{IS}$  = purity value of the *Internal standard*, based on quantitative  $^{31}\text{P}$  NMR analysis (% by weight)

$MW_{IS}$  = molecular weight of the *Internal standard*, 326.28 g/mol (for triphenyl phosphate)

$$mmol_{PL} = (I_{PL} \times A_{IS} \times mmol_{IS}) / (I_{IS} \times A_{PL})$$

$mmol_{PL}$  = millimoles of the phospholipid of interest in the *Sample solution* (mmol)

$I_{PL}$  = integrated area under the phospholipid signal of interest obtained from the spectrum of the *Sample solution*

$A_{IS}$  = number of phosphorus atoms per molecule expected from the *Internal standard*, 1 (for triphenyl phosphate)

$I_{IS}$  = integrated area under the *Internal standard* obtained from the spectrum of the *Sample solution*

$A_{PL}$  = number of phosphorus atoms per molecule expected from the phospholipid of interest, 1 (for any phospholipid listed in [Table 3](#))

$$\Delta C_{PL} = 1/n \times MW_{PL} \times mmol_{PL} \times V/a$$

$C_{PL}$  = content of the phospholipid of interest in the Capsules taken (mg/Capsule)

$n$  = number of Capsules used to prepare the *Sample stock solution*

$MW_{PL}$  = molecular weight of the phospholipid of interest (mg/mmol, from [Table 3](#))

$mmol_{PL}$  = millimoles of the phospholipid of interest in the *Sample solution* (mmol)

$V$  = volume of volumetric flask used to prepare the *Sample stock solution* (mL)

$a$  = volume of the aliquot of *Sample stock solution* used to prepare the *Sample solution* (mL)  $\blacktriangle$  (USP 1-May-2019)

**Table 3**

Component	Approximate Chemical Shift (ppm) in Reference to Triphenyl Phosphate	Molecular Weight (g/mol)
Triphenyl phosphate ( <i>Internal standard</i> )	-17.8	—
Phosphatidylcholine, including ether (PC)	-0.89	791
1-Lysophosphatidylcholine (1-LPC) <sup>a</sup>	-0.48	534.5
2-Lysophosphatidylcholine (2-LPC) <sup>a</sup>	-0.4	534.5
Phosphatidylethanolamine (PE)	-0.24	770
<i>N</i> -Acylphosphatidylethanolamine (NAPE)	0	1032
Lysophosphatidylethanolamine (LPE)	0.25	492.5
Other	—	800

<sup>a</sup> Ability to resolve the signals of 1-LPC and 2-LPC will depend upon the applied magnetic field strength of the NMR spectrometer used for the test procedure.

$\blacktriangle$  Calculate the percentage of the labeled amount of total phospholipids per Capsule taken:

$$\text{Result} = \Sigma C_{PL} \times 100/L$$

$C_{PL}$  = sum of the individual amounts of phospholipids of interest in the Capsules taken (mg/Capsule)

$L$  = label claim of total phospholipids (mg/Capsule)  $\blacktriangle$  (USP 1-May-2019)

**Acceptance criteria:**  $\blacktriangle$ NLT 95.0% of the labeled amount of total phospholipids  $\blacktriangle$  (USP 1-May-2019)

**Change to read:**

• **CONTENT OF ASTAXANTHIN**

[NOTE—Perform this analysis in subdued light using low-actinic glassware.]

**Sample solution:** 0.005 g/mL of Krill Oil in chloroform using the portion of oil from NLT 10 Capsules. [NOTE—If the solution is not clear, centrifuge it with an appropriate centrifuge to obtain a clear supernatant.]

**Instrumental conditions**

(See [Ultraviolet-Visible Spectroscopy \(857\)](#).)

**Analytical wavelength:** 486 nm

**Cell:** 1 cm

**Blank:** Chloroform

**Analysis**

**Sample:** *Sample solution*

Calculate the percentage of astaxanthin in the portion of Krill Oil taken from the Capsules:

$$\text{Result} = A/(F \times C)$$

$A$  = absorbance of the *Sample solution*

$F$  = coefficient of extinction ( $E^{1\%}$ ) of pure astaxanthin in chloroform ( $100 \text{ mL} \cdot \text{g}^{-1} \cdot \text{cm}^{-1}$ ), 1692

**Acceptance criteria:** NLT  $\blacktriangle$ 0.005%  $\blacktriangle$  (USP 1-May-2019)

#### PERFORMANCE TESTS

- [DISINTEGRATION AND DISSOLUTION \(2040\)](#), [Rupture Test for Soft Shell Capsules](#): Meet the requirements
- [WEIGHT VARIATION \(2091\)](#): Meet the requirements

#### CONTAMINANTS

##### LIMIT OF DIOXINS, FURANS, AND POLYCHLORINATED BIPHENYLS

**Analysis:** Determine the content of polychlorinated dibenzo-para-dioxins (PCDDs) and polychlorinated dibenzofurans (PCDFs) by Method 1613, Revision B of the Environmental Protection Agency (EPA). Determine the content of polychlorinated biphenyls (PCBs) by Method 1668, Revision A of the EPA.

**Acceptance criteria:** The sum of PCDDs and PCDFs is NMT 2.0 pg/g of World Health Organization (WHO) toxic equivalents. The sum of PCDDs, PCDFs, and dioxin-like PCBs [non-ortho International Union of Pure and Applied Chemistry (IUPAC) congeners PCB-77, PCB-81, PCB-126, and PCB-169; and mono-ortho IUPAC congeners PCB-105, PCB-114, PCB-118, PCB-123, PCB-156, PCB-157, PCB-167, and PCB-189] is NMT 10.0 pg/g of WHO toxic equivalents.

- [MICROBIAL ENUMERATION TESTS \(2021\)](#): The total aerobic microbial count does not exceed  $10^3$  cfu/g, and the combined molds and yeasts count does not exceed  $10^2$  cfu/g.
- [ABSENCE OF SPECIFIED MICROORGANISMS \(2022\)](#), [Test Procedures](#), [Test for Absence of Salmonella Species](#) and [Test for Absence of Escherichia coli](#): Meet the requirements

#### SPECIFIC TESTS

##### ASTAXANTHIN ESTERIFICATION

**Standard solution A:** 10 mg/mL of [USP Astaxanthin Esters from Haematococcus pluvialis RS](#) in acetone

**Standard solution B:** 10 mg/mL of [USP Astaxanthin \(Synthetic\) RS](#) in acetone

**Sample solution:** Using the portion of oil from NLT 10 Capsules, prepare a solution of 250 mg/mL in acetone.

##### Chromatographic system

(See [Chromatography \(621\)](#), [General Procedures](#), [Thin-Layer Chromatography](#).)

**Mode:** TLC

**Adsorbent:** 0.25-mm layer of chromatographic silica gel. [NOTE—Dry silica gel at 110° for 1 h before use.]

**Application volume:** 5  $\mu$ L

**Developing solvent system:** Hexane and acetone (70:30)

##### Analysis

**Samples:** *Standard solution A*, *Standard solution B*, and *Sample solution*

Develop the chromatogram in the *Developing solvent system* until the solvent front has moved about 15 cm of the length of the plate. Remove the plate from the chamber, and allow to dry.

**Acceptance criteria:** The principal spot from *Standard solution B*, located in the bottom half of the plate, is free astaxanthin. The *Sample solution* may exhibit a light, minor spot in the same location. The principal spots from *Standard solution A* are from monoesters (primary spot, located slightly above the middle of the plate) and diesters (secondary spot, located in the top third of the plate). The principal spot from the *Sample solution* should correspond in color and  $R_f$  value to the diester spot from *Standard solution A*. The secondary spot from the *Sample solution* should correspond in color and approximately the same  $R_f$  value to the monoester spot from *Standard solution A*. [NOTE—Slight differences in  $R_f$  values within monoester spots and within diester spots may exist because of different intensities.]

- [FATS AND FIXED OILS \(401\)](#), [Procedures](#), [Peroxide Value](#): NMT 5.0 mEq peroxide/kg

#### ADDITIONAL REQUIREMENTS

- **PACKAGING AND STORAGE:** Preserve in tight containers, and store at room temperature. Protect from light.

##### Change to read:

- **LABELING:** The label states the amount of docosahexaenoic acid (DHA), eicosapentaenoic acid (EPA), and total phospholipids  $\blacktriangle$  in mg/Capsule.  $\blacktriangle$  (USP 1-May-2019)

##### Change to read:

- [USP REFERENCE STANDARDS \(11\)](#)  
[USP Astaxanthin \(Synthetic\) RS](#)  
[USP Astaxanthin Esters from Haematococcus pluvialis RS](#)

$\blacktriangle$  (USP 1-May-2019)

[USP Krill Oil RS](#)

$\blacktriangle$  (USP 1-May-2019)

**Auxiliary Information-** Please [check for your question in the FAQs](#) before contacting USP.

Topic/Question	Contact	Expert Committee
KRILL OIL CAPSULES	<a href="#">Natalia Davydova</a> Scientific Liaison +1 (301) 816-8328	NBDS2015 Non-botanical Dietary Supplements 2015

**Chromatographic Columns Information:** [Chromatographic Columns](#)

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