Title: SMB PROCESS FOR PRODUCING HIGHLY PURE EPA FROM FISH OIL

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Figure 2

2A Feed mixture 1st intermediate product 2nd intermediate product

1st intermediate product 2nd intermediate product PUFA product

FIRST SEPARATION STEP SECOND SEPARATION STEP THIRD SEPARATION STEP

(57) Abstract: The present invention provides a chromatographic separation process for recovering a polyunsaturated fatty acid (PUFA) product from a feed mixture which is a fish oil or which is derived from fish oil, which process comprises the steps of: (i) purifying the feed mixture in a chromatographic separation step, to obtain a first intermediate product; and (ii) purifying the first intermediate product obtained in (i) in a simulated or actual moving bed chromatographic separation step, to obtain a second intermediate product; and (iii) purifying the second intermediate product obtained in (ii) in a simulated or actual moving bed chromatographic separation step, to obtain the PUFA product; wherein an aqueous organic solvent is used as eluent in each separation step; saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step; the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); and the PUFA product obtained in the third separation step contains EPA or an EPA derivative in an amount greater than 90 wt%.
The present invention relates to an improved chromatographic separation process for purifying the polyunsaturated fatty acid EPA or a derivative thereof.

EPA and its derivatives are precursors for biologically important molecules, which play an important role in the regulation of biological functions such as platelet aggregation, inflammation and immunological responses. Thus, EPA and its derivatives may be therapeutically useful in treating a wide range of pathological conditions including CNS conditions; neuropathies, including diabetic neuropathy; cardiovascular diseases; general immune system and inflammatory conditions, including inflammatory skin diseases.

EPA is found in natural raw materials, and in particular fish oils. The EPA in fish oils is, however, present in such oils in admixture with saturated fatty acids and numerous other impurities.

Purification of EPA from fish oils is particularly challenging. Thus, fish oils are extremely complex mixtures containing a large number of different components with very similar retention times in chromatography apparatuses. They represent a much more challenging feedstock from which to purify EPA than, for example, an algal oil feedstock. However, a very high degree of purity of EPA is required, particularly for pharmaceutical and nutraceutical applications. Historically, therefore, distillation has been used to purify EPA for therapeutic applications.

Unfortunately, EPA is extremely fragile. Thus, when heated in the presence of oxygen, it is prone to isomerization, peroxidation and oligomerization. The fractionation and purification of EPA to prepare pure fatty acids is therefore difficult. Distillation, even under vacuum, can lead to non-acceptable product degradation.

Simulated and actual moving bed chromatography are known techniques, familiar to those of skill in the art. The principle of operation involves countercurrent movement of a liquid eluent phase and a solid adsorbent phase. This operation allows minimal usage of solvent making the process economically viable. Such separation technology
has found several applications in diverse areas, including hydrocarbons, industrial chemicals, oils, sugars and APIs.

As is well known, in a conventional stationary bed chromatographic system, a mixture whose components are to be separated percolates through a container. The container is generally cylindrical, and is typically referred to as the column. The column contains a packing of a porous material (generally called the stationary phase) exhibiting a high permeability to fluids. The percolation velocity of each component of the mixture depends on the physical properties of that component so that the components exit from the column successively and selectively. Thus, some of the components tend to fix strongly to the stationary phase and thus will percolate slowly, whereas others tend to fix weakly and exit from the column more quickly. Many different stationary bed chromatographic systems have been proposed and are used for both analytical and industrial production purposes.

In contrast, a simulated moving bed chromatography apparatus consists of a number of individual columns containing adsorbent which are connected together in series. Eluent is passed through the columns in a first direction. The injection points of the feedstock and the eluent, and the separated component collection points in the system, are periodically shifted by means of a series of valves. The overall effect is to simulate the operation of a single column containing a moving bed of the solid adsorbent, the solid adsorbent moving in a countercurrent direction to the flow of eluent. Thus, a simulated moving bed system consists of columns which, as in a conventional stationary bed system, contain stationary beds of solid adsorbent through which eluent is passed, but in a simulated moving bed system the operation is such as to simulate a continuous countercurrent moving bed.

An actual moving bed system is similar in operation to a simulated moving bed system. However, rather than shifting the injection points of the feed mixture and the eluent, and the separated component collection points by means of a system of valves, instead a series of adsorption units (i.e. columns) are physically moved relative to the feed and drawoff points. Again, operation is such as to simulate a continuous countercurrent moving bed.

Processes and equipment for actual moving bed chromatography are described in several patents, including US 6,979,402, US 5,069,883 and US 4,764,276, the entirety of which are incorporated herein by reference.

A typical simulated moving bed chromatography apparatus is illustrated with reference to Figure 1. The concept of a simulated or actual moving bed chromatographic separation process is explained by considering a vertical chromatographic column containing stationary phase S divided into sections, more precisely into four superimposed sub-zones I, II, III and IV going from the bottom to the top of the column. The eluent is introduced at the bottom at IE by means of a pump P. The mixture of the components A and B which are to be separated is introduced at IA + B between sub-zone II and sub-zone III. An extract containing mainly B is collected at SB between sub-zone I and sub-zone II, and a raffinate containing mainly A is collected at SA between sub-zone III and sub-zone IV.

In the case of a simulated moving bed system, a simulated downward movement of the stationary phase S is caused by movement of the introduction and collection points relative to the solid phase. In the case of an actual moving bed system, simulated downward movement of the stationary phase S is caused by movement of the various chromatographic columns relative to the introduction and collection points. In Figure 1, eluent flows upward and mixture A + B is injected between sub-zone II and sub-zone III. The components will move according to their chromatographic interactions with the stationary phase, for example adsorption on a porous medium. The component B that exhibits stronger affinity to the stationary phase (the slower running component) will be more slowly entrained by the eluent and will follow it with delay. The component A that exhibits the weaker affinity to the stationary phase (the faster
running component) will be easily entrained by the eluent. If the right set of parameters, especially the flow rate in each sub-zone, are correctly estimated and controlled, the component A exhibiting the weaker affinity to the stationary phase will be collected between sub-zone III and sub-zone IV as a raffinate and the component B exhibiting the stronger affinity to the stationary phase will be collected between sub-zone I and sub-zone II as an extract.

To achieve high purity EPA or EPA ethyl ester in purities of greater than 90%, for example greater than 95 or 97%, it is possible to utilise a simulated moving bed separation process which performs two simultaneous separation steps. Such a process is described in international patent application no. PCT/GB 10/002339, the entirety of which is incorporated herein by reference.

In general, all chromatographic separation techniques for separating PUFAs, including SMB processes, utilise large volumes of organic solvents as eluents. After the chromatographic separation process is completed the PUFAs must be recovered from solution in the eluent. Typically a large expenditure of time and energy is involved in recovering PUFAs from solution in the eluent. Furthermore, organic solvents used as eluents in chromatographic separation processes are frequently harmful to the environment or to the operatives handling them. Therefore, a chromatographic separation process which reduces the amount of organic solvent that needs to be used is required.

It has now been advantageously found that EPA or an EPA derivative can be produced in a similarly high purity as described in PCT/GB 10/002339 by a three-step separation process which uses a much lower volume of solvent that the two-step process. The improved process of the present invention utilises almost 50% less solvent than the two-step process described in PCT/GB 10/002339. This is clearly advantageous in terms of cost, ease of recovery of product, and environmental impact.

Summary of the invention

It has been surprisingly found that EPA or an EPA derivative can be effectively purified from commercially available feedstocks such as fish oils by simulated or
actual moving bed apparatus using a relatively low volume of an aqueous organic solvent eluent. The present invention therefore provides a chromatographic separation process for recovering a polyunsaturated fatty acid (PUFA) product from a feed mixture which is a fish oil or which is derived from fish oil, which process comprises the steps of:

(i) purifying the feed mixture in a chromatographic separation step, to obtain a first intermediate product; and

(ii) purifying the first intermediate product obtained in (i) in a simulated or actual moving bed chromatographic separation step, to obtain a second intermediate product; and

(iii) purifying the second intermediate product obtained in (ii) in a simulated or actual moving bed chromatographic separation step, to obtain the PUFA product; wherein an aqueous organic solvent is used as eluent in each separation step; saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step;

the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); and

the PUFA product obtained in the third separation step contains EPA or an EPA derivative in an amount greater than 90 wt%.

Also provided is a PUFA product obtainable by the process of the present invention.

Description of the Figures

Figure 1 illustrates the basic principles of a simulated or actual moving bed process for separating a binary mixture.

Figure 2 illustrates three ways in which the chromatographic separation process of the invention may be carried out.

Figure 3 illustrates a preferred embodiment of the invention which is suitable for producing high purity EPA.

Figure 4 illustrates in more detail the embodiment of Figure 2.
Figure 5 illustrates a more preferred embodiment of the embodiment shown in Figure 2.

Figure 6 illustrates a two-stage separation process for producing EPA (not in accordance with the present invention).

Figure 7 shows a GC trace of a suitable feed stock for use in accordance with the process of the present invention.

Figure 8 shows a GC trace of a first intermediate product produced in accordance with the process of the present invention.

Figure 9 shows a GC trace of a second intermediate product produced in accordance with the process of the present invention.

Figure 10 shows a GC trace of a PUFA product produced in accordance with the process of the present invention.

**Detailed description of the invention**

As used herein, the term "PUFA product" refers to a product comprising one or more polyunsaturated fatty acids (PUFAs), and/or derivatives thereof, typically of nutritional or pharmaceutical significance. The PUFA product obtained in the process of the present invention contains EPA or an EPA derivative in an amount greater than 90 wt%, i.e. EPA or an EPA derivative is present at 90 wt% purity relative to all of the components in the final PUFA product not including the aqueous organic solvent eluent. Thus EPA or an EPA derivative is present in the PUFA product in an amount of at least 90 wt% on the basis of all of the components of the PUFA product which originated in the feed mixture.

An EPA derivative is EPA in the form of a mono-, di- or tri-glyceride, ester, phospholipid, amide, lactone, or salt. Triglycerides and esters are preferred. Esters are more preferred. Esters are typically alkyl esters, preferably C1-C6 alkyl esters,
more preferably C1-C4 alkyl esters. Examples of esters include methyl and ethyl esters. Ethyl esters are most preferred.

Typically, the PUFA product contains EPA or an EPA derivative in an amount greater than 95 wt%, preferably greater than 97 wt%.

In one embodiment, the PUFA product contains EPA in an amount greater than 90 wt%, preferably greater than 95 wt%, more preferably greater than 97 wt%. As explained above, EPA is present at the specified wt% relative to the total amount of all of the components of the PUFA product which originated in the feed mixture.

In another embodiment, the PUFA product contains EPA ethyl ester in an amount greater than 90 wt%, preferably greater than 95 wt%, more preferably greater than 97 wt%. As explained above, EPA is present at the specified wt% relative to the total amount of all of the components of the PUFA product which originated in the feed mixture.

Suitable feed mixtures for fractionating by the process of the present invention are fish oils, or feed stocks derived from fish oils. Suitable fish oils for use in the process of the present invention are well known to the skilled person. Typical fish oils contain EPA, DHA, SDA, and typically a range of other PUFAs both more and less polar than EPA, saturated fatty acids and monounsaturated fatty acids.

The feed mixture may undergo chemical treatment before fractionation by the process of the invention. For example, it may undergo glyceride transesterification or glyceride hydrolysis followed in certain cases by selective processes such as crystallisation, molecular distillation, urea fractionation, extraction with silver nitrate or other metal salt solutions, iodolactonisation or supercritical fluid fractionation. Alternatively, a feed mixture may be used directly with no initial treatment step.

The feed mixtures typically contain the PUFA product and at least one more polar component and at least one less polar component. The less polar components have a stronger adherence to the adsorbent used in the process of the present invention than does the PUFA product. During operation, such less polar components typically
move with the solid adsorbent phase in preference to the liquid eluent phase. The more polar components have a weaker adherence to the adsorbent used in the process of the present invention than does the PUFA product. During operation, such more polar components typically move with the liquid eluent phase in preference to the solid adsorbent phase. In general, more polar components will be separated into a raffinate stream, and less polar components will be separated into an extract stream.

Examples of the more and less polar components include (1) other compounds occurring in natural oils (e.g. marine oils), (2) byproducts formed during storage, refining and previous concentration steps and (3) contaminants from solvents or reagents which are utilized during previous concentration or purification steps.

Examples of (1) include other unwanted PUFAs; saturated fatty acids; sterols, for example cholesterol; vitamins; and environmental pollutants, such as polychlorobiphenyl (PCB), polyaromatic hydrocarbon (PAH) pesticides, chlorinated pesticides, dioxines and heavy metals. PCB, PAH, dioxines and chlorinated pesticides are all highly non-polar components.

Examples of (2) include isomers and oxidation or decomposition products from the PUFA product, for instance, auto-oxidation polymeric products of fatty acids or their derivatives.

Examples of (3) include urea which may be added to remove saturated or mono-unsaturated fatty acids from the feed mixture.

Preferably, the feed mixture is a PUFA-containing marine oil (e.g. a fish oil), more preferably a marine oil (e.g. a fish oil) comprising EPA and/or DHA.

A typical feed mixture for preparing concentrated EPA (EE) by the process of the present invention comprises 50-75% EPA (EE), 0 to 10% DHA (EE), and other components including other essential ω-3 and ω-6 fatty acids.
A preferred feed mixture for preparing concentrated EPA (EE) by the process of the present invention comprises 55% EPA (EE), 5% DHA (EE), and other components including other essential ω-3 and ω-6 fatty acids. DHA (EE) is less polar than EPA(EE).

The process of the present invention involves multiple chromatography separation steps.

The first separation step is effective to remove saturated and/or monounsaturated fatty acids present in the feed mixture and may be carried out using a stationary bed or simulated or actual moving bed chromatography apparatus.

When the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, there are several ways in which the three separation steps may be realised. Four preferred ways of carrying out the process are given as first, second, third and fourth embodiments below.

In a first embodiment, the first, second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the first, second and third separation steps being carried out in first, second and third zones respectively, wherein each zone has one or more injection points for a feed mixture stream, one or more injection points for water and/or organic solvent, a raffinate take-off stream from which liquid can be collected from said zone, and an extract take-off stream from which liquid can be collected from said zone.

Typically, each zone has only one injection point for a feed mixture. In one embodiment, each zone has only one injection point for the aqueous organic solvent eluent. In another embodiment, each zone has two or more injection points for water and/or organic solvent.
Typically each zone used has a single array of chromatography columns linked in series containing, as eluent, an aqueous organic solvent. Typically, each of the chromatography columns in a zone are linked to the two columns in the apparatus adjacent to that column. Thus, the output from a given column in a zone is connected to the input of the adjacent column, for example in the zone, which is downstream with respect to the flow of eluent in the system. Typically, none of the chromatography columns in a zone are linked to non-adjacent columns in the same zone.

The term "raffinate" is well known to the person skilled in the art. In the context of actual and simulated moving bed chromatography it refers to the stream of components that move more rapidly with the liquid eluent phase compared with the solid adsorbent phase. Thus, a raffinate stream is typically enriched with more polar components, and depleted of less polar components compared with a feed stream.

The term "extract" is well known to the person skilled in the art. In the context of actual and simulated moving bed chromatography it refers to the stream of components that move more rapidly with the solid adsorbent phase compared with the liquid eluent phase. Thus, an extract stream is typically enriched with less polar components, and depleted of more polar components compared with a feed stream.

As used herein the term "nonadjacent" refers to columns, in for example the same apparatus, separated by one or more columns, preferably 3 or more columns, more preferably 5 or more columns, most preferably about 5 columns.

In a second embodiment, the first and second separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the first and second separation steps being carried out in first and second zones respectively, wherein each zone is as defined herein, and wherein the third separation step is carried out in a separate simulated or actual moving bed chromatography apparatus.
In the second embodiment, the third separation step is typically carried out in a simulated or actual moving bed chromatography apparatus comprising a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, and having one or more injection points for a feed mixture stream, one or more injection points for water and/or organic solvent, a raffinate take-off stream from which liquid can be collected from said plurality of linked chromatography columns, and an extract take-off stream from which liquid can be collected from said plurality of linked chromatography columns. This chromatography apparatus typically has only one injection point for a feed mixture. In one embodiment, this chromatography apparatus has only one injection point for the aqueous organic solvent eluent. In another embodiment, this chromatography apparatus has two or more injection points for water and/or organic solvent.

The chromatography apparatus used in the third separation step in the second embodiment typically has a single array of chromatography columns linked in series containing, as eluent, an aqueous organic solvent. Typically, each of the chromatography columns are linked to the two columns in the apparatus adjacent to that column. Thus, the output from a given column is connected to the input of the adjacent column, which is downstream with respect to the flow of eluent in the system. Typically, none of the chromatography columns are linked to non-adjacent columns in the chromatography apparatus.

The chromatography apparatus used in the third separation step in the second embodiment is a separate apparatus from the apparatus used in the first and second separation steps. Thus, two separate apparatuses are used. Eluent circulates separately in the separate chromatographic apparatuses. Thus, eluent is not shared between the separate chromatographic apparatuses other than what eluent may be present as solvent in the second intermediate product which is produced in the second step, and which is then introduced into the chromatographic apparatus used in the third separation step. Chromatographic columns are not shared between the separate chromatographic apparatuses.

After the second intermediate product is obtained in the second separation step, the aqueous organic solvent eluent may be partly or totally removed before the second
intermediate product is purified further in the third separation step. Alternatively, the intermediate product may be purified further in the third step without the removal of any solvent present.

The chromatography apparatus used in the third separation step in the second embodiment is similar to the chromatography apparatus illustrated in Figure 1.

In a third embodiment, the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined herein, and wherein the first separation step is carried out in a separate simulated or actual moving bed chromatography apparatus.

In the third embodiment, the first separation step is typically carried out in a simulated or actual moving bed chromatography apparatus comprising a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, and having one or more injection points for a feed mixture stream, one or more injection points for water and/or organic solvent, a raffmate take-off stream from which liquid can be collected from said plurality of linked chromatography columns, and an extract take-off stream from which liquid can be collected from said plurality of linked chromatography columns. This chromatography apparatus typically has only one injection point for a feed mixture. In one embodiment, this chromatography apparatus has only one injection point for the aqueous organic solvent eluent. In another embodiment, this chromatography apparatus has two or more injection points for water and/or organic solvent.

The chromatography apparatus used in the first separation step in the third embodiment typically has a single array of chromatography columns linked in series containing, as eluent, an aqueous organic solvent. Typically, each of the chromatography columns are linked to the two columns in the apparatus adjacent to that column. Thus, the output from a given column is connected to the input of the adjacent column, which is downstream with respect to the flow of eluent in the
Typically, none of the chromatography columns are linked to non-adjacent columns in the chromatography apparatus.

The chromatography apparatus used in the first separation step in the third embodiment is a separate apparatus from the apparatus used in the second and third separation steps. Thus, two separate apparatuses are used. Eluent is not shared between the separate chromatographic apparatuses other than what eluent may be present as solvent in the first intermediate product which is produced in the first step, and which is introduced into the chromatographic apparatus used in the second separation step. Chromatographic columns are not shared between the separate chromatographic apparatuses.

After the first intermediate product is obtained in the first separation step, the aqueous organic solvent eluent may be partly or totally removed before the intermediate product is purified further in the next separation step. Alternatively, the first intermediate product may be purified further in the second separation step without the removal of any solvent present.

The chromatography apparatus used in the first separation step in the third embodiment is similar to the chromatography apparatus illustrated in Figure 1.

It will be appreciated that in the first, second and third embodiments above two or more separation steps may take place simultaneously in a single simulated or actual moving bed chromatography apparatus having two or three zones, wherein a zone is as defined above. A typical chromatography apparatus having two or more zones, for example two or three zones, is as described in, for example, PCT/GB 10/002339, which is incorporated herein by reference.

In a fourth embodiment, either (a) the first, second and third separation steps are carried out sequentially on the same chromatography apparatus, first and second intermediate products being recovered between the first and second, and second and third separation steps respectively, and the process conditions in the chromatography apparatus being adjusted between the first and second, and second and third separation steps such that saturated and/or monounsaturated fatty acids present in the
feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); or (b) the second separation step is carried out using a different chromatographic apparatus to that used in the first separation step, and/or the third separation step is carried out using a different chromatographic apparatus to that used in the second separation step.

In the fourth embodiment, each of the chromatography apparatuses used to carry out the first, second and third separation steps is typically as defined above for the third separation step in embodiment (2).

In option (b) of the fourth embodiment, all three steps are carried out on separate chromatographic apparatuses. Two or three of the first, second and third separation steps are carried out on two or three different separate chromatographic apparatuses. These may be operated sequentially or simultaneously.

In particular, in option (b) of the fourth embodiment two separate chromatography apparatuses may be operated sequentially to carry out the first and second separation steps. In this case, the first intermediate product is recovered between the first and second separation steps and the process conditions in the first and second chromatography apparatuses are adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

In particular, in option (b) of the fourth embodiment two separate chromatography apparatuses may be operated sequentially to carry out the second and third separation steps. In this case, the second intermediate product is recovered between the second and third separation steps and the process conditions in the second and third chromatography apparatuses are adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).
In particular, in option (b) of the fourth embodiment three separate chromatography apparatuses may be operated sequentially to carry out the first, second and third separation steps. In this case, the first intermediate product is recovered between the first and second separation steps, the second intermediate product is recovered between the second and third separation steps and the process conditions in the first, second and third chromatography apparatuses are adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

In particular, in option (b) of the fourth embodiment, two separate chromatography apparatuses may be operated simultaneously to carry out the first and second separation steps. The first and second separation steps are carried out on separate chromatography apparatuses, the first intermediate product obtained in the first step being introduced into the chromatography apparatus used in the second separation step, and the process conditions in the chromatography apparatuses being adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

In particular, in option (b) of the fourth embodiment, two separate chromatography apparatuses may be operated simultaneously to carry out the second and third separation steps. The second and third separation steps are carried out on separate chromatography apparatuses, the second intermediate product obtained in the second step being introduced into the chromatography apparatus used in the third separation step, and the process conditions in the chromatography apparatuses being adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

In particular, in option (b) of the fourth embodiment, three separate chromatography apparatuses may be operated simultaneously to carry out the first, second and third separation steps. The first, second and third separation steps are carried out on separate chromatography apparatuses, the first intermediate product obtained in the
first step being introduced into the chromatography apparatus used in the second separation step, the second intermediate product obtained in the second step being introduced into the chromatography apparatus used in the third separation step, and the process conditions in the chromatography apparatuses being adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

In particular, in option (b) of the fourth embodiment, two or three separate chromatographic apparatuses are operated. Eluent circulates separately in the separate chromatographic apparatuses. Thus, eluent is not shared between the separate chromatographic apparatuses other than what eluent may be present as solvent in the intermediate product which is purified in the first and/or second step, and which is introduced into the chromatographic apparatus used in the next separation step.

Chromatographic columns are not shared between the separate chromatographic apparatuses used in the first and second and/or second and third separation steps.

After the intermediate product is obtained in the first and/or second separation step, the aqueous organic solvent eluent may be partly or totally removed before the intermediate product is purified further in the next separation step. Alternatively, the intermediate product may be purified further without the removal of any solvent present. These considerations also apply for the second intermediate product obtained in the second separation step in embodiment (2) above, and for the first intermediate product obtained in the first separation step in embodiment (3) above.

In general, any known stationary bed or simulated or actual moving bed chromatography apparatus may be utilised for the purposes of the method of the present invention, as long as the apparatus is used in accordance with the process of the present invention. Those apparatuses described in PCT/GB 10/002339, US 2,985,589, US 3,696,107, US 3,706,812, US 3,761,533, FR-A-2103302, FR-A-265 1148, FR-A-265 1149, US 6,979,402, US 5,069,883 and US 4,764,276 may all be used if configured in accordance with the process of the present invention.
The second, third and fourth embodiments above are preferred. The third and fourth embodiments are more preferred. For certain applications, the third embodiment will be most suitable. In other applications, the fourth embodiment will be most suitable.

The first to fourth embodiments are illustrated in more detail with reference to Figure 2. In all four embodiments in Figure 2 the flow of eluent is from right to left, and the effective flow of adsorbent is from left to right. It can be seen in all cases that the first intermediate product obtained from the first separation step is used as the feed mixture for the second separation step, and the second intermediate product is used as the feed mixture for the third separation step.

Referring now to Figure 2A, this illustrates the first embodiment above, i.e. where the first, second and third separation steps are carried out in a single simulated or actual moving bed chromatography apparatus in first, second and third zones respectively.

The first separation step takes place in the first zone. Then the first intermediate product from the first separation step carried out in the first zone is passed into the second zone as the feed mixture. The second separation step is then carried out in the second zone. The second intermediate product is then passed from the second separation step carried out in the second zone into the third zone as the feed mixture.

The third separation step is then carried out in the third zone.

Referring now to Figure 2B, this illustrates the second embodiment above, i.e. where the first and second separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus in first and second zones respectively, and the third separation step is carried out in a separate simulated or actual moving bed chromatography apparatus. The first separation step takes place in the first zone. Then the first intermediate product from the first separation step carried out in the first zone is passed into the second zone as the feed mixture. The second separation step is carried out in the second zone. The second intermediate product is collected from the second zone. This is then introduced into a chromatography apparatus as the feed mixture for the third separation step.

Referring now to Figure 2C, this illustrates the third embodiment above, i.e. where the second and third separation steps are carried out simultaneously in a single simulated
or actual moving bed chromatography apparatus in first and second zones respectively, and the first separation step is carried out in a separate simulated or actual moving bed chromatography apparatus. The first separation step takes place in a chromatography apparatus. The first intermediate product is collected from the first apparatus. This is then introduced into a separate chromatography apparatus as the feed mixture for the second separation step. The second separation step is carried out in the first zone of the chromatographic apparatus in which the second and third separation steps take place. The second intermediate product from the second separation step carried out in the first zone is passed into the second zone as the feed mixture for the third separation step. The third separation step takes place in the second zone.

Referring now to Figure 2D, this illustrates the fourth embodiment above, i.e. where (a) the first, second and third separation steps are carried out sequentially on the same chromatography apparatus, first and second intermediate products being recovered between the first and second, and second and third separation steps respectively, and the process conditions in the chromatography apparatus being adjusted between the first and second, and second and third separation steps such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); or (b) two or three of the first, second and third separation steps are carried out on two or three different separate apparatuses; wherein the second separation step is carried out using a different chromatographic apparatus to that used in the first separation step, and/or the third separation step is carried out using a different chromatographic apparatus to that used in the second separation step.

When the first separation step comprises purifying the feed mixture in a stationary bed chromatography apparatus, there are several ways in which the three separation steps may be realised. Thus typically, (a) the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined herein; or
(b) the second and third separation steps are carried out sequentially on the same chromatography apparatus, the second intermediate product being recovered between the second and third separation steps and the process conditions in the chromatography apparatus being adjusted between the second and third separation steps such that the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); or
(c) the second and third separation steps are carried out on separate chromatography apparatuses respectively, the intermediate product obtained from the second separation step being introduced into the chromatography apparatus used in the third separation step.

Embodiment (a) above is carried out in a similar manner to the second and third separation steps in embodiment (3) above.

The chromatography apparatuses used in embodiments (b) and (c) above is typically as defined above for the third separation step in embodiment (2). Embodiments (b) and (c) are typically carried out in a similar manner to embodiment (4) above.

It will be appreciated that in certain embodiments, two or three separation steps may be carried out simultaneously in a single chromatography apparatus having two or three zones respectively. In simulated or actual moving bed chromatography apparatuses in which two separation steps are carried out simultaneously in two zones, a raffinate or extract stream is typically collected from a column in the first zone and introduced to a nonadjacent column in the second zone. In simulated or actual moving bed chromatography apparatuses in which three separation steps are carried out simultaneously in three zones, a raffinate or extract stream is typically collected from a column in the first zone and introduced to a nonadjacent column in the second zone, and a raffinate or extract stream is typically collected from a column in the second zone and introduced to a nonadjacent column in the third zone. This enables the first and/or second intermediate products collected in the first and/or second separation steps to be used as the feed mixture for the next separation step.

Typically, the second intermediate product is collected as the raffinate stream in the second separation step, and the PUFA product is collected as the extract stream in the
third separation step; or the second intermediate product is collected as the extract stream in the second separation step, and the PUFA product is collected as the raffmate stream in the third separation step.

5 Preferably, the second intermediate product is collected as the raffmate stream in the second separation step, and the PUFA product is collected as the extract stream in the third separation step.

Typically, in embodiments where the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus in first and second zones respectively, (a) the second intermediate product is collected as a raffmate stream containing the PUFA product together with more polar components from a column in the first zone and introduced to a nonadjacent column in the second zone, where the PUFA product is then collected as the extract stream in the third separation step carried out in the second zone; or (b) the second intermediate product is collected as an extract stream containing the PUFA product together with less polar components from a column in the first zone and introduced to a nonadjacent column in the second zone, where the PUFA product is then collected as the raffmate stream in the third separation step carried out in the second zone.

20 Preferably, in embodiments where the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus in first and second zones respectively, the second intermediate product is collected as a raffmate stream containing the PUFA product together with more polar components from a column in the first zone and introduced to a nonadjacent column in the second zone, where the PUFA product is then collected as the extract stream in the third separation step which is carried out in the second zone.

When the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, the first intermediate product is typically collected as the raffmate stream in the first separation step.

When the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, the first intermediate product is
typically collected as the raffmate stream in the first separation step and (a) the second intermediate product is collected as the raffmate stream in the second separation step, and the PUFA product is collected as the extract stream in the third separation step; or (b) the second intermediate product is collected as the extract stream in the second separation step, and the PUFA product is collected as the raffmate stream in the third separation step.

Typically, the first intermediate product obtained in the first separation step is enriched in the PUFA product compared to the feed mixture; and/or the second intermediate product obtained in the second separation step is enriched in the PUFA product compared to the first intermediate product.

Preferably the first intermediate product obtained in the first separation step is enriched in the PUFA product compared to the feed mixture and the second intermediate product obtained in the second separation step is enriched in the PUFA product compared to the first intermediate product.

Typically, the first intermediate product obtained in the first separation step is depleted in saturated and/or monounsaturated fatty acids compared to the feed mixture.

Typically, in the first step the PUFA product is separated from components of the feed mixture which are less polar than the PUFA product, in the second step the PUFA product is separated from components of the feed mixture which are less polar than the PUFA product but more polar than the components separated in the first separation step, and in the third separation step the PUFA product is separated from components which are more polar than the PUFA product.

Alternatively, in the first step the PUFA product is separated from components of the feed mixture which are less polar than the PUFA product, in the second step the PUFA product is separated from components of the feed mixture which are more polar than the PUFA product, and in the third separation step the PUFA product is separated from components which are less polar than the PUFA product but more polar than the components separated in the first separation step.
The components of the feed mixture separated in the first step which are less polar than the PUFA product are typically unsaturated and/or monounsaturated fatty acids.

The components of the feed mixture which are less polar than the PUFA product but more polar than the components separated in the first separation step typically include DHA or a DHA derivative and/or other PUFAs or PUFA derivatives which are less polar than the PUFA product but more polar than the components separated in the first separation step.

The components of the feed mixture which are more polar than the PUFA product include SDA or an SDA derivative and/or other PUFAs which are more polar than the PUFA product.

PUFAs other than EPA are well known and include ω-3 and ω-6 PUFAs. Examples of ω-3 PUFAs include alpha-linolenic acid (ALA), stearidonic acid (SDA), eicosatrienoic acid (ETE), eicosatetraenoic acid (ETA), docosapentaenoic acid (DPA) and docosahexaenoic acid (DHA). Examples of ω-6 PUFAs include linoleic acid (LA), gamma-linolenic acid (GLA), eicosadienoic acid, dihomogamma-linolenic acid (DGLA), arachidonic acid (ARA), docosadienoic acid, adrenic acid and docosapentaenoic (co-6) acid.

The number of columns used in each separation step is not particularly limited. A skilled person would easily be able to determine an appropriate number of columns to use. The number of columns is typically 4 or more, preferably 6 or more, more preferably 8 or more, for example 4, 5, 6, 7, 8, 9, or 10 columns. In preferred embodiment, 5 or 6 columns, more preferably 6 columns are used. In another preferred embodiment, 7 or 8 columns, more preferably 8 columns are used. Typically, there are no more than 25 columns, preferably no more than 20, more preferably no more than 15.

In embodiments where two separation steps take place simultaneously in a single chromatography apparatus in first and second zones respectively, the number of
columns in each zone is typically 4 or more, preferably 6 or more, more preferably 8 or more, for example 4, 5, 6, 7, 8, 9, or 10 columns.

In embodiments where three separation steps take place simultaneously in a single chromatography apparatus in first, second and third zones respectively, the number of columns in each zone is typically 4 or more, preferably 6 or more, more preferably 8 or more, for example 4, 5, 6, 7, 8, 9, or 10 columns.

The first, second and third separation steps typically involve the same number of columns. For certain applications they may have different numbers of columns.

The dimensions of the columns used are not particularly limited, and will depend on the volume of feed mixture to be purified. A skilled person would easily be able to determine appropriately sized columns to use. The diameter of each column is typically between 10 and 1000mm, preferably between 10 and 500mm, more preferably between 25 and 250mm, even more preferably between 50 and 100 mm, and most preferably between 70 and 80 mm. The length of each column is typically between 10 and 300 cm, preferably between 10 and 200 cm, more preferably between 25 and 150cm, even more preferably between 70 and 110 cm, and most preferably between 80 and 100 cm.

The first, second and third separation steps typically involve columns having identical dimensions but may, for certain applications, they have different dimensions.

The flow rates to the column are limited by maximum pressures across the series of columns and will depend on the column dimensions and particle size of the solid phases. One skilled in the art will easily be able to establish the required flow rate for each column dimension to ensure efficient desorption. Larger diameter columns will in general need higher flows to maintain linear flow through the columns.

For the typical column sizes outlined above, typically the flow rate of eluent into the chromatographic apparatus used in the first or second separation step is from 1 to 4.5L/min, preferably from 1.5 to 2.5 L/min. Typically, the flow rate of the extract from the chromatographic apparatus used in the first or second separation step is from
0.1 to 2.5 L/min, preferably from 0.5 to 2.25 L/min. In embodiments where part of the extract from the first or second separation step is recycled back into the apparatus used in the first or second separation step, the flow rate of recycle is typically from 0.7 to 1.4 L/min, preferably about 1 L/min. Typically, the flow rate of the raffinate from the chromatographic apparatus used in the first or second separation step is from 0.2 to 2.5 L/min, preferably from 0.3 to 2.0 L/min. In embodiments where part of the raffinate from the first or second separation step is recycled back into the apparatus used in the first or second separation step, the flow rate of recycle is typically from 0.3 to 1.0 L/min, preferably about 0.5 L/min. Typically, the flow rate of introduction of the feed mixture into the chromatographic apparatus used in the first or second separation step is from 5 to 150 mL/min, preferably from 10 to 100 mL/min, more preferably from 20 to 60 mL/min.

For the typical column sizes outlined above, typically the flow rate of eluent into the chromatographic apparatus used in the third separation step is from 1 to 4 L/min, preferably from 1.5 to 3.5 L/min. Typically, the flow rate of the extract from the chromatographic apparatus used in the third separation step is from 0.5 to 2 L/min, preferably from 0.7 to 1.9 L/min. In embodiments where part of the extract from the third separation step is recycled back into the apparatus used in the third separation step, the flow rate of recycle is typically from 0.6 to 1.4 L/min, preferably from 0.7 to 1.1 L/min, more preferably about 0.9 L/min. Typically, the flow rate of the raffinate from the chromatographic apparatus used in the third separation step is from 0.5 to 2.5 L/min, preferably from 0.7 to 1.8 L/min, more preferably about 1.4 L/min. In embodiments where part of the raffinate from the third separation step is recycled back into the apparatus used in the third separation step, the flow rate of recycle is typically from 0.3 to 1.0 L/min, preferably about 0.5 L/min.

As the skilled person will appreciate, references to rates at which liquid is collected or removed via the various extract and raffinate streams refer to volumes of liquid removed in an amount of time, typically L/minute. Similarly, references to rates at which liquid is recycled back into an apparatus, typically to an adjacent column in the apparatus, refer to volumes of liquid recycled in an amount of time, typically L/minute.
The step time, i.e. the time between shifting the points of injection of the feed mixture and eluent, and the various take off points of the collected fractions, is not particularly limited, and will depend on the number and dimensions of the columns used, and the flow rate through the apparatus. A skilled person would easily be able to determine appropriate step times to use in the process of the present invention. The step time is typically from 100 to 1000 seconds, preferably from 200 to 800 seconds, more preferably from about 250 to about 750 seconds. In some embodiments, a step time of from 100 to 400 seconds, preferably 200 to 300 seconds, more preferably about 250 seconds, is appropriate. In other embodiments, a step time of from 600 to 900 seconds, preferably 700 to 800 seconds, more preferably about 750 seconds is appropriate.

In the process of the present invention, actual moving bed chromatography is preferred.

Conventional adsorbents known in the art for actual and simulated moving bed systems may be used in the process of the present invention. Each chromatographic column may contain the same or a different adsorbent. Typically, each column contains the same adsorbent. Examples of such commonly used materials are polymeric beads, preferably polystyrene reticulated with DVB (divinylbenzene); and silica gel, preferably reverse phase bonded silica gel with C8 or C18 alkanes, especially C18. C18 bonded reverse phase silica gel is preferred. The adsorbent used in the process of the present invention is preferably non-polar.

The shape of the adsorbent stationary phase material may be, for example, spherical or nonspherical beads, preferably substantially spherical beads. Such beads typically have a diameter of 5 to 500 microns, preferably 10 to 500 microns, more preferably 15 to 500 microns, more preferably 40 to 500 microns, more preferably 100 to 500 microns, more preferably 250 to 500 microns, even more preferably 250 to 400 microns, most preferably 250 to 350 microns. In some embodiments, beads with a diameter of 5 to 35 microns may be used, typically 10 to 30 microns, preferably 15 to 25 microns. Some preferred particle sizes are somewhat larger than particle sizes of beads used in the past in simulated and actual moving bed processes. Use of larger particles enables a lower pressure of eluent to be used in the system. This, in turn, has
advantages in terms of cost savings, efficiency and lifetime of the apparatus. It has surprisingly been found that adsorbent beads of large particle size may be used in the process of the present invention (with their associated advantages) without any loss in resolution.

The adsorbent typically has a pore size of from 10 to 50 nm, preferably 15 to 45 nm, more preferably 20 to 40 nm, most preferably 25 to 35 nm.

Typically, the process of the present invention is conducted at from 15 to 55°C, preferably at from 20 to 40°C, more preferably at about 30°C. Thus, the process is typically carried out at room temperature, but may be conducted at elevated temperatures.

As mentioned above, saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii). This is typically effected by adjusting the process conditions in the chromatography apparatus, or zone in a chromatography apparatus in which the first, second and third separation steps are carried out.

Thus, the process conditions in the first, second and third separation steps typically vary. The process conditions which vary may include, for example, the size of the columns used, the number of columns used, the packing used in the columns, the step time of the SMB apparatus, the temperature of the apparatus, the eluent used in the separation steps, or the flow rates used in the apparatus, in particular the recycle rate of liquid collected via the extract or raffmate streams.

Preferably the process conditions which vary are the water:organic solvent ratio of the eluent used in the separation steps, and/or the recycle rate of liquid collected via the extract or raffmate streams in the separation steps. Both of these options are discussed in more detail below.

Typically, part of the extract stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and/or part
of the raffinate stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and/or part of the extract stream from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step; and/or part of the raffinate stream from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step.

Preferably, part of the extract stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and part of the raffinate stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and part of the extract stream from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step; and part of the raffinate stream from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step.

When the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, typically part of the extract stream from the apparatus used in the first separation step is recycled back into the apparatus used in the first separation step; and/or part of the raffinate stream from the apparatus used in the first separation step is recycled back into the apparatus used in the first separation step.

When the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, preferably part of the extract stream from the apparatus used in the first separation step is recycled back into the apparatus used in the first separation step; and part of the raffinate stream from the apparatus used in the first separation step is recycled back into the apparatus used in the first separation step; and part of the extract stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and part of the raffinate stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and part of the extract stream from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step; and part of the raffinate stream
from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step.

This recycle involves feeding part of the extract or raffmate stream out of the chromatography apparatus used in the first, second or third separation step back into the apparatus used in that step, typically into an adjacent column. This adjacent column is the adjacent column which is downstream with respect to the flow of eluent in the system.

When two or three separation steps are carried out simultaneously in a single chromatography in two or three zones respectively, this recycle involves recycling the particular extract or raffmate stream removed from a zone back into the same zone.

The rate at which liquid collected via the extract or raffmate stream in a particular separation step is recycled back into a chromatography apparatus or zone used in that separation step is the rate at which liquid collected via that stream is fed back into the apparatus used in that step, typically into an adjacent column, i.e. the downstream column with respect to the flow of eluent in the system.

This can be seen with reference to a preferred embodiment in Figure 5. The rate of recycle of extract in the first separation step is the rate at which extract collected from the bottom of column 2 of the chromatographic apparatus used in the first separation step is fed into the top of column 3 of the chromatographic apparatus used in the first separation step, i.e. the flow rate of liquid into the top of column 3 of the chromatographic apparatus used in the first separation step.

The rate of recycle of extract in the second separation step is the rate at which extract collected at the bottom of column 10 of the chromatographic apparatus used in the second separation step is fed into the top of column 11 of the chromatographic apparatus used in the second separation step, i.e. the flow rate of liquid into the top of column 11 of the chromatographic apparatus used in the second separation step.

The rate of recycle of extract in the third separation step is the rate at which extract collected at the bottom of column 19 of the chromatographic apparatus used in the
second separation step is fed into the top of column 19 of the chromatographic apparatus used in the second separation step, i.e. the flow rate of liquid into the top of column 19 of the chromatographic apparatus used in the second separation step.

Recycle of the extract and/or raffinate streams in the first, second and/or third separation steps is typically effected by feeding the liquid collected via that stream in that separation step into a container, and then pumping an amount of that liquid from the container back into the apparatus or zone used in that separation step, typically into an adjacent column. In this case, the rate of recycle of liquid collected via a particular extract or raffinate stream in the first and/or second separation steps, typically back into an adjacent column, is the rate at which liquid is pumped out of the container back into the chromatography apparatus or zone, typically into an adjacent column.

As the skilled person will appreciate, the amount of liquid being introduced into a chromatography apparatus via the eluent and feedstock streams is balanced with the amount of liquid removed from the apparatus, and recycled back into the apparatus.

Thus, with reference to Figure 5, for the extract stream, the flow rate of eluent (desorbent) into the chromatographic apparatus(es) used in the second and third separation steps (D) is equal to the rate at which liquid collected via the extract stream in that separation step accumulates in a container (E2 and E3) added to the rate at which extract is recycled back into the chromatographic apparatus used in that particular separation step (D-E2 and D-E3).

For the raffinate stream from a separation step, the rate at which extract is recycled back into the chromatographic apparatus used in that particular separation step (D-El and D-E2) added to the rate at which feedstock is introduced into the chromatographic apparatus used in that particular separation step (F and R1) is equal to the rate at which liquid collected via the raffinate stream in that particular separation step accumulates in a container (R1 and R2) added to the rate at which raffinate is recycled back into the chromatographic apparatus used in that particular separation step (D+F-El-R1 and D+R1-E2-R2).
The rate at which liquid collected from a particular extract or raffinate stream from a chromatography apparatus or zone accumulates in a container can also be thought of as the net rate of removal of that extract or raffinate stream from that chromatography apparatus.

Typically, the rate at which liquid collected via one or both of the extract and raffinate streams in the second separation step is recycled back into the apparatus used in that separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); and/or wherein the rate at which liquid collected via one or both of the extract and raffinate streams in the third separation step is recycled back into the apparatus used in that separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

Preferably, the rate at which liquid collected via one or both of the extract and raffinate streams in the second separation step is recycled back into the apparatus used in that separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); and wherein the rate at which liquid collected via one or both of the extract and raffinate streams in the third separation step is recycled back into the apparatus used in that separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

When the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, the rate at which liquid collected via one or both of the extract and raffinate streams in the first separation step is recycled back into the apparatus used in that separation step is typically adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed
in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

Typically, the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step differs from the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatography apparatus used in the third separation step; and/or the rate at which liquid collected via the raffinate stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step differs from the rate at which liquid collected via the raffinate stream in the third separation step is recycled back into the chromatography apparatus used in the third separation step.

When the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, the rate at which liquid collected via the extract stream in the first separation step is recycled back into the chromatography apparatus used in the first separation step typically differs from the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step; and/or the rate at which liquid collected via the raffinate stream in the first separation step is recycled back into the chromatography apparatus used in the first separation step typically differs from the rate at which liquid collected via the raffinate stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step.

Varying the rate at which liquid collected via the extract and/or raffinate streams in the first, second and/or third separation steps is recycled back into the apparatus used in that particular separation step has the effect of varying the amount of more polar and less polar components present in the extract and raffinate streams. Thus, for example, a lower extract recycle rate results in fewer of the less polar components in that separation step being carried through to the raffinate stream. A higher extract recycle rate results in more of the less polar components in that separation step being carried through to the raffinate stream.
This can be seen, for example, in the specific embodiment of the invention shown in Figure 5. The rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in that separation step (D-E2) will affect to what extent any of component A is carried through to the raffinate stream in the second separation step (R2).

Typically, the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step. Preferably, a raffinate stream containing the PUFA product together with more polar components is collected from the second separation step and purified in the third separation step, and the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

Alternatively, the rate at which liquid collected via the raffinate stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the raffinate stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step. Preferably, an extract stream containing the PUFA product together with less polar components is collected from the second separation step and purified in the third separation step, and the rate at which liquid collected via the raffinate stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the raffinate stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

Where recycle rates are adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and
the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii), the water:organic solvent ratio of the eluents used in the separation steps where the recycle rates differ may be the same or different.

The eluent used in the process of the present invention is an aqueous organic solvent.

The aqueous organic solvent typically comprises water and one or more alcohols, ethers, esters, ketones or nitriles, or mixtures thereof.

Alcohol solvents are well known to the person skilled in the art. Alcohols are typically short chain alcohols. Alcohols typically are of formula ROH, wherein R is a straight or branched Ci-C₆ alkyl group. The Ci-C₆ alkyl group is preferably unsubstituted. Examples of alcohols include methanol, ethanol, n-propanol, i-propanol, n-butanol, i-butanol, s-butanol and t-butanol. Methanol and ethanol are preferred. Methanol is more preferred.

Ether solvents are well known to the person skilled in the art. Ethers are typically short chain ethers. Ethers typically are of formula R-O-R, wherein R and R are the same or different and represent a straight or branched Ci-C₆ alkyl group. The Ci-C₆ alkyl group is preferably unsubstituted. Preferred ethers include diethylether, diisopropylether, and methyl t-butyl ether (MTBE).

Ester solvents are well known to the person skilled in the art. Esters are typically short chain esters. Esters typically are of formula R-(C=O)0-R, wherein R and R are the same or different and represent a straight or branched Ci-C₆ alkyl group. Preferred esters include methylacetate and ethylacetate.

Ketone solvents are well known to the person skilled in the art. Ketones are typically short chain ketones. Ketones typically are of formula R-(C=O)-R, wherein R and R are the same or different and represent a straight or branched Ci-C₆ alkyl group. The Ci-C₆ alkyl group is preferably unsubstituted. Preferred ketones include acetone, methylethylketone and methyl isobutyl ketone (MIBK).
Nitrile solvents are well known to the person skilled in the art. Nitriles are typically short chain nitriles. Nitriles typically are of formula R-CN, wherein R represents a straight or branched C$_i$-C$_6$ alkyl group. The C$_i$-C$_6$ alkyl group is preferably unsubstituted. Preferred nitriles include acetonitrile.

Typically, the aqueous organic solvent is aqueous alcohol or aqueous acetonitrile. The aqueous organic solvent is preferably aqueous methanol or aqueous acetonitrile. Aqueous methanol is more preferred.

Typically, the eluent is not in a supercritical state. Typically, the eluent is a liquid.

Typically, the average water:organic solvent ratio, for example water:methanol ratio, of the eluent in the entire apparatus is from 0.1:99.9 to 9:91 wt%, preferably from 0.25:99.75 to 7:93 wt%, more preferably from 0.5:99.5 to 6:94 wt%.

When the aqueous organic solvent is aqueous acetonitrile, the eluent typically contains up to 30 wt% water, remainder acetonitrile. Preferably, the eluent contains from 5 to 25 wt% water, remainder acetonitrile. More preferably, the eluent contains from 10 to 20 wt% water, remainder acetonitrile. Even more preferably, the eluent contains from 15 to 25 wt% water, remainder acetonitrile.

Typically, the water:organic solvent ratio used in each separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step; and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

Typically, the aqueous organic solvent eluent used in two or more of the separation steps has a different water:organic solvent ratio. In one embodiment, the water:organic solvent ratio used in each separation step has a different water:organic solvent ratio.

The eluting power of the eluent used in two or more of the separation steps is typically different. Depending on the choice of organic solvent, they may be more
powerful desorbers than water. Alternatively, they may be less powerful desorbers than water. Acetonitrile and alcohols, for example, are more powerful desorbers than water.

In a preferred embodiment, the aqueous organic solvent eluent used in the second and third separation steps has the same water:organic solvent ratio, and the aqueous organic solvent eluent used in the first separation step has a different water:organic solvent ratio from the organic solvent eluent used in the second and third separation steps.

In this preferred embodiment, the eluting power of the eluent used in the second and third separation steps is the same; and/or the eluting power of the eluent used in the first separation step is greater than that of the eluent used in the second separation step. Preferably in this embodiment, the eluting power of the eluent used in the second and third separation steps is the same; and the eluting power of the eluent used in the first separation step is greater than that of the eluent used in the second and third separation steps. In this embodiment, when the aqueous organic solvent is aqueous alcohol or acetonitrile, the amount of alcohol or acetonitrile in the eluent used in the second and third separation steps is typically the same, and the amount of alcohol or acetonitrile in the eluent used in the first separation step is typically greater than the amount of alcohol or acetonitrile in the eluent used in the second and third separation steps. Thus, in this embodiment, the waternorganic solvent ratio of the eluent in the second and third separation steps is typically the same, and the water:organic solvent ratio of the eluent in the first separation step is typically lower than the water:organic solvent ratio of the eluent in the second and third separation steps.

In this preferred embodiment, the water:organic solvent ratio of the eluent in the first separation step is typically from 0:100 to 5:95 wt%, preferably from 0.1:99.9 to 2.5:97.5 wt%, more preferably from 0.1:99.9 to 2.98 wt%, even more preferably from 0.1:99.9 to 1:99 wt%, even more preferably from 0.25:99.75 to 0.75:99.25 wt%, and most preferably about 0.5:99.5. In this preferred embodiment, the waternorganic solvent ratio of the eluent in the second and third separation steps is typically from 5:95 to 11:89 wt%, preferably 6:94 to 10:90 wt%, more preferably from 7:93 to 9:91
wt%, even more preferably from 7.5:92.5 to 8.5:91.5 wt%, and most preferably about
8:92 wt%.

In this preferred embodiment, the water:organic solvent ratio of the eluent used in the first
separation step is preferably from 0.1:99.9 to 1:99 wt%, and the water:organic solvent ratio
of the eluent used in the second and third separation steps is preferably from 7:93 to 9:91 wt%.

In an alternative embodiment, the aqueous organic solvent eluent used in each separation step has a different water:organic solvent ratio.

In this alternative embodiment, the eluting power of the eluent used in the first separation step is greater than that of the eluent used in the second separation step; and/or the eluting power of the eluent used in the second separation step is greater than that of the eluent used in the third separation step. Preferably, a raffinate stream containing the PUFA product together with more polar components is collected from the second separation step and purified in the third separation step and the eluting power of the eluent used in the second separation step is greater than that of the eluent used in the third separation step. Alternatively, an extract stream containing the PUFA product together with less polar components is collected from the second separation step and purified in the third separation step and the eluting power of the eluent used in the second separation step is lower than that of the eluent used in the third separation step.

In practice this is achieved by varying the relative amounts of water and organic solvent used in each separation step. In this embodiment, when the aqueous organic solvent is aqueous alcohol or acetonitrile, the amount of alcohol or acetonitrile in the eluent used in the first separation step is typically greater than the amount of alcohol or acetonitrile in the eluent used in the second separation step; and/or the amount of alcohol or acetonitrile in the eluent used in the second separation step is typically greater than the amount of alcohol or acetonitrile in the eluent used in the third separation step. Thus, in this embodiment, the water:organic solvent ratio of the eluent in the first separation step is typically lower than the water:organic solvent ratio of the eluent in the second separation step; and/or the water:organic solvent ratio of
the eluent in the second separation step is typically lower than the water:organic solvent ratio of the eluent in the third separation step.

It will be appreciated that the ratios of water and organic solvent in each separation step referred to above are average ratios within the totality of the chromatographic apparatus.

Typically, the water:organic solvent ratio of the eluent in each separation step is controlled by introducing water and/or organic solvent into one or more columns in the chromatographic apparatuses used in the separation steps. Thus, for example, to achieve a lower water:organic solvent ratio in the first separations step than in the second and third separation steps, water is typically introduced more slowly into the chromatographic apparatus used in the first separation step than in the second and third separation steps.

In some embodiments, essentially pure organic solvent and essentially pure water may be introduced at different points in the chromatographic apparatus used in each separation step. The relative flow rates of these two streams will determine the overall solvent profile in the chromatographic apparatus. In other embodiments, different organic solvent/water mixtures may be introduced at different points in each chromatographic apparatus used in each separation step. That will involve introducing two or more different organic solvent/water mixtures into the chromatographic apparatus used in a particular separation step, each organic solvent/water mixture having a different organic solvent:water ratio. The relative flow rates and relative concentrations of the organic solvent/water mixtures in this embodiment will determine the overall solvent profile in the chromatographic apparatus used in that separation step.

Preferably, the aqueous organic solvent eluent used in the second and third separation steps has the same water:organic solvent ratio, and the aqueous organic solvent eluent used in the first separation step has a different water:organic solvent ratio from the organic solvent eluent used in the second and third separation steps; and the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step differs
from the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatography apparatus used in the third separation step.

More preferably, the water:organic solvent ratio of the eluent in the second and third separation steps is the same, and the water:organic solvent ratio of the eluent in the first separation step is lower than the water:organic solvent ratio of the eluent in the second and third separation steps; and the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

Even more preferably,

the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus;

the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined herein, and wherein the first separation step is carried out in a separate simulated or actual moving bed chromatography apparatus;

the first intermediate product is collected as the raffinate stream in the first separation step, the second intermediate product is collected as the raffinate stream in the second separation step, and the PUFA product is collected as the extract stream in the third separation step;

the second intermediate product raffinate stream containing the PUFA product together with more polar components is collected from a column in the first zone and introduced into a nonadjacent column in the second zone;

the aqueous organic solvent eluent used in the second and third separation steps has the same water:organic solvent ratio, and the water:organic solvent ratio of the eluent used in the first separation step is lower than the water:organic solvent ratio of the eluent used in the second and third separation steps; and
the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

It is preferred that the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus; and the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined herein, and wherein the first separation step is carried out in a separate simulated or actual moving bed chromatography apparatus. A preferred embodiment of this is illustrated in Figure 3.

A feed mixture F comprising the PUFA product (B) and more polar (C) and less polar (A’) and (A) components is purified in the first separation step. In the first separation step, the least polar components (e.g. saturates and/or monounsaturates) (A) are removed as extract stream E1. The PUFA product (B), more polar components (C) and less polar (but more polar than (A’)) components (A) are collected as raffmate stream R1. Raffmate stream R1 is the intermediate product which is then purified in the second separation step.

In the second separation step, the less polar components (A) are removed as extract stream E2. The PUFA product (B) and more polar components (C) are collected as raffmate stream R2. Raffmate stream R2 is the intermediate product which is then purified in the third separation step.

In the third separation step, more polar components (C) are removed as raffmate stream R3. The PUFA product (B) is collected as extract stream E3. The second and third separation steps take place in two zones in a single SMB chromatographic apparatus.
This embodiment is illustrated in more detail in Figure 4. Figure 4 is identical to Figure 2, except that the points of introduction of the aqueous organic solvent desorbent (D) into each chromatographic apparatus are shown.

Typical solvents for use in this most preferred embodiment are aqueous alcohols or aqueous acetonitrile, preferably aqueous methanol.

Typically in this preferred embodiment, the aqueous organic solvent eluent used in the second and third separation steps has the same water:organic solvent ratio, and the aqueous organic solvent eluent used in the first separation step has a different water:organic solvent ratio from the organic solvent eluent used in the second and third separation steps; and the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step differs from the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatography apparatus used in the third separation step.

Preferably in this preferred embodiment, the water:organic solvent ratio of the eluent in the second and third separation steps is the same, and the water:organic solvent ratio of the eluent in the first separation step is lower than the water:organic solvent ratio of the eluent in the second and third separation steps; and the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

In this preferred embodiment the first raffmate stream in the first separation step is typically removed downstream of the point of introduction of the feed mixture into the chromatographic apparatus used in the first separation step, with respect to the flow of eluent.

In this particularly preferred embodiment, the first extract stream in the first separation step is typically removed upstream of the point of introduction of the feed
mixture into the chromatographic apparatus used in the first separation step, with respect to the flow of eluent.

In this particularly preferred embodiment, the second raffinate stream in the second separation step is typically removed downstream of the point of introduction of the first intermediate product into the chromatographic apparatus used in the second separation step, with respect to the flow of eluent.

In this particularly preferred embodiment, the second extract stream in the second separation step is typically collected upstream of the point of introduction of the first intermediate product into the chromatographic apparatus used in the second separation step, with respect to the flow of eluent.

In this particularly preferred embodiment, the third raffinate stream in the third separation step is typically removed downstream of the point of introduction of the second intermediate product into the chromatographic apparatus used in the third separation step, with respect to the flow of eluent.

In this particularly preferred embodiment, the third extract stream in the third separation step is typically collected upstream of the point of introduction of the second intermediate product into the chromatographic apparatus used in the third separation step, with respect to the flow of eluent.

Typically in this preferred embodiment, the aqueous organic solvent is introduced into the chromatographic apparatus used in the first separation step upstream of the point of removal of the first extract stream, with respect to the flow of eluent.

Typically in this preferred embodiment, the aqueous organic solvent is introduced into the chromatographic apparatus used in the second separation step upstream of the point of removal of the second extract stream, with respect to the flow of eluent.

Typically in this preferred embodiment, the aqueous organic solvent is introduced into the chromatographic apparatus used in the third separation step upstream of the point of removal of the third extract stream, with respect to the flow of eluent.
A more preferred embodiment of the invention illustrated in Figures 3 and 4 is shown in Figure 5. This illustrates the number of columns used in each separation step, and shows typical points of introduction of feed mixtures and eluents, and typical points of removal of extract and raffinate streams.

Thus, in this more preferred embodiment, the SMB chromatography apparatus used in the first separation step consists of eight chromatographic columns, 1 to 8. The SMB chromatography apparatus used in the second separation step consists of eight chromatographic columns, 9 to 16. The SMB chromatography apparatus used in the third separation step consists of seven chromatographic columns, 17 to 23.

In each apparatus the columns are typically arranged in series so that (in the case of the first separation step) the bottom of column 1 is linked to the top of column 2, the bottom of column 2 is linked to the top of column 3... etc. ..and the bottom of column 8 is linked to the top of column 1. These linkages may optionally be via a holding container, with a recycle stream into the next column. The flow of eluent through the system is from column 1 to column 2 to column 3 etc. The effective flow of adsorbent through the system is from column 8 to column 7 to column 6 etc.

In this more preferred embodiment, a feed mixture F comprising the PUFA product (B) and more polar (C) and less polar (A') and (A) components is introduced into the top of column 5 in the chromatographic apparatus used in the first separation step. Aqueous organic solvent desorbent is introduced into the top of column 1 of the chromatographic apparatus used in the first separation step. In the first separation step, the least polar components (e.g. saturates and/or monounsaturates) (A) are removed as extract stream E1 from the bottom of column 2. The PUFA product (B), more polar components (C) and less polar (but more polar than (A')) components (A) are collected as raffinate stream R1 from the bottom of column 6.

Raffinate stream R1 is the first intermediate product which is then purified in the second separation step, by being introduced into the chromatographic apparatus used in the second separation step at the top of column 13. Aqueous organic solvent
Desorbent is introduced into the top of column D in the chromatographic apparatus used in the second separation step.

In the second separation step, the less polar components (A) are removed as extract stream E2 at the bottom of column 10. The PUFA product (B) and more polar components (C) are collected as raffinate stream R2 at the bottom of column 14. Raffinate stream R2 is the intermediate product which is then purified in the third separation step, by being introduced into the chromatographic apparatus used in the second separation step at the top of column 21.

In the third separation step, more polar components (C) are removed as raffinate stream R3 at the bottom of column 22. The PUFA product (B) is collected as extract stream E3 at the bottom of column 18. The second and third separation steps take place in two zones in a single SMB chromatographic apparatus.

In this more preferred embodiment, aqueous organic solvent is typically introduced into the top of column 1 of the chromatographic apparatus used in the first separation step.

In this more preferred embodiment, aqueous organic solvent is typically introduced into the top of column 9 of the chromatographic apparatus used in the second separation step.

In this more preferred embodiment, aqueous organic solvent is typically introduced into the top of column 17 of the chromatographic apparatus used in the third separation step.

In this more preferred embodiment, the feed stream is typically introduced into the top of column 5 of the chromatographic apparatus used in the first separation step.

In this more preferred embodiment, a first raffinate stream is typically collected as the first intermediate product from the bottom of column 6 of the chromatographic apparatus used in the first separation step. This first intermediate product is then purified in the second separation step and is typically introduced into the top of
column 13 of the chromatographic apparatus used in the second separation step. The first raffinate stream may optionally be collected in a container before being purified in the second separation step.

In this more preferred embodiment, a first extract stream is typically removed from the bottom of column 2 of the chromatographic apparatus used in the first separation step. The first extract stream may optionally be collected in a container and reintroduced into the top of column 3 of the chromatographic apparatus used in the first separation step.

In this more preferred embodiment, a second raffinate stream is typically collected as the second intermediate product from the bottom of column 14 of the chromatographic apparatus used in the second separation step. This second intermediate product is then purified in the third separation step and is typically introduced into the top of column 21 of the chromatographic apparatus used in the third separation step. The second raffinate stream may optionally be collected in a container before being purified in the second separation step.

In this more preferred embodiment, a second extract stream is typically removed from the bottom of column 10 of the chromatographic apparatus used in the second separation step.

In this more preferred embodiment, a third extract stream is typically collected from the bottom of column 18 of the chromatographic apparatus used in the third separation step. This third extract stream typically contains the purified PUFA product. The third extract stream may optionally be collected in a container and reintroduced into the top of column 19 of the chromatographic apparatus used in the third separation step.

In this more preferred embodiment, a third raffinate stream is typically removed from the bottom of column 22 of the chromatographic apparatus used in the third separation step.
Typically in this more preferred embodiment, the aqueous organic solvent eluent used in the second and third separation steps has the same water:organic solvent ratio, and the aqueous organic solvent eluent used in the first separation step has a different water:organic solvent ratio from the organic solvent eluent used in the second and third separation steps; and the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step differs from the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatography apparatus used in the third separation step.

Preferably in this more preferred embodiment, the water:organic solvent ratio of the eluent in the second and third separation steps is the same, and the water:organic solvent ratio of the eluent in the first separation step is lower than the water:organic solvent ratio of the eluent in the second and third separation steps; and the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

In this more preferred embodiment, the water:organic solvent ratio of the eluent used in the second and third separation steps is the same and is from 7:93 to 9:91 wt%, and the water:organic solvent ratio of the eluent in the first separation step is from 0.1:99.9 to 1:99 wt%.

Although these preferred and more preferred embodiments are shown as for Figure 2C discussed above, they may also be carried out with apparatuses configured such that:

the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, and the first, second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the first, second and third
separation steps being carried out in first, second and third zones respectively, wherein each zone is as defined herein; or

the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, and the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined herein, and wherein the first separation step is carried out in a separate simulated or actual moving bed chromatography apparatus; or

the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, and (a) the first, second and third separation steps are carried out sequentially on the same chromatography apparatus, first and second intermediate products being recovered between the first and second, and second and third separation steps respectively, and the process conditions in the chromatography apparatus being adjusted between the first and second, and second and third separation steps such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); or

(b) the second separation step is carried out using a different chromatographic apparatus to that used in the first separation step, and/or the third separation step is carried out using a different chromatographic apparatus to that used in the second separation step; or

the first separation step comprises purifying the feed mixture in a stationary bed chromatography apparatus, and the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined herein; or

the first separation step comprises purifying the feed mixture in a stationary bed chromatography apparatus, and the second and third separation steps are carried out sequentially on the same chromatography apparatus, the second intermediate product being recovered between the second and third separation steps and the process
conditions in the chromatography apparatus being adjusted between the second and third separation steps such that the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); or the first separation step comprises purifying the feed mixture in a stationary bed chromatography apparatus, and the second and third separation steps are carried out on separate chromatography apparatuses respectively, the intermediate product obtained from the second separation step being introduced into the chromatography apparatus used in the third separation step.

The process of the invention allows much higher purities of PUFA product to be achieved than have been possible with conventional chromatographic techniques. PUFA products produced by the process of the invention also have particularly advantageous impurity profiles, which are quite different from those observed in oils prepared by known techniques. The present invention therefore also relates to compositions comprising a PUFA product, for example one obtainable by the process of the present invention.

In practice, the process of the present invention will generally be controlled by a computer. The present invention therefore also provides a computer program for controlling a chromatographic apparatus as defined herein, the computer program containing code means that when executed instruct the apparatus to carry out the process of the invention.

The following Examples illustrate the invention.
EXAMPLES

Example 1

A fish oil derived feedstock (55 weight% EPA EE, 5 weight% DHA EE) is fractionated using an actual moving bed chromatography system using bonded C18 silica gel as stationary phase and aqueous methanol as eluent according to the system schematically illustrated in Figure 5. A GC trace of the feed mixture is shown as Figure 7.

In a first separation step, the feed mixture was passed through an SMB apparatus having 8 columns 1 to 8 (diameter: 152mm, length: 813mm) connected in series as shown Figure 5. Process conditions were adjusted to remove saturated and monounsaturated components from the feed mixture as the extract stream. A 0.5:99.5 wt% water:methanol eluent was used. The raffinate stream was retained as the first intermediate product. A GC trace of the first intermediate product is shown as Figure 8.

The first intermediate product was passed through an SMB apparatus having two zones with eight columns, columns 9 to 16, in the first zone and seven columns, columns 17 to 23, in the second zone. An 8:92 wt% water:methanol eluent was used in both first and second zones, i.e. in both the second and third separation steps. The process conditions in the first zone were adjusted to purify EPA from the slower running components such as DHA, which were removed as the extract stream. The raffinate stream was retained as the second intermediate product. A GC trace of the second intermediate product is shown as Figure 9.

The second intermediate product was then introduced into the second zone and separated from the faster running components, which were removed as a raffinate stream. High purity EPA was collected as the extract stream from the second zone. A GC trace of the EPA PUFA product is shown as Figure 10.

EPA was produced with a final purity of greater than 97%.
It can be seen that for the three separation steps taken together, the overall rate of accumulation of extract (E1+E2+E3) 3876 ml/min.

The process conditions for each separation step are as follows:

**First separation step**
- Feedstock feed rate: 94 ml/min
- Desorbent feed rate: 6250 ml/min
- Extract accumulation rate: 1250 ml/min
- Extract recycle rate: 5000 ml/min
- Raffmator accumulation rate: 1688 ml/min
- Cycle time: 600 sees

**Second separation step**
- First intermediate product feed rate: 40 ml/min
- Desorbent feed rate: 6313 ml/min
- Extract accumulation rate: 1188 ml/min
- Extract recycle rate: 5125 ml/min
- Raffmator accumulation rate: 1625 ml/min
- Cycle time: 1200 sees

**Third separation step**
- Second intermediate product feed rate: 40 ml/min
- Desorbent feed rate: 6189 ml/min
- Extract accumulation rate: 1438 ml/min
- Extract recycle rate: 4750 ml/min
- Raffmator accumulation rate: 1438 ml/min
- Cycle time: 1080 sees
Comparative Example 1

An experiment was carried out to produced a PUFA product containing greater than 97% EPA from the same feed mixture as was used in Example 1. However, instead of using a three step separation process in accordance with the present invention, only two separation steps were used. Thus, the process was carried out in accordance with the process disclosed in PCT/GB 10/002339, and as illustrated in Figure 6.

A single chromatographic apparatus having two zones was used as shown in Figure 6. The first zone contains 8 columns (diameter: 24", length: 32") and the second zone 7 columns (diameter: 24", length: 32"). Process conditions were adjusted to separate the EPA PUFA product from less polar components of the feed mixture in the first zone, and more polar components of the feed mixture in the second zone. An 8:92 wt% water methanol eluent was used in both zones.

EPA was produced with a final purity of greater than 97%.

It can be seen that for the two separation steps taken together, the overall rate of accumulation of extract (E1+E2) was 10571 ml/min. Thus, it can be seen that a much higher volume of aqueous organic solvent is required to recover the PUFA product compared with the three step process of the invention.

The process conditions for the separation steps are as follows:

First separation step

Feed mixture feed rate: 34 ml/min
Desorbent feed rate: 14438 ml/min
Extract accumulation rate: 9313 ml/min
Extract recycle rate: 5125 ml/min
Raffinate accumulation rate: 1688ml/min
Cycle time: 1200 sees
Third separation step

Intermediate product feed rate: 40 ml/min
Desorbent feed rate: 6189 ml/min
Extract accumulation rate: 1438 ml/min
Extract recycle rate: 4750 ml/min
Raffinate accumulation rate: 1438 ml/min
Cycle time: 1080 sees
CLAIMS

1. A chromatographic separation process for recovering a polyunsaturated fatty acid (PUFA) product from a feed mixture which is a fish oil or which is derived from fish oil, which process comprises the steps of:
   (i) purifying the feed mixture in a chromatographic separation step, to obtain a first intermediate product; and
   (ii) purifying the first intermediate product obtained in (i) in a simulated or actual moving bed chromatographic separation step, to obtain a second intermediate product; and
   (iii) purifying the second intermediate product obtained in (ii) in a simulated or actual moving bed chromatographic separation step, to obtain the PUFA product; wherein an aqueous organic solvent is used as eluent in each separation step; saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step; the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); and the PUFA product obtained in the third separation step contains EPA or an EPA derivative in an amount greater than 90 wt%.

2. A process according to claim 1, wherein the first separation step comprises purifying the feed mixture in a stationary bed or simulated or actual moving bed chromatography apparatus.

3. A process according to claim 1 or 2, wherein the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus; and wherein the first, second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the first, second and third separation steps being carried out in first, second and third zones respectively, wherein each zone has one or more injection points for a feed mixture stream, one or more injection points for water and/or organic solvent,
a raffinate take-off stream from which liquid can be collected from said zone, and an extract take-off stream from which liquid can be collected from said zone.

5. A process according to claim 1 or 2, wherein the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus; and wherein the first and second separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the first and second separation steps being carried out in first and second zones respectively, wherein each zone is as defined in claim 3, and wherein the third separation step is carried out in a separate simulated or actual moving bed chromatography apparatus.

6. A process according to claim 1 or 2, wherein the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus; and wherein (a) the first, second and third separation steps are carried out sequentially on the same chromatography apparatus, first and second intermediate products being recovered between the first and second, and second and third separation steps respectively, and the process conditions in the chromatography apparatus being adjusted between the first and second, and second and third separation
steps such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); or

(b) the second separation step is carried out using a different chromatographic apparatus to that used in the first separation step, and/or the third separation step is carried out using a different chromatographic apparatus to that used in the second separation step.

7. A process according to claim 1 or 2, wherein the first separation step comprises purifying the feed mixture in a stationary bed chromatography apparatus; and wherein

(a) the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined in claim 3; or

(b) the second and third separation steps are carried out sequentially on the same chromatography apparatus, the second intermediate product being recovered between the second and third separation steps and the process conditions in the chromatography apparatus being adjusted between the second and third separation steps such that the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); or

(c) the second and third separation steps are carried out on separate chromatography apparatuses respectively, the intermediate product obtained from the second separation step being introduced into the chromatography apparatus used in the third separation step.

8. A process according to any one of claims 3 to 7, wherein in simulated or actual moving bed chromatography apparatuses in which two separation steps are carried out simultaneously in two zones, a raffmate or extract stream is collected from a column in the first zone and introduced to a nonadjacent column in the second zone; and/or
wherein in simulated or actual moving bed chromatography apparatuses in which three separation steps are carried out simultaneously in three zones, a raffmate or extract stream is collected from a column in the first zone and introduced to a nonadjacent column in the second zone, and a raffmate or extract stream is collected from a column in the second zone and introduced to a nonadjacent column in the third zone.

9. A process according to any one of the preceding claims, wherein the first intermediate product obtained in the first separation step is enriched in the PUFA product compared to the feed mixture; and the second intermediate product obtained in the second separation step is enriched in the PUFA product compared to the first intermediate product.

10. A process according to any one of the previous claims, wherein in the first step the PUFA product is separated from components of the feed mixture which are less polar than the PUFA product, in the second step the PUFA product is separated from components of the feed mixture which are less polar than the PUFA product but more polar than the components separated in the first separation step, and in the third separation step the PUFA product is separated from more polar components of the feed mixture.

11. A process according to any one of the preceding claims, wherein the components separated from the PUFA product in the second separation step include DHA or a DHA derivative and/or other PUFAs or PUFA derivatives which are less polar than the PUFA product; and/or the components separated from the PUFA product in the third separation step include SDA or an SDA derivative and/or other PUFAs which are more polar than the PUFA product.

12. A process according to any one of the previous claims, wherein the second intermediate product is collected as the raffmate stream in the second separation step, and the PUFA product is collected as the extract stream in the third separation step.
13. A process according to claim 12, wherein the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, and wherein the first intermediate product is collected as the raffinate stream in the first separation step.

14. A process according to any one of the preceding claims, wherein the eluent is a mixture of water and an alcohol, an ether, an ester, a ketone or a nitrile.

15. A process according to claim 14, wherein the eluent is a mixture of water and methanol.

16. A process according to any one of the preceding claims, wherein the PUFA product contains EPA or an EPA derivative in an amount greater than 95 wt%, preferably 97 wt%.

17. A process according to any one of the preceding claims, wherein the EPA derivative is EPA ethyl ester (EE).

18. A process according to any one of the preceding claims, wherein
- part of the extract stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and/or
- part of the raffinate stream from the apparatus used in the second separation step is recycled back into the apparatus used in the second separation step; and/or
- part of the extract stream from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step; and/or
- part of the raffinate stream from the apparatus used in the third separation step is recycled back into the apparatus used in the third separation step.

19. A process according to claim 18, wherein the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, and wherein
part of the extract stream from the apparatus used in the first separation step
is recycled back into the apparatus used in the first separation step; and/or
part of the raffinate stream from the apparatus used in the first separation step
is recycled back into the apparatus used in the first separation step.

20. A process according to any one of the previous claims, wherein the
water:organic solvent ratio used in each separation step is adjusted such that
saturated and/or monounsaturated fatty acids present in the feed mixture are
removed in the first separation step; and
the PUFA product is separated from different components of the feed mixture
in steps (ii) and (iii).

21. A process according to any one of the previous claims, wherein the aqueous
organic solvent eluent used in each separation step has a different
water:organic solvent ratio.

22. A process according to any one of claims 1 to 20, wherein the aqueous organic
solvent eluent used in the second and third separation steps has the same
water:organic solvent ratio, and the aqueous organic solvent eluent used in the
first separation step has a different water:organic solvent ratio from the organic
solvent eluent used in the second and third separation steps.

23. A process according to claim 22, wherein the water:organic solvent ratio of the
aqueous organic solvent eluent used in the first separation step is lower than
the watenorganic solvent ratio of the aqueous organic solvent eluent used in
the second and third separation steps.

24. A process according to claim 23, wherein the watenorganic solvent ratio of the
eluent used in the first separation step is from 0.1:99.9 to 1:99 wt%, and the
water:organic solvent ratio of the eluent used in the second and third
separation steps is from 7:93 to 9:91 wt%.

25. A process according to any one of claims 18 to 24, wherein the rate at which
liquid collected via one or both of the extract and raffinate streams in the
second separation step is recycled back into the apparatus used in that separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii); and/or

wherein the rate at which liquid collected via one or both of the extract and raffinate streams in the third separation step is recycled back into the apparatus used in that separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

26. A process according to claim 22, wherein the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, and wherein the rate at which liquid collected via one or both of the extract and raffinate streams in the first separation step is recycled back into the apparatus used in that separation step is adjusted such that saturated and/or monounsaturated fatty acids present in the feed mixture are removed in the first separation step, and the PUFA product is separated from different components of the feed mixture in steps (ii) and (iii).

27. A process according to any one of claims 18 to 26, wherein the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatography apparatus used in the second separation step differs from the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatography apparatus used in the third separation step.

28. A process according to any one of claims 18 to 27, wherein the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus, and wherein the rate at which liquid collected via the extract stream in the first separation step is recycled back into the chromatography apparatus used in the first separation step differs from the rate at which liquid collected via the extract stream in the second separation step is
recycled back into the chromatography apparatus used in the second separation step.

29. A process according to any one of claims 18 to 28, wherein the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

30. A process according to claim 15, wherein:
   - the first separation step comprises purifying the feed mixture in a simulated or actual moving bed chromatography apparatus;
   - the second and third separation steps are carried out simultaneously in a single simulated or actual moving bed chromatography apparatus having a plurality of linked chromatography columns containing, as eluent, an aqueous organic solvent, the second and third separation steps being carried out in first and second zones respectively, wherein each zone is as defined in claim 3, and wherein the first separation step is carried out in a separate simulated or actual moving bed chromatography apparatus;
   - the first intermediate product is collected as the raffmate stream in the first separation step, the second intermediate product is collected as the raffmate stream in the second separation step, and the PUFA product is collected as the extract stream in the third separation step;
   - the second intermediate product raffmate stream containing the PUFA product together with more polar components is collected from a column in the first zone and introduced into a nonadjacent column in the second zone;
   - the aqueous organic solvent eluent used in the second and third separation steps has the same water:organic solvent ratio, and the water:organic solvent ratio of the eluent used in the first separation step is lower than the water:organic solvent ratio of the eluent used in the second and third separation steps; and
- the rate at which liquid collected via the extract stream in the second separation step is recycled back into the chromatographic apparatus used in the second separation step is faster than the rate at which liquid collected via the extract stream in the third separation step is recycled back into the chromatographic apparatus used in the third separation step.

31. A computer program for controlling a chromatography apparatus as defined in any one of the preceding claims, which computer program contains code means that, when executed, instructs the apparatus to carry out a process according to any one of the preceding claims.
Figure 2

2A

Feed mixture

FIRST SEPARATION STEP  
SECOND SEPARATION STEP  
THIRD SEPARATION STEP

1st intermediate product  
2nd intermediate product  
PUFA product

2B

Feed mixture

FIRST SEPARATION STEP  
SECOND SEPARATION STEP  
THIRD SEPARATION STEP

1st intermediate product  
2nd intermediate product  
PUFA product

2C

Feed mixture

FIRST SEPARATION STEP  
SECOND SEPARATION STEP  
THIRD SEPARATION STEP

1st intermediate product  
2nd intermediate product  
PUFA product

2D

Feed mixture

FIRST SEPARATION STEP  
SECOND SEPARATION STEP  
THIRD SEPARATION STEP

1st intermediate product  
2nd intermediate product  
PUFA product
Figure 3

FIRST SEPARATION STEP

F (A' + A + B + C)

E1 (A')

R1 (A + B + C)

FLOW OF ELUENT

FLOW OF ADSORBENT

SECOND SEPARATION STEP

ZONE 1

R1 (A + B + C)

E2 (A)

R2 (B + C)

THIRD SEPARATION STEP

ZONE 2

R2 (B + C)

E3 (B)

R3 (C)

FLOW OF ELUENT

FLOW OF ADSORBENT
Figure 4

FLOW OF ELUENT

FLOW OF ADSORBENT

FLOW OF ELUENT

FLOW OF ADSORBENT
Figure 8

EPA 85 EE intermediate from first separation step
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

INV. B01D15/18 C11B3/10 C11C1/00 C11C1/08

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)

B01D C11B C11C

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

Electronic database consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
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<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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</table>

Further documents are listed in the continuation of Box C.

Date of the actual completion of the international search

19 September 2012

Name and mailing address of the ISA/

European Patent Office, P.B. 5818 Patentlaan 2

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Fax: +31-70) 340-3016

Fourgeaud, Damien

Date of mailing of the international search report

27/09/2012

Authorized officer

Form PCT/ISA/210 (second sheet) (April 2005)
INTERNATIONAL SEARCH REPORT

Box No. II  Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. X Claims Nos.: 31 because they relate to subject matter not required to be searched by this Authority, namely:

   see FURTHER INFORMATION sheet PCT/ISA/210

2. ☐ Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. ☐ Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. III  Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

1. ☐ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. ☐ As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.

3. ☐ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:

4. ☒ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

☒ 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.

Form PCT/ISA/210 (continuation of first sheet (2)) (April 2005)
Claim 31 is directed to a computer program that should control the chromatographic process of claims 1 to 30. Such a program fails, however, under the exceptions listed in Rule 39.1 PCT, see point (vi). Consequently, in accordance with Article 17(2)(a)(i) PCT, no international search report is established for the subject-matter of this claim.
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