(12)





(11) **EP 1 576 166 B1**

EUROPEAN PATENT SPECIFICATION

- (45) Date of publication and mention of the grant of the patent:15.10.2014 Bulletin 2014/42
- (21) Application number: 03813566.1
- (22) Date of filing: 11.12.2003

(51) Int Cl.: C12N 15/82^(2006.01) C12N 9/02^(2006.01)

C12P 7/64 ^(2006.01) C12N 9/10 ^(2006.01)

- (86) International application number: PCT/EP2003/014054
- (87) International publication number: WO 2004/057001 (08.07.2004 Gazette 2004/28)

(54) NOVEL METHOD FOR THE PRODUCTION OF POLYUNSATURATED FATTY ACIDS

VERFAHREN ZUR HERSTELLUNG VON MEHRFACH UNGESÄTTIGTEN FETTSÄUREN NOUVEAU PROCEDE DE PRODUCTION D'ACIDES GRAS POLYINSATURES

- (84) Designated Contracting States:
 AT BE BG CH CY CZ DE DK EE ES FI FR GB GR
 HU IE IT LI LU MC NL PT RO SE SI SK TR
- (30) Priority: **19.12.2002 GB 0229578 21.07.2003 GB 0316989**
- (43) Date of publication of application: 21.09.2005 Bulletin 2005/38
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- (56) References cited: WO-A-00/34439 WO-A-02/057464 WO-A-02/077213 US-A1- 2002 138 874 US-B1- 7 256 033
 - WALLIS & BROWSE: "The delta-8-desaturase of Euglena gracilis: An alternate pathway for synthesis of 20-carbon polyunsaturated fatty acis", ARCHIVES OF BIOCHEMISTRY AND BIOPHYSICS, vol. 365, 1999, pages 307-316,

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Description

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[0001] The present invention relates to an improved process for the specific production of poly-unsaturated ω -3 and ω -6 fatty acids and a process for the production of triglycerides having an increased content of unsaturated fatty acids, in particular ω -3 and ω -6 fatty acids having at least two double bonds and a 20 or 22 carbon atom chain length. The invention relates to the production of a transgenic oil producing a plant, having an increased content of fatty acids, oils or lipids containing C₂₀- or C₂₂- fatty acids with a Δ 5, 7, 8, 10 double bond, respectively due to the expression of a Δ 8- desaturase and a Δ 9- elongase from organisms such as plants preferably Algae like Isochrysis galbana or Euglena

- gracilis. In addition the invention relates to a process for the production of poly unsaturated fatty acids such as Eicos apentaenoic, Arachidonic, Docosapentaenoic or Docosahexaenoic acid through the co- expression of a Δ-8-desaturase, a Δ-9-elongase and a Δ-5 desaturase in organisms such as plants.
 [0002] The invention additionally relates to the use of specific nucleic acid sequences encoding for the aforementioned proteins with Δ-8-desaturase-, Δ-9-elongase- or Δ-5-desaturase-activity, nucleic acid constructs, vectors and organisms containing said nucleic acid sequences. Also disclosed are unsaturated fatty acids and triglycerides having an increased
- ¹⁵ content of at least 1 % by weight of unsaturated fatty acids and use thereof. [0003] Fatty acids and triglycerides have numerous applications in the food industry, animal nutrition, cosmetics and in the drug sector. Depending on whether they are free saturated or unsaturated fatty acids or triglycerides with an increased content of saturated or unsaturated fatty acids, they are suitable for the most varied applications; thus, for example, <u>polyunsaturated fatty acids</u> (= PUFAs) are added to infant formula to increase its nutritional value. The various
- fatty acids and triglycerides are mainly obtained from microorganisms such as Mortierella or from oil-producing plants such as soybean, oilseed rape, sunflower and others, where they are usually obtained in the form of their triacylglycerides. Alternatively, they are obtained advantageously from animals, such as fish. The free fatty acids are prepared advantageously by hydrolysis.
- [0004] Whether oils with unsaturated or with saturated fatty acids are preferred depends on the intended purpose; thus, for example, lipids with unsaturated fatty acids, specifically polyunsaturated fatty acids, are preferred in human nutrition since they have a positive effect on the cholesterol level in the blood and thus on the possibility of heart disease. They are used in a variety of dietetic foodstuffs or medicaments. In addition PUFAs are commonly used in food, feed and in the cosmetic industry. Poly unsaturated ω-3-and/or ω-6-fatty acids are an important part of animal feed and human food. Because of the common composition of human food poly unsaturated ω-3-fatty acids, which are an essential
- ³⁰ component of fish oil, should be added to the food to increase the nutritional value of the food; thus, for example, poly unsaturated fatty acids such as Docosahexaenoic acid (= DHA, $C_{22:6}^{\Delta4,7,10,13,16,19}$) or Eicosapentaenoic acid (= EPA, $C_{20:5}^{\Delta5,8,11,14,17}$) are added as mentioned above to infant formula to increase its nutritional value. Whereas DHA has a positive effect of the brain development of babies. The addition of poly unsaturated ω -3-fatty acids is preferred as the addition of poly unsaturated ω -6-fatty acids like Arachidonic acid (= ARA, $C_{20:4}^{\Delta5,8,11,14}$) to common food have an
- ³⁵ undesired effect for example on rheumatic diseases such as rheumatoid arthritis. Poly unsaturated w-3- and ω-6-fatty acids are precursor of a family of paracrine hormones called eicosanoids such as prostaglandins which are products of the metabolism of Dihomo-γ-linoleic acid, ARA or EPA. Eicosanoids are involved in the regulation of lipolysis, the initiation of inflammatory responses, the regulation of blood circulation and pressure and other central functions of the body. Eicosanoids comprise prostaglandins, leukotrienes, thromboxanes, and prostacyclins. ω-3-fatty acids seem to prevent
- artherosclerosis and cardiovascular diseases primarily by regulating the levels of different eicosanoids. Other Eicosanoids are the thromboxanes and leukotrienes which are products of the metabolism of ARA or EPA.
 [0005] Principally microorganisms such as Mortierella or oil producing plants such as soybean, rapeseed or sunflower or algae such as Crytocodinium or Phaeodactylum are a common source for oils containing PUFAs, where they are usually obtained in the form of their triacyl glycerides. Alternatively, they are obtained advantageously from animals,
- ⁴⁵ such as fish. The free fatty acids are prepared advantageously by hydrolysis with a strong base such as potassium or sodium hydroxide. Higher poly unsaturated fatty acids such as DHA, EPA, ARA, Dihomo- γ -linoleic acid (C_{20:3}^{Δ8,11,14}) or Docosapentaenoic acid (= DPA, C_{22:5}^{Δ7,10,13,16,19}) are not produced by oil producing plants such as soybean, rapeseed, safflower or sunflower. A natural sources for said fatty acids are fish for example herring, salmon, sardine, redfish, eel, carp, trout, halibut, mackerel, pike-perch or tuna or algae.
- 50 [0006] On account of their positive properties there has been no shortage of attempts in the past to make available genes which participate in the synthesis of fatty acids or triglycerides for the production of oils in various organisms having a modified content of unsaturated fatty acids. Thus, in WO 91/13972 and its US equivalent a Δ-9-desaturase is described. In WO 93/11245 a Δ-15-desaturase and in WO 94/11516 a Δ-12-desaturase is claimed. WO 00/34439 discloses a Δ-5- and a Δ-8-desaturase. Other desaturases are described, for example, in EP-A-0 550162, WO 94/18337,
- ⁵⁵ WO 97/30582, WO 97/21340, WO 95/18222, EP-A-0 794 250, Stukey et al., J. Biol. Chem., 265,1990: 20144-20149, Wada et al., Nature 347, 1990: 200-203 or Huang et al., Lipids 34, 1999: 649-659. To date, however, the various desaturases have been only inadequately characterized biochemically since the enzymes in the form of membrane-bound proteins are isolable and characterizable only with very great difficulty (McKeon et al., Methods in Enzymol. 71, 100).

1981: 12141-12147, Wang et al., Plant Physiol. Biochem., 26,1988: 777-792). Generally, membrane-bound desaturases are characterized by introduction into a suitable organism which is then investigated for enzyme activity by means of analysis of starting materials and products. Δ -6-Desaturases are described in WO 93/06712, US 5,614,393, US 5614393, WO 96/21022, WO0021557 and WO 99/27111 and their application to production in transgenic organisms is also

⁵ described, e.g. in WO 9846763, WO 9846764 and WO 9846765. At the same time the expression of various fatty acid biosynthesis genes, as in WO 9964616 or WO 9846776, and the formation of poly-unsaturated fatty acids is also described and claimed.

[0007] From US2002/0138874 A1 the identification and uses of several genes involved in the elongation of monounsaturated fatty acids, especially a functionally active elongase, which utilize a monounsaturated fatty acid as a substrate

- is known. WO 02/077213 A3 describes an elongase gene encoding a polypeptide which elongates α-linolenic acid by at least two carbon atoms whereas γ-linolenic acid is not elongated.
 [0008] With regard to the effectiveness of the expression of desaturases and their effect on the formation of polyun-saturated fatty acids it may be noted that through expression of desaturases and elongases as described to date only low contents of poly-unsaturated fatty acids/lipids, such as by way of example eicosapentaenoic or archidonic acid, have
- ¹⁵ been achieved. Therefore, an alternative and more effective pathway with higher product yield is desirable. [0009] Accordingly, ther is still a great demand for new and more suitable genes which encode enzymes which participate in the biosynthesis of unsaturated fatty acids and make it possible to produce certain fatty acids specifically on an industrial scale without unwanted byproducts forming. In the selection of genes for biosynthesis two characteristics above all are particularly important. On the one hand, there is as ever a need for improved processes for obtaining the
- ²⁰ highest possible contents of polyunsaturated fatty acids. [0010] Accordingly, it is an object of the present invention to provide further genes of desaturase and elongase enzymes for the synthesis of polyunsaturated fatty acids in organisms preferably in microorganisms and plants and to use them in a commercial process for the production of poly unsaturated fatty acids. Said process should increase PUFA content in organisms as much as possible preferably in seeds of an oil producing plant.
- ²⁵ **[0011]** We have found that this object is achieved by a process for the production of compounds of the following general formula

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in transgenic oil producing plant with a content of at least 1 % by weight of said compounds referred to the total lipid content of said oil producing plant which comprises the following steps:

a) introduction of at least one nucleic acid sequence in a transgenic oil producing plant, which encodes a ∆-9elongase having a nucleic acid sequence depicted in SEQ ID NO:3, and

b) introduction of at least one second nucleic acid sequence which encodes a Δ -8-desaturase having a nucleic acid sequence depicted in SEQ ID NO:1, and

c) if necessary introduction of at least a third nucleic acid sequence, which encodes a ∆-5-desaturase having a nucleic acid sequence depicted in SEQ ID NO:5, 7 or 9, and

d) cultivating and harvesting of said oil producing plant; and where the variables and substituents in formula I have the following meanings:

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- R¹ = hydroxyl-, Coenzyme A-(Thioester), phosphatidylcholine-, phosphatidylethanolamine-, phosphatidylglycerol-, diphosphatidylglycerol-, phosphatidylserine-, phosphatidylinositol-, sphingolipid-, glycoshingolipid- or a residue of the general formula II:

$$H_{2}C-O-R^{2}$$

 $HC-O-R^{3}$ (II)
 $H_{2}C-O-f--$

where the substituents in formula II have the following meanings:

 $R^{2} = \frac{10}{10} R^{2} = \frac{10}{10} R^{2} = \frac{10}{10} R^{2} = \frac{10}{10} R^{2} R^{2$

 R^3 = hydrogen-, saturated or unsaturated C_2 - C_{24} -alkylcarbonyl-, or

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 R^2 and R^3 independent of each other a residue of the formula la:

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n =

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3,4 or 6, m = 3, 4 or 5 and p = 0 or 3, preferably n = 3, m = 4 or 5 and p = 0 or 3.

R¹ indicates in the formula I hydroxyl-, Acetyl-Coenzyme A-, phosphatidylcholine-, phosphatidylethanolamine-, phosphatidylglycerol-, diphosphatidylglycerol-, phosphatidylserine-, phosphatidylinositol-, sphingolipid-, glycoshingolipid- or a residue of the general formula II

 $H_{2}C-O-R^{2}$ $HC-O-R^{3}$ $H_{2}C-O-f^{-1}$ (II)

[0012] The abovementioned residues for R¹ are always coupled to compounds of the general formula I in the form of their ester or thioester.

[0013] R^2 indicates in structures of the general formula II hydrogen, phosphatidylcholine-, phosphatidylethanolamine-, phosphatidylglycerol-, diphosphatidylglycerol-, phosphatidylserine-, phosphatidylinositol-, shingolipid-, glycoshingolipid-, glycoshingolipid- or saturated or unsaturated C_2 - C_{24} -alkylcarbonyl-residues,

- [0014] Alkyl radicals which may be mentioned are substituted or unsubstituted, saturated or unsaturated C₂-C₂₄-alkylcarbonyl- chains such as ethylcarbonyl-, n-propylcarbonyl-, n-butylcarbonyl-, n-pentylcarbonyl-, n-hexylcarbonyl-, nheptylcarbonyl-, n-octyl-carbonyl-, n-nonylcarbonyl-, n-decylcarbonyl-, n-undecylcarbonyl-, n-dodecylcarbonyl-, n-tridecylcarbonyl-, n-tetradecylcarbonyl-, n-pentadecylcarbonyl-, n-hexadecylcarbonyl-, n-heptadecylcarbonyl-, n-octadecylcarbonyl-, n-nonadecylcarbonyl-, n-eicosylcarbonyl-, n-docosanylcarbonyl- or n-tetracosanylcarbonyl-, that contain one or more double bonds. Saturated or unsaturated C₁₀-C₂₂-Alkylcarbonylresidues such as n-decylcarbonyl-, n-undecyl-
- 50 carbonyl-, n-dodecylcarbonyl-, n-tridecylcarbonyl-, n-tetra-decylcarbonyl-, n-pentadecylcarbonyl-, n-hexadecylcarbonyl-, n-heptadecylcarbonyl-, n-octadecylcarbonyl-, n-nonadecylcarbonyl-, n-eicosylcarbonyl-, n-docosanyl-carbonyl- or n-tetracosanylcarbonyl- are preferred, which contain one ore more double bonds. In particular privileged are saturated or unsaturated C₁₀-C₂₂-alkylcarbonyl-residue as C₁₀-alkylcarbonyl-, C₁₁-alkylcarbonyl-, C₁₂-alkylcarbonyl-, C₁₃-alkylcarbonyl-, C₁₄-alkylcarbonyl-, C₁₆-alkylcarbonyl-, C₁₈-alkylcarbonyl-, C₂₀-alkylcarbonyl-, C₂₂-alkylcarbonyl- or C₂₄-alkyl-
- ⁵⁵ carbonyl-residue, that contain one ore more double bonds. In particular privileged are saturated or unsaturated C₁₆-C₂₂-alkylcarbonylresidue as C₁₆-alkylcarbonyl-, C₁₈-alkylcarbonyl-, C₂₀-alkylcarbonyl- or C₂₂-alkylcarbonyl-residue, that contain one ore more double bonds. The residues contain in particular two, three, four or five double bonds. Particularly preferred are residues of 20 or 22 carbon atoms having up to five double bonds, preferably three, four or five

double bonds. All residues are derived from the mentioned corresponding fatty acids.

[0015] R³ indicates in structures of the general formula II hydrogen, saturated or unsaturated C₂-C₂₄-alkylcarbonyl.

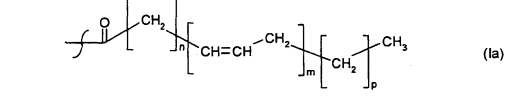
- [0016] Substituted or unsubstituted, saturated or unsaturated C₂-C₂₄-alkylcarbonyl- residues are e. g. ethylcarbonyl-, n-propylcarbonyl-, n-butylcarbonyl-, n-pentylcarbonyl-, n-hexyl-carbonyl-, n-hetylcarbonyl-, n-octylcarbonyl-, n-nonyl-
- ⁵ carbonyl-, n-decylcarbonyl-, n-undecylcarbonyl-, n-dodecylcarbonyl-, n-tridecylcarbonyl-, n-tetradecylcarbonyl-, n-pentadecylcarbonyl-, n-hexadecylcarbonyl-, n-heptadecylcarbonyl-, n-octadecylcarbonyl-, n-nonadecylcarbonyl-, n-eicosylcarbonyl-, n-docosanylcarbonyl- or n-tetracosanylcarbonyl-, having one or more double bonds. Preferred are saturated or unsaturated C₁₀-C₂₄-alkylcarbonyl residues as n-decylcarbonyl-, n-undecylcarbonyl-, n-dodecylcarbonyl-, n-tridecylcarbonyl-, n-tetradecylcarbonyl-, n-pentadecylcarbon yl-, n-hexadecylcarbonyl-, n-heptadecylcarbonyl-, n-octadecylcar-
- ¹⁰ bonyl-, n-nonadecylcarbonyl-, n-eicosylcarbonyl-, n-docosanylcarbonyl- or n-tetracosanylcarbonyl-, with one ore more double bonds. In particular saturated or unsaturated C_{10} - C_{24} -alkylcarbonyl residues as C_{10} -alkylcarbonyl-, C_{11} -alkyl-carbonyl-, C_{12} -alkylcarbonyl-, C_{13} -alkylcarbonyl-, C_{14} -alkylcarbonyl-, C_{16} -alkylcarbonyl-, C_{18} -alkylcarbonyl-, C_{20} -alkylcarbonyl-, C_{22} -alkylcarbonyl-, C_{22} -alkylcarbonyl-, C_{16} -alkylcarbonyl-, C_{18} -alkylcarbonyl-, C_{20} -alkylcarbonyl- or C_{24} -alkylcarbonyl-residues with one or more double bonds. In particular preferred are saturated or unsaturated C_{16} - C_{22} -alkylcarbonylresidue as C_{16} -alkylcarbonyl-, C_{18} -alkylcarbonyl-, C_{20} -alkylcarbonyl- or
- ¹⁵ C₂₂-alkylcarbonyl-residues, with multiple double bonds. C₁₈-alkylcarbonyl-residues are particularly preferred, which contain one, two, three or four double bonds and C₂₀-alkylcarbonyl-residues, with three, four or five double bonds. All residues are derived from the corresponding fatty acids.

[0017] R² and R³ indicates in structures of the general formula II independent of each other a residue of the general formula la



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whereas the variables in the formula I and Ia are defined as: n = 3,4 or 6, m = 3, 4 or 5 and p = 0 or 3. In particular: n = 3, m = 4 or 5 and p = 0 or 3.

[0018] The abovementioned residues R¹, R² and R³ can be substituted with hydoxyl- or epoxy-groups or might contain also triple bonds.

[0019] According to the invention the used nucleic acid sequences are isolated nucleic sequences coding for polypeptides having C₂₀- Δ 5- or Δ -8 desaturase or C₁₈- Δ 9-elongase activity.

- ³⁵ **[0020]** The according to inventive process synthesized substances of formula I which contain as residue R¹ the residue of formula II contain preferentially a mixture of different residues R² or R³. The residues are derived from different fatty acid molecules as short chain fatty acids with 4 to 6 C-atoms, mid-chain fatty acids having 8 to 12 C-atoms and longchain fatty acids with 14 to 24 C-atoms, whereas the long-chain fatty acids are preferred. Said long chain fatty acids are derived preferentially from C₁₈- or C₂₀-poly unsaturated fatty acids having advantageously between two and five double
- ⁴⁰ bonds. In addition the backbone of formula I is also derived from such a aforementioned fatty acid which advantageously is also different from R² and R³. That means compounds which are produced by the inventive process are in one aspect of the invention triglycerides of different substituted or unsubstituted, saturated or unsaturated fatty acid ester or thioesters.
 [0021] Also disclosed are poly-unsaturated fatty acid esters (of the formula I) with 18, 20 or 22 fatty acid carbon atoms chain length with at least two double bonds, preferably three, four or five are particularly preferred.
- In particular fatty acid molecules with three, four or five double bonds are preferred for the synthesis of eicosadienoic, eicosatrienoic, eicosatetranoic (arachidonic-acid) and eicosapentanoic acid (C20:2n-6, Δ11, 14; C20:3n-6, Δ8, 11, 14; C20:4n-6, Δ5, 8, 11, 14, C20:3n-3, Δ11, 14, 17; C20:4n-3, Δ8, 11, 14, 17; C20:5n-3, Δ5, 8, 11, 14, 17) in the inventive process, whereas arachidonic acid and eicosapentaenoic acid are most preferred. We have found that this object is advantageously achieved by the combined expression of three isolated nucleic acid sequences according to
- 50 the invention which encode for polypeptides having the following activities: a polypeptides with C20-Δ-8-desaturase activity, a C18-Δ-9-elongase activity, and a C20-Δ-5 desaturase activity. This objective was achieved in particular by the co-expression of the isolated nucleic acid sequences according to the invention. C18 fatty acids with a double bond in Δ-9-position are elongated by the Δ-9-elongase advantageously used in the inventive process. By the Δ-8-desaturase used in the process a double in Δ-8-position is introduced into C20 fatty acids. In addition a double bond can be introduced 55 into the fatty acid molecules in Δ-5-position by the Δ-5-desaturase.
- into the fatty acid molecules in Δ-5-position by the Δ-5-desaturase.
 [0023] The fatty acid ester of C₁₈-, C₂₀- and/or C₂₂-poly unsaturated fatty acids synthesized in the inventive process advantageously in form of their triglycerides as ester or thioesters can be isolated from the producing organism for example from a microorganism or a plant in the form of an oil, lipid or lipid mixture for example as sphingolipids, phos-

phoglycerides, lipids, glycolipids such as glycosphingolipids, phospholipids such as phosphatidylethanolamine, phosphatidylcholine, phosphatidylserine, phosphatidylglycerol, phosphatidylinositol or diphosphatidylglycerol, or as monoacylglyceride, diacylglyceride or triacylglyceride or as other fatty acid esters such as acetyl-Coenzym A thioester, which contain saturated or unsaturated fatty acids preferably poly unsaturated fatty acids with at least two preferably at least

⁵ three double bonds in the fatty acid molecule. In addition to the in form of the aforementioned esters bound fatty acids also fatty acids bound in other compounds can be produced or also free fatty acids can be produced by the inventive process.

[0024] In general the transgenic oil producing plants used in the inventive process contain fatty acid esters or fatty acids in a distribution of nearly 80 to 90 % by weight of triacyl glycerides, 2 to 5 % by weight diacyl glycerides, 5 to 10

- % by weight monoacyl glycerides, 1 to 5 % by weight free fatty acids and 2 to 8 % by weight phospholipids, whereas the total amount of the aforementioned compounds are all together a 100 % by weight.
 [0025] In the inventive process(es) [the singular shall include the plural and vice versa] at least 1 % by weight, preferably at least 2, 3, 4 or 5 % by weight, more preferably at least 6, 7, 8, or 9 % by weight, most preferably 10, 20 or 30 % by weight of the compounds of formula I referred to the total lipid content of the organism used in the process are produced.
- ¹⁵ Preferred starting material for the inventive process are linoleic acid (C18:2) and/or linolenic acid (C18:3) which are transformed to the preferred end products ARA or EPA. As for the inventive process organisms are used the product of the process is not a product of one pure substance per se. It is a mixture of different substances of formula I where one or more compounds are the major product and others are only contained as side products. In the event that in an organism used in the process linoleic and linolenic acid are available the end product is a mixture of ARA and EPA.
- Advantageously the side products shall not exceed 20 % by weight referred to the total lipid content of the organism, preferably the side products shall not exceed 15 % by weight, more preferably they shall not exceed 10 % by weight, most preferably they shall not exceed 5 % by weight. Preferably organisms are used in the process which contain as starting material either linoleic or linolenic acid so that as end product of the process only ARA or EPA are produced. In the event EPA and ARA are produced together, they should be produced in a ratio of at least 1:2 (EPA:ARA), preferably
- of at least 1:3, more preferably of at least 1:4, most preferably of at least 1:5. In the event that a mixture of different fatty acids such as ARA and EPA are the product of the inventive process said fatty acids can be further purified by method known by a person skilled in the art such as distillation, extraction, crystallization at low temperatures, chromatography or a combination of said methods.
- [0026] In principle all host organisms can be used in the inventive process for example transgenic organisms such as plants like mosses; green, red, brown or blue algae; monocotyledons or dicotyledones. Advantageously oil producing Transgenic algae, mosses or plants are used in the inventive processes described herein (for the invention the singular shall include the plural and vice versa), Additional advantageously organisms are plants or parts thereof. Plants are preferably used, very particularly preferably plants such as oilseed plants containing high amounts of lipid compounds such as rapeseed, poppy, mustard, hemp, castor bean, sesame, olive, calendula, punica, hazel nut, almond, macadamia,
- ³⁵ avocado, pumpkin, walnut, laurel, pistachio, primrose, canola, peanut, linseed, soybean, safflower, sunflower, borage or plants such as maize, wheat, rye, oat, triticale, rice, barley, cotton, manihot, pepper, tagetes, solanaceaous plants such as potato, tobacco, eggplant, and tomato, Vicia species, pea, alfalfa, bushy plants (coffee, cacao, tea), Salix species, trees (oil palm, coconut) and perennial grasses and forage crops. Particularly preferred plants of the invention are oilseed plants rapeseed, poppy, mustard, hemp, castor bean, sesame, olive, calendula, punica, hazel nut, almond,
- macadamia, avocado, pumpkin, laurel, pistachio, primrose, canola, peanut, linseed, soybean, safflower, sunflower, borage or trees (oil palm, coconut). Most preferred are C₁₈₋₂- and/or C_{18:3}-fatty acid rich plants such as hemp, sesame, linseed, poppy, pumpkin, walnut, tobacco, cotton, safflower or sunflower.
 [0027] Depending on the nucleic acid and/or the organism used in the inventive processes different compounds of
- the general formula I can be synthesized. In addition depending on the plant used in the process different mixtures of formula I compounds or single compounds such as arachidonic acid or eicosapentaenoic acid in free or bound form can be produced. In the event that in the inventive processes organism are used which have as precursor of the fatty acid synthesis preferably C_{18:2}- or C_{18:3}-fatty acids different poly unsaturated fatty acids can be synthesized for example starting from C_{18:2}-fatty acids γ-linoleic acid, dihomo-γ-linoleic acid or arachidonic acid can be produced. By influencing the
- ⁵⁰ activity of the different genes or their gene products different single compounds or compound mixtures can be produced. As living organisms are used in the inventive process the crude material that means crude lipids and/or oils isolated from the organisms preferably contain at least some starting compounds such as C_{18:2}- or C_{18:3}-fatty acids or their combination in the product and depending on the activity of the nucleic acid sequences and their gene products fatty acid intermediates of the biosynthesis chain. Said starting compounds or intermediates are in the product in a concen-
- ⁵⁵ tration of less than 20 or 15 % by weight, preferably less than 10, 9, 8, 7 or 6 % by weight, more preferably less than 5, 4, 3, 2 or 1 % by weight of the total fatty acids isolated from the used organism. **100291** Transport of the total fatty acids isolated as meaning single plant cells and their cultures on solid media or in

[0028] Transgenic plants are to be understood as meaning single plant cells and their cultures on solid media or in liquid culture, parts of plants and entire plants such as plant cell cultures, protoplasts from plants, callus cultures or plant

tissues such as leafs, shoots, seeds, flowers, roots etc. Said transgenic plants can be cultivated for example on solid or liquid culture medium, in soil or in hydroponics.

[0029] After cultivation transgenic plants which are used in the inventive process can be brought to the market without isolating compounds of the general formula I. Preferably the compounds of the general formula I are isolated from the

- ⁵ organisms in the form of their free fatty acids, their lipids or oils. The purification can be done by conventional methods such as squeezing and extraction of the plants or other methods instead of the extraction such as distillation, crystallization at low temperatures, chromatography or a combination of said methods. Advantageously the plants are grinded, heated and/or vaporized before the squeezing and extraction procedure. As solvent for the extraction solvents such as hexane are used. The isolated oils are further purified by acidification with for example phosphoric acid. The free fatty acids are
- ¹⁰ produced from said oils or lipids by hydrolysis. Charcoal or diatom earth are used to remove dyes from the fluid. In another preferred embodiment of the inventive process the alkyl ester of the fatty acids are produced from the oils and lipids by transesterification with an enzyme of with conventional chemistry. A preferred method is the production of the alkyl ester in the presence of alcohalates of the corresponding lower alcohols (C1 to C10 alcohols such as methanol, ethanol, propanol, butanol, hexanol etc.) such as methanolate or ethanolate. Therefore as the skilled worker knows the
- ¹⁵ alcohol in the presence of a catalytic amount of a base such as NaOH or KOH is added to the oils or lipids. [0030] It is also disclosed that the lipids can be obtained in the usual manner after the organisms have been grown. To this end, the organisms can first be harvested and then disrupted, or they can be used directly. It is advantageous to extract the lipids with suitable solvents such as apolar solvents, for example hexane, or polar solvents, for example ethanol, isopropanol, or mixtures such as hexane/isopropanol, phenol/chloroform/isoamyl alcohol, at temperatures be-
- 20 tween 0°C and 80°C, preferably between 20°C and 50°C. As a rule, the biomass is extracted with an excess of solvent, for example with an excess of solvent to biomass of 1:4. The solvent is subsequently removed, for example by distillation. The extraction may also be carried out with supercritical CO₂. After the extraction, the remainder of the biomass can be removed, for example, by filtration. Standard methods for the extraction of fatty acids from plants and microorganisms are described in Bligh et al. (Can. J. Biochem. Physiol. 37, 1959: 911-917) or Vick et al. (Plant Physiol. 69, 1092:1102, 1109)
- ²⁵ 1982:1103-1108).

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[0031] The crude oil thus obtained can then be purified further, for example by removing cloudiness by adding polar solvents such as acetone or apolar solvents such as chloroform, followed by filtration or centrifugation. Further purification via columns or other techniques is also possible.

[0032] To obtain the free fatty acids from the triglycerides, the latter are hyrolyzed in the customary manner, for example using NaOH or KOH.

[0033] In the process oils, lipids and/or free fatty acids or fractions thereof are produced. Said products can be used for the production of feed and food products, cosmetics or pharmaceuticals.

[0034] In principle all nucleic acids encoding polypeptides with Δ -8-desaturase, Δ -9-elongase and/or Δ -5-desaturase activity can be used in the inventive process. Preferably the nucleic acid sequences can be isolated for example from

- ³⁵ microorganism or plants such as fungi like Mortierella, algae like Euglena, Crypthecodinium or Isochrysis, diatoms like Phaeodactylum or mosses like Physcomitrella or Ceratodon, but also non-human animals such as Caenorhabditis are possible as source for the nucleic acid sequences. Advantageous nucleic acid sequences which encode polypeptides having a Δ-8-desaturase, Δ-9-elongase and/or Δ-5-desaturase activity are originate from microorganisms or plants, advantageously Phaeodactylum tricomutum, Ceratodon purpureus, Physcomitrella patens, Euglena gracilis or Isochrysis
- 40 galbana. Euglena gracilis or Isochrysis galbana are specific for the conversion of ω-3- or ω-6 fatty acids. Thus, the co expression of a Δ-9 elongase and a C20-specific Δ-8-desaturase leads to the formation of eicosatrienoic acid (C20:6n-3, Δ8, 11, 14) and eicosatetraenoic acid (C20:3n-4, Δ8, 11, 14, 17). Co-expression of a third gene coding for a C20-Δ5 specific desaturase leads to the production of Arachidonic acid (C20:6n-4, Δ5, 8, 11, 14) or Eicosapentaenoic acid (C20:3n-5, Δ5, 8, 11, 14, 17).
- ⁴⁵ [0035] The invention discloses the nucleotide sequence SEQ ID, No: 1 and the polypeptide sequence SEQ ID No: 2. Derivative(s) of the sequences according to the invention are for example, functional homologues of the polypeptides or enzymes encoded by SEQ ID NO: 2, but are not part of the invention. Also disclosed are derivatives of SEQ ID NO: 4, SEQ ID NO: 6, SEQ ID NO: 8 or SEQ ID NO: 10 which exhibit the same said specific enzymatic activity, but are not part of the invention. This specific enzymatic activity allows advantageously the synthesis of unsaturated fatty acids
- ⁵⁰ having more than three double bonds in the fatty acid molecule. By unsaturated fatty acids is meant in what follows diunsaturated or polyunsaturated fatty acids which possess double bonds. The double bonds may be conjugated or non conjugated. The said sequences encode enzymes which exhibit Δ-9 elongase, Δ-8-desaturase or -Δ5-desaturase activity.
 [0036] The enzymes, Δ-9 elongase, Δ-8-desaturase or Δ5-desaturase, advantageously either elongates fatty acid chains with 18 carbon atoms (see SEQ ID NO: 2) or introduces a double bond into fatty acid residues of glycerolipids,
- ⁵⁵ free fatty acids or acyl-CoA fatty acids at position C_8 - C_9 (see SEQ ID NO: 4) or at position C_5 - C_6 (see SEQ ID NO: 6, SEQ ID NO: 8 or SEQ ID NO: 10).

[0037] The nucleic acid sequence(s) according to the invention (for purposes of the application the singular encompasses the plural and vice versa) or fragments thereof may advantageously be used for isolating other genomic sequences

via homology screening.

[0038] Allele variants include in particular functional variants obtainable by deletion, insertion or substitution of nucleotides in the sequences depicted in SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9 the enzymatic activity of the derived synthesized proteins being retained.

- ⁵ **[0039]** Starting from the DNA sequence described in SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9 or parts of said sequences such DNA sequences can be isolated using, for example, normal hybridization methods or the PCR technique from other eukaryotes such as those identified above for example. These DNA sequences hybridize under standard conditions with the said sequences. For hybridization use is advantageously made of short oligonucleotides of the conserved regions of an average length of about 15 to 70 bp, preferably of about 17 to 60 bp,
- ¹⁰ more preferably of about 19 to 50 bp, most preferably of about 20 to 40 bp, for example, which can be determined by comparisons with other desaturase or elongase genes in the manner known to those skilled in the art. The histidine box sequences are advantageously employed. However, longer fragments of the nucleic acids according to the invention or the complete sequences may also be used for hybridization. Depending on the nucleic acid employed: oligonucleotide, longer fragment or complete sequence, or depending on which type of nucleic acid, DNA or RNA, is used for hybridization
- these standard conditions vary. Thus, for example, the melting temperatures of DNA:DNA hybrids are approximately 10 °C lower than those of DNA:RNA hybrids of the same length.
 [0040] By standard conditions is meant, for example, depending on the nucleic acid in question temperatures between 42 °C and 58 °C in an aqueous buffer solution having a concentration of between 0.1 and 5 x SSC (1 X SSC = 0.15 M
- NaCl, 15 mM sodium citrate, pH 7.2) or additionally in the presence of 50 % formamide, such as by way of example 42
 °C in 5 x SSC, 50 % formamide. Hybridization conditions for DNA:DNA hybrids are advantageously 0.1 x SSC and temperatures between approximately 20 °C and 45 °C, preferably between approximately 30 °C and 45 °C. For DNA:RNA hybrids the hybridization conditions are advantageously 0.1 x SSC and temperatures between approximately 30 °C and 55 °C, preferably between approximately 45 °C and 55 °C. These specified temperatures for hybridization are melting temperature values calculated by way of example for a nucleic acid having a length of approximately 100 nucleotides
- ²⁵ and a G + C content of 50 % in the absence of formamide. The experimental conditions for DNA hybridization are described in relevant genetics textbooks such as by way of example Sambrook et al., "Molecular Cloning", Cold Spring Harbor Laboratory, 1989, and may be calculated by formulae known to those skilled in the art, for example as a function of the length of the nucleic acids, the nature of the hybrids or the G + C content. Those skilled in the art may draw on the following textbooks for further information on hybridization: Ausubel et al. (eds), 1985, Current Protocols in Molecular
- ³⁰ Biology, John Wiley & Sons, New York; Hames and Higgins (eds), 1985, Nucleic Acids Hybridization: A Practical Approach, IRL Press at Oxford University Press, Oxford; Brown (ed), 1991, Essential Molecular Biology: A Practical Approach, IRL Press at Oxford University Press, Oxford. [0041] By homologues of the sequences SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7 and SEQ ID
- IDVAT By nomologues of the sequences SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7 and SEQ ID NO: 9 is meant derivatives such as by way of example promoter variants. These variants may be modified by one or
 more nucleotide exchanges, by insertion(s) and/or deletion(s) without, however, adversely affecting the functionality or efficiency of the promoters. Furthermore, the promoters can have their efficiency increased by altering their sequence
- or be completely replaced by more effective promoters even of foreign organisms. [0042] By derivatives is advantageously meant variants whose nucleotide sequence has been altered in the region
- from -1 to -2000 ahead of the start codon in such a way that the gene expression and/or the protein expression is modified, preferably increased. Furthermore, by derivatives is also meant variants which have been modified at the 3' end.
- **[0043]** The nucleic acid sequences according to the invention which encode a Δ -8-desaturase, a Δ -5-desaturase and/or a Δ -9-elongase may be produced by synthesis or obtained naturally or contain a mixture of synthetic and natural DNA components as well as consist of various heterologous Δ -8-desaturase, Δ -5-desaturase and/or Δ -9-elongase gene segments from different organisms. In general, synthetic nucleotide sequences are produced with codons which are preferred
- ⁴⁵ by the corresponding host organisms, plants for example. This usually results in optimum expression of the heterologous gene. These codons preferred by plants may be determined from codons having the highest protein frequency which are expressed in most of the plant species of interest. An example concerning Corynebacterium glutamicum is provided in Wada et al. (1992) Nucleic Acids Res. 20:2111-2118). Such experiments can be carried out using standard methods and are known to the person skilled in the art.
- 50 [0044] In addition, artificial DNA sequences are suitable, provided, as described above, they mediate the desired property, for example an increase in the content of Δ-8 and/or Δ-5 double bonds in fatty acids, oils or lipids in organisms such as in a plant by over-expression of the Δ-8-and/or Δ-5-desaturase gene in preferably in crop plants. Such artificial DNA sequences can exhibit Δ-8 and/or Δ-5-desaturase and/or Δ-9-elongase activity, for example by back-translation of proteins constructed by means of molecular modeling, or be determined by in vitro selection. Possible techniques for in
- vitro evolution of DNA to modify or improve the DNA sequences are described in Patten, P.A. et al., Current Opinion in Biotechnology 8, 724-733(1997) or in Moore, J.C. et al., Journal of Molecular Biology 272, 336-347 (1997). Particularly suitable are encoding DNA sequences which are obtained by back-translation of a polypeptide sequence in accordance with the codon use specific to the host plant. Those skilled in the art familiar with the methods of plant genetics can

easily determine the specific codon use by computer analyses of other known genes of the plant to be transformed. **[0045]** Other suitable equivalent nucleic acid sequences which may be mentioned are sequences that encode fusion proteins, a component of the fusion protein being a Δ -8-and/or a Δ -5-desaturase polypeptide and/or a Δ -9 elongase polypeptide or a functionally equivalent part thereof. The second part of the fusion protein can be, for example, another

- ⁵ polypeptide having enzymatic activity or an antigenic polypeptide sequence by means of which it is possible to demonstrate Δ -8- and/or Δ -5-desaturase or Δ -9-elongase expression (e.g. myc tag or his tag). Preferably, however, this is a regulatory protein sequence, such as by way of example a signal sequence for the endoplasmic reticulum (= ER) which directs the Δ -8- and/or Δ -5-desaturase protein and/or the Δ -9-elongase protein to the desired point of action, or regulatory sequences which influence the expression of the nucleic acid sequence according to the invention, such as promoters
- ¹⁰ or terminators. In another preferred embodiment the second part of the fusion protein is a plastidial targeting sequence as described by Napier J.A. [Targeting of foreign proteins to the chloroplast, Methods Mol. Biol., 49, 1995: 369 - 376]. A preferred used vector comprising said plastidial targeting sequence is disclosed by Colin Lazarus [Guerineau F., Woolston S., Brooks L., Mullineaux P. "An expression cassette for targeting foreign proteins into chloroplast; Nucleic. Acids Res., Dec 9, 16 (23), 1988: 11380].
- ¹⁵ [0046] Advantageously, the Δ-8-desaturase and Δ-9-elongase and/or the Δ-5-desaturase genes in the method according to the invention may be combined with other genes for fatty acid biosynthesis. Examples of such genes are the acyl transferases, other desaturases or elongases such as Δ-4-, Δ-5- or Δ-6-desaturases or w-3- and/or ω-6-specific desaturases such as Δ-12 (for C₁₈ fatty acids), Δ-15 (for C₁₈ fatty acids) or Δ-19 (for C₂₂ fatty acids) and/or such as Δ-5- or Δ-6-elongases. For in vivo and especially in vitro synthesis combination with e.g. NADH cytochrome B5 reductases
- which can take up or release reduction equivalents is advantageous.
 [0047] By the amino acid sequences according to the invention is meant proteins which contain an amino acid sequence depicted in the sequence SEQ ID NO: 2. Also disclosed are the sequences SEQ ID NO: 4, SEQ ID NO: 6, SEQ ID NO: 8 and SEQ ID NO: 10. For the estimation of an enzymatic activity which is "not substantially reduced" or which has the "same enzymatic activity" the enzymatic activity of the derived sequences are determined and compared with the wild
- 25 type enzyme activities. In doing this, for example, certain amino acids may be replaced by others having similar physicochemical properties (space filling, basicity, hydrophobicity, etc.). For example, arginine residues are exchanged for lysine residues, valine residues for isoleucine residues or aspartic acid residues for glutamic acid residues. However, one or more amino acids may also be swapped in sequence, added or removed, or a plurality of these measures may be combined with one another. The present invention also encompasses those nucleotide sequences which are obtained
- ³⁰ by modification of the Δ-8-desaturase nucleotide sequence, the Δ-5-desaturase nucleotide sequence and/or the Δ-9-elongase nucleotide sequence used in the inventive processes. The aim of such a modification may be, e.g., to further bound the encoding sequence contained therein or also, e.g., to insert further restriction enzyme interfaces.
 [0048] Functional equivalents also include those variants whose function by comparison as described above with the
- ³⁵ activity of the initial enzyme, that is activity is higher than 100 %, preferably higher than 110 %, particularly preferably higher than 130 %).

[0049] At the same time the nucleic acid sequence may, for example, advantageously be a DNA or cDNA sequence. Suitable encoding sequences for insertion into an expression cassette according to the invention include by way of example those which encode a Δ -8-desaturase, a Δ -5-desaturase and/or a Δ -9-elongase with the sequences described

above and lend the host the ability to overproduce fatty acids, oils or lipids having double bonds in the Δ-8-position and Δ-5-position, it being advantageous when at the same time fatty acids having at least four double bonds are produced. These sequences may be of homologous or heterologous origin.
 [0050] By the expression cassette (= nucleic acid construct or fragment or gene construct) according to the invention

is meant the sequences specified in SEQ ID NO: 1. Also disclosed are expression cassette with SEQ ID NO: 3, SEQ
 ⁴⁵ ID NO: 5, SEQ ID NO: 7 and/or SEQ ID NO: 9 which result from the genetic code and/or derivatives thereof which are functionally linked with one or more regulation signals advantageously to increase the gene expression and which control the expression of the encoding sequence in the host cell. These regulatory sequences should allow the selective expression of the genes and the protein expression. Depending on the host organism this may mean, for example, that

- the gene is expressed and/or overexpressed only after induction or that it is expressed and/or overexpressed immediately. Examples of these regulatory sequences are sequences to which inductors or repressors bind and in this way regulate the expression of the nucleic acid. In addition to these new regulation sequences or instead of these sequences the natural regulation of these sequences ahead of the actual structural genes may still be present and optionally have been genetically modified so that natural regulation was switched off and the expression of the genes increased. However, the gene construct can also be built up more simply, that is no additional regulation signals have been inserted ahead
- ⁵⁵ of the nucleic acid sequence or derivatives thereof and the natural promoter with its regulation has not been removed. Instead of this the natural regulation sequence was mutated in such a way that no further regulation ensues and/or the gene expression is heightened. These modified promoters in the form of part sequences (= promoter containing parts of the nucleic acid sequences according to the invention) can also be brought on their own ahead of the natural gene

to increase the activity. In addition, the gene construct may advantageously also contain one or more so-called enhancer sequences functionally linked to the promoter which allow enhanced expression of the nucleic acid sequence. At the 3' end of the DNA sequences additional advantageous sequences may also be inserted, such as further regulatory elements or terminators. The Δ -8- and/or Δ -5-desaturase gene and/or the Δ -9-elongase gene may be present in one or more

- ⁵ copies in the expression cassette (= gene construct). [0051] As described above, the regulatory sequences or factors can preferably positively influence and so increase the gene expression of the introduced genes. Thus, reinforcement of the regulatory elements advantageously on the transcription level may be effected by using powerful transcription signals such as promoters and/or enhancers. However, in addition reinforcement of translation is also possible, for example by improving the stability of the mRNA.
- 10 [0052] Suitable promoters in the expression cassette are in principle all promoters which can control the expression of foreign genes in organisms such as microorganisms like protozoa such as ciliates, algae such as green, brown, red or blue algae such as Euglenia, bacteria such as gram-positive or gram-negative bacteria, yeasts such as Saccharomyces, Pichia or Schizosaccharomyces or fungi such as Mortierella, Thraustochytrium or Schizochytrium or plants such as Aleuritia, advantageously in plants or fungi. Use is preferably made in particular of plant promoters or promoters
- ¹⁵ derived from a plant virus. Advantageous regulation sequences for the method according to the invention are found for example in promoters such as cos, tac, trp, tet, trp-tet, lpp, lac, lpp-lac, lacl^{q-,} T7, T5, T3, gal, trc, ara, SP6, λ-P_R or in λ-P_L promoters which are employed advantageously in gram-negative bacteria. Other advantageous regulation sequences are found, for example, in the gram-positive promoters amy and SPO2, in the yeast or fungal promoters ADC1, MFα, AC, P-60, CYC1, GAPDH, TEF, rp28, ADH or in the plant promoters CaMV/35S [Franck et al., Cell 21(1980)]
- ²⁰ 285-294], SSU, OCS, lib4, STLS1, B33, nos (= Nopalin Synthase Promoter) or in the ubiquintin or phaseolin promoter. The expression cassette may also contain a chemically inducible promoter by means of which the expression of the exogenous $\Delta 8$ - and/or Δ -5-desaturase gene and/or the Δ -9-elongase gene in the organisms can be controlled advantageously in the plants at a particular time. Advantageous plant promoters of this type are by way of example the PRP1 promoter [Ward et al., Plant. Mol. Biol.22(1993), 361-366], a promoter inducible by benzenesulfonamide (EP 388 186),
- ²⁵ a promoter inducible by tetracycline [Gatz et al., (1992) Plant J. 2,397-404], a promoter inducible by salicylic acid (WO 95/19443), a promoter inducible by abscisic acid (EP 335 528) and a promoter inducible by ethanol or cyclohexanone (WO93/21334). Other examples of plant promoters which can advantageously be used are the promoter of cytosolic FBPase from potato, the ST-LSI promoter from potato (Stockhaus et al., EMBO J. 8 (1989) 2445-245), the promoter of phosphoribosyl pyrophosphate amidotransferase from Glycine max (see also gene bank accession number U87999)
- 30 or a nodiene-specific promoter as described in EP 249 676. Particularly advantageous are those plant promoters which ensure expression in tissues or plant parts/organs in which fatty acid biosynthesis or the precursor stages thereof occurs, as in endosperm or in the developing embryo for example. Particularly noteworthy are advantageous promoters which ensure seed-specific expression such as by way of example the USP promoter or derivatives thereof, the LEB4 promoter, the phaseolin promoter or the napin promoter. The particularly advantageous USP promoter cited according to the
- ³⁵ invention or its derivatives mediate very early gene expression in seed development [Baeumlein et al., Mol Gen Genet, 1991, 225 (3): 459-67]. Other advantageous seed-specific promoters which may be used for monocotylodonous or dicotylodonous plants are the promoters suitable for dicotylodons such as napin gene promoters, likewise cited by way of example, from oilseed rape (US 5,608,152), the oleosin promoter from Arabidopsis (WO 98/45461), the phaseolin promoter from Phaseolus vulgaris (US 5,504,200), the Bce4 promoter from Brassica (WO 91/13980) or the leguminous
- ⁴⁰ B4 promoter (LeB4, Baeumlein et al., Plant J., 2, 2, 1992: 233-239) or promoters suitable for monocotylodons such as the promoters of the lpt2 or lpt1 gene in barley (WO 95/15389 and WO 95/23230) or the promoters of the barley hordeine gene, the rice glutelin gene, the rice oryzin gene, the rice prolamin gene, the wheat gliadin gene, the white glutelin gene, the corn zein gene, the oats glutelin gene, the sorghum kasirin gene or the rye secalin gene which are described in WO99/16890.
- ⁴⁵ [0053] Furthermore, particularly preferred are those promoters which ensure the expression in tissues or plant parts in which, for example, the biosynthesis of fatty acids, oils and lipids or the precursor stages thereof takes place. Particularly noteworthy are promoters which ensure a seed-specific expression. Noteworthy are the promoter of the napin gene from oilseed rape (US 5,608,152), the USP promoter from Vicia faba (USP = unknown seed protein, Baeumlein et al., Mol Gen Genet, 1991, 225 (3): 459-67), the promoter of the oleosin gene from Arabidopsis (WO98/45461), the phaseolin
- ⁵⁰ promoter (US 5,504,200) or the promoter of the legumin B4 gene (LeB4; Baeumlein et al., 1992, Plant Journal, 2 (2): 233-9). Other promoters to be mentioned are that of the lpt2 or lpt1 gene from barley (WO95/15389 and WO95/23230) which mediate seed-specific expression in monocotyledonous plants. Other advantageous seed specific promoters are promoters such as the promoters from rice, corn or wheat disclosed in WO 99/16890 or Amy32b, Amy6-6 or aleurain (US 5,677,474), Bce4 (rape, US 5,530,149), glycinin (soy bean, EP 571 741), phosphoenol pyruvat carboxyl-ase (soy
- ⁵⁵ bean, JP 06/62870), ADR12-2 (soy bean, WO 98/08962), isocitratlyase (rape, US 5,689,040) or β-amylase (barley, EP 781 849).

[0054] As described above, the expression construct (= gene construct, nucleic acid construct) may contain yet other genes which are to be introduced into the organisms. These genes can be subject to separate regulation or be subject

to the same regulation region as the Δ -8- and/or Δ -5-desaturase gene and/or the Δ -9-elongase gene. These genes are by way of example other biosynthesis genes, advantageously for fatty acid biosynthesis, which allow increased synthesis. Examples which may be mentioned are the genes form Δ -15- Δ -12-, Δ -9-, Δ -5-, Δ -4-desaturase, α -ketoacyl reductases, α -ketoacyl synthases, elongases or the various hydroxylases and acyl-ACP thioesterases. The desaturase genes are advantageously used in the nucleic acid construct.

- ⁵ advantageously used in the nucleic acid construct.
 [0055] In principle all natural promoters with their regulation sequences can be used like those named above for the expression cassette according to the invention and the method according to the invention. Over and above this, synthetic promoters may also advantageously be used.
- [0056] In the preparation of an expression cassette various DNA fragments can be manipulated in order to obtain a nucleotide sequence which usefully reads in the correct direction and is equipped with a correct reading raster. To connect the DNA fragments (= nucleic acids according to the invention) to one another adaptors or linkers may be attached to the fragments.

[0057] The promoter and the terminator regions can usefully be provided in the transcription direction with a linker or polylinker containing one or more restriction points for the insertion of this sequence. Generally, the linker has 1 to 10,

- ¹⁵ mostly 1 to 8, preferably 2 to 6, restriction points. In general the size of the linker inside the regulatory region is less than 100 bp, frequently less than 60 bp, but at least 5 bp. The promoter may be both native or homologous as well as foreign or heterologous to the host organism, for example to the host plant. In the 5'-3' transcription direction the expression cassette contains the promoter, a DNA sequence which encodes a Δ-8-desaturase gene, a Δ-5-desaturase gene and/or a Δ-9-elongase gene and a region for transcription termination. Different termination regions can be exchanged for one
- 20 another in any desired fashion.

[0058] Furthermore, manipulations which provide suitable restriction interfaces or which remove excess DNA or restriction interfaces can be employed. Where insertions, deletions or substitutions, such as transitions and transversions, come into consideration, *in vitro* mutagenesis, primer repair, restriction or ligation may be used. In suitable manipulations such as restriction, chewing back or filling of overhangs for blunt ends complementary ends of the fragments can be provided for the ligation

²⁵ provided for the ligation.

[0059] For an advantageous high expression the attachment of the specific ER retention signal SEKDEL inter alia can be of importance (Schouten, A. et al., Plant Mol. Biol. 30 (1996), 781-792). In this way the average expression level is tripled or even quadrupled. Other retention signals which occur naturally in plant and animal proteins located in the ER may also be employed for the construction of the cassette. In another preferred embodiment a plastidial targeting

³⁰ sequence is used as described by Napier J.A. [Targeting of foreign proteins to the chloroplast, Methods Mol. Biol., 49, 1995: 369-376]. A preferred used vector comprising said plastidial targeting sequence is disclosed by Colin Lazarus [Guerineau F., Woolston S., Brooks L., Mullineaux P. "An expression cassette for targeting foreign proteins into chloroplast; Nucleic. Acids Res., Dec 9, 16 (23), 1988: 11380].

[0060] Preferred polyadenylation signals are plant polyadenylation signals, preferably those which substantially correspond to T-DNA polyadenylation signals from Agrobacterium tumefaciens, in particular gene 3 of the T-DNA (octopin

- respond to T-DNA polyadenylation signals from Agrobacterium tumefaciens, in particular gene 3 of the T-DNA (octopin synthase) of the Ti plasmid pTiACH5 (Gielen et al., EMBO J.3 (1984), 835 et seq.) or corresponding functional equivalents.
 [0061] An expression cassette is produced by fusion of a suitable promoter with a suitable Δ-8- and/or Δ-5-desaturase DNA sequence and/or a suitable Δ-9-elongase DNA sequence together with a polyadenylation signal by common recombination and cloning techniques as described, for example, in T. Maniatis, E.F. Fritsch and J. Sambrook, Molecular
- 40 Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY (1989) as well as in T.J. Silhavy, M.L. Berman and L.W. Enquist, Experiments with Gene Fusions, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY (1984) and in Ausubel, F.M. et al., Current Protocols in Molecular Biology, Greene Publishing Assoc. and Wiley-Interscience (1987).

[0062] In the preparation of an expression cassette various DNA fragments can be manipulated to produce a nucleotide sequence which usefully reads in the correct direction and is equipped with a correct reading raster. Adapters or linkers can be attached to the fragments for joining the DNA fragments.
[0062] The preparation of the terminator regions can usefully be previded in the transcription direction with a linker or

[0063] The promoter and the terminator regions can usefully be provided in the transcription direction with a linker or polylinker containing one or more restriction points for the insertion of this sequence. Generally, the linker has 1 to 10, mostly 1 to 8, preferably 2 to 6, restriction points. In general the size of the linker inside the regulatory region is less than

- 50 100 bp, frequently less than 60 bp, but at least 5 bp. The promoter may be both native or homologous as well as foreign or heterologous to the host organism, for example to the host plant. In the 5'-3' transcription direction the expression cassette contains the promoter, a DNA sequence which either encodes a Δ-8- and/or Δ-5-desaturase gene and/or a Δ-9-elongase gene and a region for transcription termination. Different termination regions can be exchanged for one another in any desired fashion.
- ⁵⁵ **[0064]** In the preparation of an expression cassette various DNA fragments can be manipulated to produce a nucleotide sequence which usefully reads in the correct direction and is equipped with a correct reading raster. Adapters or linkers can be attached to the fragments for joining the DNA fragments.

[0065] The DNA sequences encoding the nucleic acid sequences used in the inventive processes such as the Δ -8-

desaturase from Euglenia gracilis, the Δ -9-elongase from Isochrysis galbana and/or the Δ -5-desaturase for example from Caenorhabditis elegans, Mortierella alpina, Borage officinalis or Physcomitrella patens contain all the sequence characteristics needed to achieve correct localization of the site of fatty acid, lipid or oil biosynthesis. Accordingly, no further targeting sequences are needed per se. However, such a localization may be desirable and advantageous and

⁵ hence artificially modified or reinforced so that such fusion constructs are also a preferred advantageous embodiment of the invention.

[0066] Particularly preferred are sequences which ensure targeting in plastids. Under certain circumstances targeting into other compartments (reported in: Kermode, Crit. Rev. Plant Sci. 15, 4 (1996), 285-423) may also be desirable, e.g. into vacuoles, the mitochondrium, the endoplasmic reticulum (ER), peroxisomes, lipid structures or due to lack of corresponding operative sequences retention in the compartment of origin, the cytosol.

[0067] Advantageously, the nucleic acid sequences according to the invention or the gene construct together with at least one reporter gene are cloned into an expression cassette which is introduced into the organism via a vector or directly into the genome. This reporter gene should allow easy detection via a growth, fluorescence, chemical, bioluminescence or resistance assay or via a photometric measurement. Examples of reporter genes which may be mentioned

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- ¹⁵ are antibiotic- or herbicide-resistance genes, hydrolase genes, fluorescence protein genes, bioluminescence genes, sugar or nucleotide metabolic genes or biosynthesis genes such as the Ura3 gene, the Ilv2 gene, the luciferase gene, the β-galactosidase gene, the gfp gene, the 2-desoxyglucose-6-phosphate phosphatase gene, the β-glucuronidase gene, β-lactamase gene, the neomycin phosphotransferase gene, the hygromycin phosphotransferase gene or the BASTA (= gluphosinate-resistance) gene. These genes permit easy measurement and quantification of the transcription
- 20 activity and hence of the expression of the genes. In this way genome positions may be identified which exhibit differing productivity.

[0068] In a preferred embodiment an expression cassette comprises upstream, i.e. at the 5' end of the encoding sequence, a promoter and downstream, i.e. at the 3' end, a polyadenylation signal and optionally other regulatory elements which are operably linked to the intervening encoding sequence for Δ -8-desaturase, Δ -9-elongase and/or Δ -

- 5-desaturase DNA sequence. By an operable linkage is meant the sequential arrangement of promoter, encoding sequence, terminator and optionally other regulatory elements in such a way that each of the regulatory elements can fulfill its function in the expression of the encoding sequence in due manner. The sequences preferred for operable linkage are targeting sequences for ensuring subcellular localization in plastids. However, targeting sequences for ensuring subcellular localization in the mitochondrium, in the endoplasmic reticulum (= ER), in the nucleus, in oil corpuscles or
- other compartments may also be employed as well as translation promoters such as the 5' lead sequence in tobacco mosaic virus (Gallie et al., Nucl. Acids Res. 15 (1987), 8693 -8711).
 [0069] An expression cassette may, for example, contain a constitutive promoter or a tissue-specific promoter (preferably the USP or napin promoter) the gene to be expressed and the ER retention signal. For the ER retention signal the KDEL amino acid sequence (lysine, aspartic acid, glutamic acid, leucine) or the KKX amino acid sequence (lysine).
- ³⁵ Iysine-X-stop, wherein X means every other known amino acid) is preferably employed. [0070] For expression in a prokaryotic or eukaryotic host organism, for example a microorganism such as a fungus or a plant the expression cassette is advantageously inserted into a vector such as by way of example a plasmid, a phage or other DNA which allows optimum expression of the genes in the host organism. Examples of suitable plasmids are: in E. coli pLG338, pACYC184, pBR series such as e.g. pBR322, pUC series such as pUC18 or pUC19, M113mp
- 40 series, pKC30, pRep4, pHS1, pHS2, pPLc236, pMBL24, pLG200, pUR290, pIN-III¹¹³-B1, λgt11 or pBdCl; in Streptomyces pIJ101, pIJ364, pIJ702 or pIJ361; in Bacillus pUB110, pC194 or pBD214; in Corynebacterium pSA77 or pAJ667; in fungi pALS1, pIL2 or pBB116; other advantageous fungal vectors are described by Romanos, M.A. et al., [(1992) "Foreign gene expression in yeast: a review", Yeast 8: 423-488] and by van den Hondel, C.A.M.J.J. et al. [(1991) "Heterologous gene expression in filamentous fungi" as well as in More Gene Manipulations in Fungi [J.W. Bennet & L.L. Lasure,
- ⁴⁵ eds., pp. 396-428: Academic Press: San Diego] and in "Gene transfer systems and vector development for filamentous fungi" [van den Hondel, C.A.M.J.J. & Punt, P.J. (1991) in: Applied Molecular Genetics of Fungi, Peberdy, J.F. et al., eds., pp. 1-28, Cambridge University Press: Cambridge]. Examples of advantageous yeast promoters are 2μM, pAG-1, YEp6, YEp13 or pEMBLYe23. Examples of algal or plant promoters are pLGV23, pGHlac⁺, pBIN19, pAK2004, pVKH or pDH51 (see Schmidt, R. and Willmitzer, L., 1988). The vectors identified above or derivatives of the vectors identified
- ⁵⁰ above are a small selection of the possible plasmids. Further plasmids are well known to those skilled in the art and may be found, for example, in the book Cloning Vectors (Eds. Pouwels P.H. et al. Elsevier, Amster-dam-New York-Oxford, 1985, ISBN 0 444 904018). Suitable plant vectors are described inter alia in "Methods in Plant Molecular Biology and Biotechnology" (CRC Press), Ch. 6/7, pp. 71-119. Advantageous vectors are known as shuttle vectors or binary vectors which replicate in E. coli and Agrobacterium.
- ⁵⁵ **[0071]** By vectors is meant with the exception of plasmids all other vectors known to those skilled in the art such as by way of example phages, viruses such as SV40, CMV, baculovirus, adenovirus, transposons, IS elements, phasmids, phagemids, cosmids, linear or circular DNA. These vectors can be replicated autonomously in the host organism or be chromosomally replicated, chromosomal replication being preferred.

[0072] In a further embodiment of the vector the expression cassette according to the invention may also advantageously be introduced into the organisms in the form of a linear DNA and be integrated into the genome of the host organism by way of heterologous or homologous recombination. This linear DNA may be composed of a linearized plasmid or only of the expression cassette as vector or the nucleic acid sequences according to the invention.

⁵ **[0073]** In a further advantageous embodiment the nucleic acid sequence according to the invention can also be introduced into an organism on its own.

[0074] If in addition to the nucleic acid sequence according to the invention further genes are to be introduced into the organism, all together with a reporter gene in a single vector or each single gene with a reporter gene in a vector in each case can be introduced into the organism, whereby the different vectors can be introduced simultaneously or successively.

¹⁰ **[0075]** The vector advantageously contains at least one copy of the nucleic acid sequences according to the invention and/or the expression cassette (= gene construct) according to the invention.

[0076] By way of example the plant expression cassette can be installed in the pRT transformation vector ((a) Toepfer et al., 1993, Methods Enzymol., 217: 66-78; (b) Toepfer et al. 1987, Nucl. Acids. Res. 15: 5890 ff.).

- [0077] Alternatively, a recombinant vector (= expression vector) can also be transcribed and translated in vitro, e.g. by using the T7 promoter and the T7 RNA polymerase.
- **[0078]** Expression vectors employed in prokaryotes frequently make use of inducible systems with and without fusion proteins or fusion oligopeptides, wherein these fusions can ensue in both N-terminal and C-terminal manner or in other useful domains of a protein. Such fusion vectors usually have the following purposes: i.) to increase the RNA expression rate; ii.) to increase the achievable protein synthesis rate; iii.) to increase the solubility of the protein; iv.) or to simplify
- ²⁰ purification by means of a binding sequence usable for affinity chromatography. Proteolytic cleavage points are also frequently introduced via fusion proteins which allows cleavage of a portion of the fusion protein and purification. Such recognition sequences for proteases are recognized, e.g. factor Xa, thrombin and enterokinase. [0079] Typical advantageous fusion and expression vectors are pGEX [Pharmacia Biotech Inc; Smith, D.B. and John-

[00/9] Typical advantageous fusion and expression vectors are pGEX [Pharmacia Biotech Inc; Smith, D.B. and Johnson, K.S. (1988) Gene 67: 31-40], pMAL (New England Biolabs, Beverly, MA) and pRIT5 (Pharmacia, Piscataway, NJ)
 which contains glutathione S-transferase (GST), maltose binding protein or protein A.

- **[0080]** Other examples of E. coli expression vectors are pTrc [Amann et al., (1988) Gene 69:301-315] and pET vectors [Studier et al., Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, California (1990) 60-89; Stratagene, Amsterdam, The Netherlands].
- [0081] Other advantageous vectors for use in yeast are pYepSec1 (Baldari, et al., (1987) Embo J. 6:229-234), pMFa
 ³⁰ (Kurjan and Herskowitz, (1982) Cell 30:933-943), pJRY88 (Schultz et al., (1987) Gene 54:113-123), and pYES derivatives
 (Invitrogen Corporation, San Diego, CA). Vectors for use in filamentous fungi are described in: van den Hondel, C.A.M.J.J.
 & Punt, P.J. (1991) "Gene transfer systems and vector development for filamentous fungi", in: Applied Molecular Genetics of Fungi, J.F. Peberdy, et al., eds., pp. 1-28, Cambridge University Press: Cambridge.
- [0082] Alternatively, insect cell expression vectors can also be advantageously utilized, e.g. for expression in Sf 9 cells. These are e.g. the vectors of the pAc series (Smith et al. (1983) Mol. Cell Biol. 3:2156-2165) and the pVL series (Lucklow and Summers (1989) Virology 170:31-39).

[0083] Furthermore, plant cells or algal cells can advantageously be used for gene expression. Examples of plant expression vectors may be found in Becker, D., et al. (1992) "New plant binary vectors with selectable markers located proximal to the left border", Plant Mol. Biol. 20: 1195-1197 or in Bevan, M.W. (1984) "Binary Agrobacterium vectors for plant transformation", Nucl. Acid. Res. 12: 8711-8721.

[0084] Furthermore, the nucleic acid sequences may also be expressed in mammalian cells, advantageously in nonhuman mammalian cells. Examples of corresponding expression vectors are pCDM8 and pMT2PC referred to in: Seed, B. (1987) Nature 329:840 or Kaufman et al. (1987) EMBO J. 6: 187-195). At the same time promoters preferred for use are of viral origin, such as by way of example promoters of polyoma, adenovirus 2, cytomegalovirus or simian virus 40.

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⁴⁵ Other prokaryotic and eukaryotic expression systems are referred to in chapters 16 and 17 of Sambrook et al., Molecular Cloning: A Laboratory Manual. 2nd, ed., Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989.

[0085] The host organism (= transgenic organism) advantageously contains at least one copy of the nucleic acid according to the invention and/or of the nucleic acid construct according to the invention.

⁵⁰ **[0086]** The introduction of the nucleic acids according to the invention, the expression cassette or the vector into organisms, plants for example, can in principle be done by all of the methods known to those skilled in the art. The introduction of the nucleic acid sequences gives rise to recombinant or transgenic organisms.

[0087] In the case of microorganisms, those skilled in the art can find appropriate methods in the textbooks by Sambrook, J. et al. (1989) Molecular cloning: A laboratory manual, Cold Spring Harbor Laboratory Press, by F.M. Ausubel et al. (1994) Current protocols in molecular biology, John Wiley and Sons, by D.M. Glover et al., DNA Cloning Vol.1, (1995), IRL Press (ISBN 019-963476-9), by Kaiser et al. (1994) Methods in Yeast Genetics, Cold Spring Harbor Laboratory

Press or Guthrie et al. Guide to Yeast Genetics and Molecular Biology, Methods in Enzymology, 1994, Academic Press. [0088] The transfer of foreign genes into the genome of a plant is called transformation. In doing this the methods

described for the transformation and regeneration of plants from plant tissues or plant cells are utilized for transient or stable transformation. Suitable methods are protoplast transformation by poly(ethylene glycol)-induced DNA uptake, the "biolistic" method using the gene cannon - referred to as the particle bombardment method, electroporation, the incubation of dry embryos in DNA solution, microinjection and gene transfer mediated by Agrobacterium. Said methods are described

- ⁵ by way of example in B. Jenes et al., Techniques for Gene Transfer, in: Transgenic Plants, Vol. 1, Engineering and Utilization, eds. S.D. Kung and R. Wu, Academic Press (1993) 128-143 and in Potrykus Annu. Rev. Plant Physiol. Plant Molec. Biol. 42 (1991) 205-225). The nucleic acids or the construct to be expressed is preferably cloned into a vector which is suitable for transforming Agrobacterium tumefaciens, for example pBin19 (Bevan et al., Nucl. Acids Res. 12 (1984) 8711). Agrobacteria transformed by such a vector can then be used in known manner for the transformation of
- ¹⁰ plants, in particular of crop plants such as by way of example tobacco plants, for example by bathing bruised leaves or chopped leaves in an agrobacterial solution and then culturing them in suitable media. The transformation of plants by means of Agrobacterium tumefaciens is described, for example, by Höfgen and Willmitzer in Nucl. Acid Res. (1988) 16, 9877 or is known inter alia from F.F. White, Vectors for Gene Transfer in Higher Plants; in Transgenic Plants, Vol. 1, Engineering and Utilization, eds. S.D. Kung and R. Wu, Academic Press, 1993, pp. 15-38.
- 15 [0089] Agrobacteria transformed by an expression vector according to the invention may likewise be used in known manner for the transformation of plants such as test plants like Arabidopsis or crop plants such as cereal crops, corn, oats, rye, barley, wheat, soybean, rice, cotton, sugar beet, canola, sunflower, flax, hemp, potatoes, tobacco, tomatoes, carrots, paprika, oilseed rape, tapioca, cassava, arrowroot, tagetes, alfalfa, lettuce and the various tree, nut and vine species, in particular of oil-containing crop plants such as soybean, peanut, castor oil plant, sunflower, com, cotton, flax,
- 20 oilseed rape, coconut, oil palm, safflower (Carthamus tinctorius) or cocoa bean, e.g. by bathing bruised leaves or chopped leaves in an agrobacterial solution and then culturing them in suitable media. For the production of PUFAs, for example stearidonic acid, eicosapentaenoic acid and docosahexaenoic acid, borage, linseed, sunflower, safflower or Primulaceae are advantageously suitable. Other suitable organisms for the production of for example γ-linoleic acid, dihomo-γ-linoleic acid or arachidonic acid are for example linseed, sunflower or safflower.
- ²⁵ **[0090]** The genetically modified plant cells may be regenerated by all of the methods known to those skilled in the art. Appropriate methods can be found in the publications referred to above by S.D. Kung and R. Wu, Potrykus or Höfgen and Willmitzer.

[0091] Accordingly, a further aspect of the invention relates to transgenic organisms transformed by at least one nucleic acid sequence, expression cassette or vector according to the invention as well as cells, cell cultures, tissue, parts -

- ³⁰ such as, for example, leaves, roots, etc. in the case of plant organisms or reproductive material derived from such organisms. The terms "host organism", "host cell", "recombinant (host) organism" and "transgenic (host) cell" are used here interchangeably. Of course these terms relate not only to the particular host organism or the particular target cell but also to the descendants or potential descendants of these organisms or cells. Since, due to mutation or environmental effects certain modifications may arise in successive generations, these descendants need not necessarily be identical with the parental cell but nevertheless are still encompassed by the term as used here.
- ³⁵ with the parental cell but nevertheless are still encompassed by the term as used here. [0092] For the purposes of the invention "transgenic" or "recombinant" means with regard for example to a nucleic acid sequence, an expression cassette (= gene construct, nucleic acid construct) or a vector containing the nucleic acid sequence according to the invention or an organism transformed by the nucleic acid sequences, expression cassette or vector according to the invention all those constructions produced by genetic engineering methods in which either
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a) the nucleic acid sequence depicted in SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7, SEQ ID NO: 9 or

b) a genetic control sequence functionally linked to the nucleic acid sequence described under (a), for example a 3'- and/or 5'- genetic control sequence such as a promoter or terminator, or

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c) (a) and (b)

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are not found in their natural, genetic environment or have been modified by genetic engineering methods, wherein the modification may by way of example be a substitution, addition, deletion, inversion or insertion of one or more nucleotide residues. Natural genetic environment means the natural genomic or chromosomal locus in the organism of origin or inside the host organism or presence in a genomic library. In the case of a genomic library the natural genetic environment of the nucleic acid sequence is preferably retained at least in part. The environment borders the nucleic acid sequence at least on one side and has a sequence length of at least 50 bp, preferably at least 500 bp, particularly preferably at least 1,000 bp, most particularly preferably at least 5,000 bp. A naturally occurring expression cassette - for example

⁵⁵ the naturally occurring combination of the natural promoter of the nucleic acid sequence according to the invention with the corresponding Δ -8-desaturase, Δ -9-elongase and/or Δ -5-desaturase gene - turns into a transgenic expression cassette when the latter is modified by unnatural, synthetic ("artificial") methods such as by way of example a mutagenation. Appropriate methods are described by way of example in US 5,565,350 or WO 00/15815.

[0093] Suitable organisms or host organisms for the nucleic acid, expression cassette or vector according to the invention are advantageously in principle all organisms which are able to synthesize fatty acids, especially unsaturated fatty acids or are suitable for the expression of recombinant genes as described above. Further examples which may be mentioned are plants such as Arabidopsis, Asteraceae such as Calendula or crop plants such as soybean, peanut,

- ⁵ castor oil plant, sunflower, corn, cotton, flax, oilseed rape, coconut, oil palm, safflower (Carthamus tinctorius) or cocoa bean. Preference is given to organisms which can naturally synthesize oils in relatively large quantities such as plants such as soybean, oilseed rape, coconut, oil palm, safflower, flax, castor oil plant, Calendula, peanut, cocoa bean or sunflower, and particular preference is given to soybean, flax, oilseed rape, sunflower or Calendula.
- [0094] Further useful host cells are identified in: Goeddel, Gene Expression Technology: Methods in Enzymology 185,
 ¹⁰ Academic Press, San Diego, CA (1990).

[0095] Usable expression strains, e.g. those exhibiting a relatively low protease activity, are described in: Gottesman, S., Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, California (1990) 119-128. **[0096]** A further object of the invention relates to the use of an expression cassette containing DNA sequences encoding a Δ -8-desaturase, a Δ -9-elongase and/or a Δ -5-desaturase gene or DNA sequences hybridizing therewith for the trans-

formation of plant cells, tissues or parts of plants. The aim of use is to increase the content of fatty acids, oils or lipids having an increased content of double bonds.

[0097] In doing so, depending on the choice of promoter, the Δ -8-desaturase, a Δ -9-elongase and/or a Δ -5-desaturase gene can be expressed specifically in the leaves, in the seeds, the nodules, in roots, in the stem or other parts of the plant. Those transgenic plants overproducing fatty acids, oils or lipids having at least three double bonds in the fatty acid

20 molecule, the reproductive material thereof, together with the plant cells, tissues or parts thereof are a further object of the present invention.

[0098] The expression cassette or the nucleic acid sequences according to the invention containing a Δ -8-desaturase, a Δ -9-elongase and/or a Δ -5-desaturase gene sequence can, moreover, also be employed for the transformation of the organisms identified by way of example above such as bacteria, cyanobacteria, yeasts, filamentous fungi, ciliates and algae with the objective of increasing the content of fatty acide, either a link processing at least three double bonds.

- algae with the objective of increasing the content of fatty acids, oils or lipids possessing at least three double bonds.
 [0099] Within the framework of the present invention, increasing the content of fatty acids, oils or lipids possessing at least three double bonds means, for example, the artificially acquired trait of increased biosynthetic performance due to functional overexpression of the Δ-8-desaturase, Δ-9-elongase and/or Δ-5-desaturase gene in the organisms according to the invention, advantageously in the transgenic plants according to the invention, by comparison with the nongenetically modified initial plants at least for the duration of at least one plant generation.
- [0100] The preferred locus of biosynthesis, of fatty acids, oils or lipids for example, is generally the seed or cell layers of the seed so that a seed-specific expression of the Δ-8-desaturase, Δ-9-elongase and/or Δ-5-desaturase gene is appropriate. It is, however, obvious that the biosynthesis of fatty acids, oils or lipids need not be limited to the seed tissue but rather can also occur in tissue-specific manner in all other parts of the plant in epidermis cells or in the nodules for example.
 - **[0101]** A constitutive expression of the exogenous Δ -8-desaturase, Δ -9-elongase and/or Δ -5-desaturase gene is, moreover, advantageous. On the other hand, however, an inducible expression may also appear desirable.

[0102] The efficiency of the expression of the Δ -8-desaturase, Δ -9-elongase and/or Δ -5-desaturase gene can be determined, for example, *in vitro* by shoot meristem propagation. In addition, an expression of the Δ -8-desaturase, Δ -9-elongase and/or Δ -5-desaturase gene modified in nature and level and its effect on fatty acid, oil or lipid biosynthesis performance can be tested on test plants in greenhouse trials.

[0103] An additional object of the invention comprises transgenic plants transformed by an expression cassette containing a Δ -8-desaturase, a Δ -9-elongase and/or a Δ -5-desaturase gene sequence according to the invention or DNA sequences hybridizing therewith, as well as transgenic cells, tissue, parts and reproduction material of such plants.

⁴⁵ Particular preference is given in this case to transgenic crop plants such as by way of example barley, wheat, rye, oats, corn, soybean, rice, cotton, sugar beet, oilseed rape and canola, sunflower, flax, hemp, thistle, potatoes, tobacco, tomatoes, tapioca, cassava, arrowroot, alfalfa, lettuce and the various tree, nut and vine species. **10101** For the purposes of the investion plante are more, and disct ideance plante measure of the investion plante.

[0104] For the purposes of the invention plants are mono- and dicotyledonous plants, mosses or algae.

[0105] A further refinement according to the invention are transgenic plants as described above which contain a nucleic acid sequence according to the invention or a expression cassette according to the invention.

[0106] Also disclosed are:

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- A method for the transformation of a plant comprising the introduction of expression cassettes according to the invention containing a Δ-8-desaturase, a Δ-9-elongase and/or a Δ-5-desaturase gene sequence derived from algae such as Euglenia or Isochrysis, fungi such as Mortierella or mosses such as Physcomitrella or DNA' sequences hybridizing therewith into a plant cell, into callus tissue, an entire plant or protoplasts of plants.
 - A method for producing PUFAs, wherein the method comprises the growing of a transgenic organism comprising

a nucleic acid as described herein or a vector encoding a Δ -8-desaturase, a Δ -9-elongase and/or a Δ -5-desaturase which specifically synthesize poly unsaturated fatty acids with at least three double bonds in the fatty acid molecule

- Use of a Δ-8-desaturase, a Δ-9-elongase and/or a Δ-5-desaturase DNA gene sequence or DNA sequences hybridizing therewith for the production of plants having an increased content of fatty acids, oils or lipids having at least three double bonds due to the expression of said Δ-8-desaturase, Δ-9-elongase and/or Δ-5-desaturase DNA sequence in plants.
- [0107] Also disclosed is a method for producing unsaturated fatty acids comprising: introducing at least one said nucleic acid sequence described herein or at least one nucleic acid construct or vector containing said nucleic acid sequence into a preferably oil-producing organism such as a plant or a fungi; growing said organism; isolating oil contained in said organism; and liberating the fatty acids present in said oil. These unsaturated fatty acids advantageously contain at least three double bonds in the fatty acid molecule. The fatty acids may be liberated from the oils or lipids, for example by basic hydrolysis, e.g. using NaOH or KOH or by acid hydrolysis preferably in the presence of an alcohol such as
- ¹⁵ methanol or ethanol. Said fatty acid liberation leads to free fatty acids or to the corresponding alkyl esters of the fatty acids. In principle an enzymatic hydrolysis for example with a lipase as enzyme is also possible. Starting from said free fatty acids or fatty acid alkyl esters mono-, di- and/or triglycerides can be synthesized either chemically or enzymatically. In another preferred embodiment of the inventive process the alkyl ester of the fatty acids are produced from the oils and lipids by transesterification with an enzyme of with conventional chemistry. A preferred method is the production of
- 20 the alkyl ester in the presence of alcohalates of the corresponding lower alcohols (C1 to C10 alcohols such as methanol, ethanol, propanol, butanol, hexanol etc.) such as methanolate or ethanolate. Therefore as the skilled worker knows the alcohol in the presence of a catalytic amount of a base such as NaOH or KOH is added to the oils or lipids.
 [0108] A method for producing triglycerides having an increased content of unsaturated fatty acids comprising: intro-
- ducing at least one said nucleic acid sequence according to the invention or at least one expression cassette according
 to the invention into an oil-producing organism; growing said organism; and isolating oil contained in said organism; is also numbered among the objects of the invention.

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[0109] Also disclosed is a method for producing triglycerides having an increased content of unsaturated fatty acids advantageously having an increased content of unsaturated fatty acids is a method wherein the fatty acids are liberated from the triglycerides with the aid of basic hydrolysis known to those skilled in the art or by means of an enzyme such as a lipase.

[0110] The methods specified above advantageously allow the synthesis of fatty acids or triglycerides having an increased content of fatty acids containing at least three double bonds in the fatty acid molecule.

[0111] The methods identified above advantageously allow the synthesis of fatty acids or triglycerides having an increased content of fatty acids containing at least three double bonds, wherein the substrate used for the reaction of the t-2 dependence of the sector of the substrate used for the reaction of the t-2 dependence of the sector of the se

- ³⁵ the Δ -8-desaturase, Δ -9-elongase and/or Δ -5-desaturase is preferably linoleic acid ($C_{20:2}^{\Delta9,12}$) acid and/or α -linolenic acid ($C_{18:2}^{\Delta9,12,15}$). In this way the method identified above advantageously allows in particular the synthesis of fatty acids derived from linoleic acid ($C_{20:2}^{\Delta9,12}$), α -linolenic acid ($C_{18:2}^{\Delta9,12,15}$), γ -linoleic acid ($C_{18:3}^{\Delta6,9,12}$), stearidonic acid ($C_{18:4}^{\Delta6,9,12,15}$), dihomo- γ -linoleic acid ($C_{20:3}^{\Delta8,11,14}$) or such as by way of example eicosapentaenoic acid and arachidonic acid.
- 40 [0112] Examples of organisms for the said methods as described above are plants such as Arabidopsis, Primulaceae, borage, barley, wheat, rye, oats, corn, soybean, rise, cotton, sugar beet, oilseed rape and canola, sunflower, flax, hemp, potatoes, tobacco, tomatoes, rape, tapioca, cassava, arrowroot, alfalfa, peanut, castor oil plant, coconut, oil palm, safflower (Carthamus tinctorius) or cocoa bean, algae or protozoa such as dinoflagellates like Crypthecodinium. Preference is given to organisms which can naturally synthesize oils in relatively large quantities such as plants such as soybean,
- oilseed rape, coconut, oil palm, safflower, castor oil plant, Calendula, peanut, cocoa bean or sunflower, and particular preference is given to soybean, oilseed rape, sunflower, flax, Primulaceae, borage or Carthamus.
 [0113] Depending on the host organism, the organisms used in the methods are grown or cultured in the manner known to those skilled in the art. Microorganisms such as fungi or algae are usually grown in a liquid medium containing a carbon source, usually in the form of sugars, a nitrogen source, usually in the form of organic nitrogen sources such
- ⁵⁰ as yeast extract or salts such as ammonium sulfate, trace elements such as iron, manganese or magnesium salts and optionally vitamins at temperatures of between 10 °C and 60 °C, preferably between 15 °C and 40 °C with exposure to gaseous oxygen. In doing so the pH of the nutrient liquid may be kept at a fixed value, that is during growth it is or is not regulated. Growth can ensue in batch mode, semibatch mode or continuously. Nutrients can be provided at the start of fermentation or be fed in semicontinuously or continuously.
- ⁵⁵ **[0114]** After transformation plants are first of all regenerated as described above and then cultured or cultivated as normal.

[0115] After growth the lipids are isolated from the organisms in the usual way. For this purpose, after harvesting the organisms may first of all be digested or used directly. The lipids are advantageously extracted using suitable solvents

such as apolar solvents like hexane or ethanol, isopropanol or mixtures such as hexane/isopropanol, phenol/chloroform/isoamyl alcohol at temperatures of between 0 °C and 80 °C, preferably between 20 °C and 50 °C. The biomass is usually extracted with an excess of solvent, for example an excess of solvent to biomass of 1:4. The solvent is then removed, for example by distillation. Extraction can also be done using supercritical CO_2 . After extraction the remaining

- ⁵ biomass may be removed, for example by filtration.
 [0116] The crude oil isolated in this way can then be further purified, for example by removing cloudiness by treatment with polar solvents such as acetone or chloroform and then filtration or centrifugation. Further purification through columns is also possible.
 - [0117] In order to obtain the free acids from the triglycerides the latter are saponified in the usual way.
- ¹⁰ **[0118]** Also disclosed are unsaturated fatty acids and triglycerides having an increased content of unsaturated fatty acids produced by the methods identified above and use thereof for producing foods, animal feeds, cosmetics or pharmaceuticals. For this purpose the latter are added in customary quantities to the foods, the animal feed, the cosmetics or pharmaceuticals.
- [0119] Said unsaturated fatty acids as well as triglycerides having an increased content of unsaturated fatty acids produced by the methods identified above are the result of the expression of the nucleic acids according to the invention in the various host organisms. This results overall in a modification of the composition of the compounds in the host cell containing unsaturated fatty acids by comparison with the original starting host cells which do not contain the nucleic acids. These modifications are more marked in host organisms, for example plant cells, which naturally do not contain the proteins or enzymes encoded by the nucleic acids than in host organisms which naturally do contain the proteins or
- enzymes encoded by the nucleic acids. This gives rise to host organisms containing oils, lipids, phospholipids, sphingolipids, glycolipids, triacylglycerols and/or free fatty acids having a higher content of PUFAs with at least three double bonds. For the purposes of the invention, by an increased content is meant that the host organisms contain at least 5 %, advantageously at least 10 %, preferably at least 20 %, particularly preferably at least 30 %, most particularly preferably at least 40 % more polyunsaturated fatty acids by comparison with the initial organism which does not contain the nucleic
- ²⁵ acids according to the invention. This is particularly the case for plants which do not naturally contain longer-chain polyunsaturated C₂₀ or C₂₂ fatty acids such as EPA or ARA. Due to the expression of the nucleic acids novel lipid compositions are produced by said means these being a further aspect of the invention. [0120] The invention is explained in more detail by the following examples.
- 30 Examples

Example 1: General cloning methods

[0121] The cloning methods, such as by way of example restriction cleavages, agarose gel electrophoresis, purification of DNA fragments, transfer of nucleic acids to nitrocellulose and nylon membranes, linkage of DNA fragments, transformation of - Escherichia coli cells, culture of bacteria and sequence analysis of recombinant DNA, were carried out as described in Sambrook et al. (1989) (Cold Spring Harbor Laboratory Press: ISBN 0-87969-309-6).

Example 2: Sequence analysis of recombinant DNA

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[0122] Sequencing of recombinant DNA molecules was done using a laser fluorescence DNA sequencer from the ABI company by the method of Sanger (Sanger et al. (1977) Proc. Natl. Acad. Sci. USA74, 5463-5467). Fragments resulting from a polymerase chain reaction were sequenced and checked to prevent polymerase errors in the constructs to be expressed.

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Example 3: Cloning of the \triangle -8-desaturase from Euglena gracilis (= SEQ ID NO: 1)

[0123] As a template for PCR amplification, cDNA from Euglena gracilis Strain Z was used. The cDNA was synthesised from total RNA extracted from cultures of E. gracilis strain Z. Unique primers to the initiating methionine and the stop codon of the Euglena Δ -8-desaturase were synthesized as shown, including restriction sites as detailed

Primer 1: EDELTA8BamF ATGGATCCACCATGAAGTCAAAGCGCCAA Primer 2: EDELTA8XhoR ATCTCGAGTTATAGAGCCTTCCCCGC

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PCR protocol

[0124]

Addition temperature: 1 min at 45 °C Denaturing temperature: 1 min at 94 °C Elongation temperature: 2 min at 72 °C Number of cycles: 30

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[0125] The PCR products were separated on an agarose gel and a 1270 bp fragment was isolated. The PCR fragment was cloned in the pGEM-T easy vector (Promega) and the insert was then sequenced. This revealed the presence of an open reading frame of 1266 base pairs, encoding a protein of 421 amino acid residues and a stop codon. The C-terminus of the cloned Δ -8-desaturase has high homologies to the Δ -8-desaturase published by Wallis and Browse (Archives of Biochem. and Biophysics, Vol. 365, No. 2, 1999) which is reported to be an enzyme of 422 residues; see also related sequence by these authors [GenBank AF139720/ AAD45877] which purports to relate to the same Δ -8-desaturase but describes an open reading frame of 419 residues]. The deduced amino acid sequence the Euglena Δ -8-desaturase described in this present invention differs from that previously described by heterogeneity at the N-terminus. In particular, the first 25 amino acid residues of LARS Δ -8-desaturase is:

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MKSKRQALP LTIDGTTYDVS AWVNF

[0126] Whereas the sequence described by Wallis & Browse is:

- 20 MKSKRQALS PLQLMEQTYDV SAWVN (as given in ABB 1999)
 - [0127] Or, alternatively

MKSKRQALSPLQLMEQTYDVVNFH (as given in GenBank AAD45877)

- [0128] Said heterogeneity present at the N-terminus of the desaturase sequence is not resultant of the PCR amplifi-
- ²⁵ cation or primers. The distinctions are true differences between the proteins.

Example 4: Construction of transgenic plants expressing the Isochrysis galbana elongase component IgASE1

- [0129] The cloning of IgASE1 cDNA is described in: Qi, B., Beaudoin, F., Fraser, T., Stobart, A. K., Napier, J.A. and Lazarus, C.M.Identification of a cDNA encoding a novel C18-Δ-9-polyunsaturated fatty acid-specific elongating activity from the docosahexaenoic acid (DHA)-producing microalga, Isochrysis galbana. FEBS Letters 510, 159-165 (2002). The cDNA was released from plasmid vector pCR2.1-TOPO by digestion with *KpnI*, and ligated into the *KpnI* site of the intermediate vector pBlueBac 4.5 (Invitrogen). Recombinant plasmids were screened for insert orientation with *Eco*RI. The insert was released from a selected plasmid with *PstI* plus *Eco*RI and ligated into binary vector plasmid pCB302-1
- 35 (Xiang *et al*, 1999) that had been cut with the same enzymes. This placed the IgASE1 coding region under the control of the CaMV 35S promoter as a translational fusion with the transit peptide of the small subunit of Rubisco (Xiang *at al.,* 1999), with the intention of targeting the elongase component to chloroplasts when expressed in transgenic plants. This recombinant binary vector was designated pCB302-1ASE. To construct a similar vector with expression of the elongase component targeted to the microsomal membrane, the IgASE1 coding region was removed from the intermediate vector
- ⁴⁰ by digestion with *Bam*HI plus Spel, and ligated into the corresponding sites of pCB302-3 (Xiang *et al.*, 1999, in which the map of pCB302-3 is incorrect: the CaMV 35S promoter (plus omega sequence) and nos terminator regions are reversed with respect to MCS2). This recombinant binary vector was designated pCB302-3ASE.
 - Example 5: Plant expression of the elongase

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[0130] Binary vectors were transferred to *Agrobacterium tumefaciens* strain GV3101 by electroporation; transformed colonies were selected on medium containing 50 μ g ml⁻¹ kanamycin. Selected colonies were gown to stationary phase at 28°C, then the cells were concentrated by centrifugation and resuspended in a dipping solution containing 5% sucrose, 0.03% Silwet-177 and 10 mM MgCl₂.

- ⁵⁰ **[0131]** Seeds of *Arabidopsis thaliana* ecotype Columbia 4 were germinated on one-half-strength Murashige and Skoog medium, and seedlings were transferred to compost in 15 cm flower pots. Plants were grown to flowering stage in a growth cabinet at 21°C, with a 23 light and 1 hour dark cycle. Plant transformation was carried out by the floral dipping method of Clough and Bent (1998, Floral dip: a simplified method for Agrobacterium-mediated transformation of Arabidopsis thaliana. Plant Journal 16, 735-743 (1998), essentially as follows:
- ⁵⁵ **[0132]** For each construct two pots containing 16 plants were inverted in the dipping solutions containing transformed *A. tumefaciens* (described above). The plants were then covered with a plastic bag and left at room temperature in the dark overnight. The bag was then removed and the plants transferred to the growth cabinet. Dipping (with fresh *A. tumefaciens* solutions) was repeated after 5 days and the plants were allowed to set seed. Bulked seed from dipped

plants (= T1 seed) was collected, and approximately 10000 seed sprinkled onto compost in a seed tray, and, after stratification at 4°C for 2 days, cultivated in the growth cabinet. When seedlings had reached the 2 to 4 true-leaf stage they were sprayed with Liberty herbicide (Aventis, 0.5g glufosinate-ammonium I⁻¹), and spraying was repeated one week later. Twelve herbicide-resistant plants were selected and potted on for each line (chloroplast or cytoplasm targeted

- ⁵ elongase component), and allowed to self fertilize. Samples of T2 seed collected from these plants were germinated on one-half-strength Murashige and Skoog medium containing Liberty (5 mg glufosinate-ammonium I⁻¹). T3 seed collected from individual surviving plants was then again germinated on Liberty plates to screen for lines that had ceased segregating for herbicide resistance. Total fatty acids extracted from leaves of such lines were analysed and those with the greatest C20 content (CB12-4 with the chloroplast-targeted elongase component and CA1-9 with the cytoplasm-targeted
- ¹⁰ elongase component) selected.

Example 6: Production of transgenic plants expressing the *Isochrysis galbana* elongase component IgASE1 and the *Euglena gracilis* ∆8 desaturase EUGD8

- 15 [0133] The △-8-desaturase coding region was removed from the yeast expression vector pESC-Trp with BamHI plus Xhol, ligated into the BamHI and Xhol sites of pBlueBac 4.5 (Invitrogen) and transformed into E. coli strain Tam1. The insert was removed from a recombinant plasmid with Bg/II and BamHI, ligated into the BamHI site of pBECKS₁₉.6 and transformed into E. coli strain Tam1. DNA minipreparations were made of the recombinant plasmids of 6 transformant colonies; these were digested with Xhol to determine the orientation of insertion of the desaturase coding region in the
- ²⁰ binary vector. One recombinant plasmid with the insert in the correct orientation for expression from the CaMV 35S promoter was transferred to *Agrobacterium tume-faciens* strain GV3101 by electroporation and a dipping solution pre-pared from a transformed colony as described above.

[0134] *Arabidopsis thaliana* lines CB12-4 and CA1-9 (see above) were subjected to floral dipping as described above. Approximately 2000 T1-seed from each line were spread on 15 cm petri dishes containing one-half-strength Murashige

- ²⁵ and Skoog (solid) medium supplemented with 50 µg ml⁻¹ kanamycin and germinated in the growth cabinet. 12 kanamycinresistant plants of the CA1-9 parental line and 3 plants of the CB12-4 parental line were transferred to potting compost and further cultivated in the growth room. Fatty acid analysis was conducted on a lea taken from each of the T2 plants, which were allowed to mature and set seed.
- 30 References

[0135]

McCormac, A.C., Eliott, M.C. and Chen, D-F.; pBECKS. A flexible series of binary vectors for Agrobacteriummediated plant transformation. Molecular Biotechnology 8, 199-213 (1997).

Xiang, C., Han, P., Lutziger, I., Wang, K. and Oliver, D.J.; A mini binary vector series for plant transformation. Plant Molecular Biology 40, 711-717 (1999).

⁴⁰ Example 7: Production of transgenic plants expressing the *Isochrysis galbana* elongase component IgASE1 and the *Euglena gracilis* Δ 8 desaturase EUGD8 and a Δ 5 desaturase

[0136] The $\Delta 5$ desaturase from Phaeodactylum tricornutum was cloned into the pGPTV plasmid (Becker, D. et al.; Plant Mol. Biol. 20 (1992), 1195-1197) habouring a hygromycin resistence selectable marker gene. For seed-specific expression the USP promoter from Vicia faber was cloned 5'-prime to the ATG of the $\Delta 5$ desaturase.

[0137] The binary vector was transferred to *Agrobacterium tumefaciens* strain GV 3101 and transformed colonies were selected on medium containing 30 μ gml⁻¹ hygromycin. Selected Agrobacteria were used for the transformation (flower transformation) of Arabidopsis plants carrying the T-DNA insertions with the Δ 9 elongase and the Δ 5 desaturase. **[0138]** Arabidopsis thaliana seedlings were germinated on Murashige and Skoog medium containing hygromycin and resistent plants were transferred to the greenhouse.

[0139] Seeds collected from individual plants were harvested and the total fatty acid profile was analyzed using GC methods.

Example 8: Cloning of expression plasmids for seed-specific expression in plants

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[0140] pBin-USP is a derivative of the plasmid pBin19. pBin-USP was produced from pBin19 by inserting a USP promoter as an EcoRI-BaMHI fragment into pBin19 (Bevan et al. (1980) Nucl. Acids Res. 12, 8711). The polyadenylation signal is that of gene 3 of the T-DNA of the Ti plasmid pTiACH5 (Gielen et al., (1984) EMBO J. 3, 835), whereby

nucleotides 11749-11939 were isolated as a Pvull-HindIII fragment and after addition of Sphl linkers to the Pvull interface between the SpHI-HindIII interface of the vector were cloned. The USP promoter corresponds to nucleotides 1-684 (gene bank accession number X56240), wherein a part of the nonencoding region of the USP gene is contained in the promoter. The promoter fragment running to 684 base pairs was amplified by standard methods by means of commercial T7 standard primer (Stratagene) and using a synthesized primer through a PCR reaction.

Primer sequence:

5'-GTCGACCCGCGGACTAGTGGGCCCTCTAGACCCGGGGGGATCC

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GGATCTGCTGGCTATGAA-3'

[0141] The PCR fragment was cut again using EcoRI/Sall and inserted into the vector pBin19 with OCS terminator. The plasmid having the designation pBinUSP was obtained. The constructs were used for transforming Arabidopsis thaliana, oilseed rape, tobacco and linseed.

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Example 9: Production of transgenic oil crops

- [0142] Production of transgenic plants (modified in accordance with Moloney et al., 1992, Plant Cell Reports, 8:238-242)
 [0143] To produce transgenic oilseed rape plants binary vectors in Agrobacterium tumefaciens C58C1:pGV2260 or
 ²⁰ Escherichia coli were used (Deblaere et al, 1984, Nucl. Acids. Res. 13, 4777-4788). For transforming oilseed rape plants (var. Drakkar, NPZ Nordeutsche Pflanzenzucht, Hohenlieth, Germany) a 1:50 dilution of an overnight culture of a positively transformed agrobacteria colony in Murashige-Skoog medium (Murashige and Skoog 1962 Physiol. Plant. 15, 473) containing 3 % of saccharose (3MS medium) was used. Petioles or hypocotyledons of freshly germinated sterile rape plants (approx. 1 cm² each) were incubated in a Petri dish with a 1:50 agrobacteria dilution for 5-10 minutes. This was
- ²⁵ followed by 3-day concubation in darkness at 25 °C on 3MS medium containing 0.8 % of Bacto-Agar. After three days, culturing was continued with 16 hours of light / 8 hours of darkness and in a weekly cycle on MS medium containing 500 mg/l of Claforan (sodium cefotaxime), 50 mg/l of kanamycin, 20 microM of benzylaminopurine (BAP) and 1.6 g/l of glucose. Growing shoots were transferred onto MS medium containing 2 % of saccharose, 250 mg/l of Claforan and 0.8 % of Bacto-Agar. If after three weeks no roots had formed 2-indolylbutyric acid was added to the medium as a growth
- ³⁰ hormone for rooting purposes.

[0144] Regenerated shoots were obtained on 2MS medium using kanamycin and Claforan, transferred into soil after rooting and after culturing grown for two weeks in a climate-controlled chamber, brought to blossom and after harvesting of ripe seed investigated for Δ -8-desaturase expression by means of lipid analyses. Lines having increased contents of double bonds at the Δ -8-position were identified. In the stably transformed transgenic lines functionally expressing the

transgene it was found that there is an increased content of double bonds at the Δ -8-position by comparison with untransformed control plants.

[0145] The same procedure was done to create plants with \triangle -9-elongase and/or \triangle -5-desaturase activity.

a) Transgenic flax plants

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[0146] Transgenic flax plants may be produced, for example by the by the method Bell et al., 1999, In Vitro Cell. Dev. Biol.-Plant. 35(6):456-465, by means of particle bombardment. Agrobacteria-mediated transformations can be produced, for example, as described by Mlynarova et al. (1994), Plant Cell Report 13: 282-285.

45 Example 10: Lipid extraction from seed and leave material

[0147] Plant material (approx 200 mg) was first of all mechanically homogenized by means of triturators in order to render it more amenable to extraction.

- [0148] The disrupted cell sediment was hydrolyzed with 1 M methanolic hydrochloric acid and 5 % dimethoxypropane for 1 h at 85 °C and the lipids were transmethylated. The resultant fatty acid methyl esters (FAMEs) were extracted in hexane. The extracted FAMEs were analyzed by gas-liquid chromatograph using a capillary column (Chrompack, WCOT fused silica, CP wax 52 CB, 25 m, 0.32 mm) and a temperature gradient of from 170 °C to 240 °C in 20 min and 5 min at 240 °C. The identity of the fatty acid methyl esters was confirmed by comparison with corresponding FAME standards (Sigma). The identity and the position of the double bond was further analyzed by means of GC-MS by suitable chemical
- derivatization of the FAME mixtures, e.g. to form 4,4-dimethoxyoxazoline derivatives (Christie, 1998).
 [0149] Figure 1 shows the fatty acid profile (FAMes) of leaf tissue from wildtype Arabidopsis thaliana as a control. Figure 2 shows the fatty acid profile (FAMes) of leaf tissue from transgenic Arabidopsis expressing the Isochrysis Δ-9-elongase (see example 4). This Arabidopsis line was subsequently re-transformed with the Euglena Δ-8-desaturase.

The fatty acid profile (FAMes) of said double transformed Arabidopsis line (Line IsoElo X Eu D8 des) is given in Figure 3. **[0150]** Furthermore this double transformed Arabidopsis line (Line IsoElo X Eu D8 des) was subsequently re-transformed with the Mortierella $\Delta 5$ desaturase (Mort $\Delta 5$) gene. The fatty acid profile (FAMes) of said triple transformed Arabidopsis line (Line IsoElo X EU D8 des x Mort $\Delta 5$) is given in Figure 4.

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Example 11: GC profiles of Arabidopsis leaf fatty acid methyl esters from different transgenics

[0151] Figure 5 shows GC profiles of Arabidopsis leaf fatty acid methyl esters extracted from wild type (WT 5a), single transgenic plants expressing Isochrysis galbana $\Delta 9$ elongase gene Ig ASE1 (5b), double transgenic plant expressing the Ig ASE1 and Euglena $\Delta 8$ desaturase (EU $\Delta 8$) genes (5c) and the triple transferic plant expressing the Ig ASE1, Eu $\Delta 8$ and the Mortierella $\Delta 5$ desaturase (Mort $\Delta 5$) genes (5d).

Table 1 shows the fatty acid composition of *Arabidopsis* plants prepared from wild type (Wt), single transgenic plant expressing the Isochrysis galbana IgASE1 elongase gene, double transgenic plants expressing the IgASE1 elongase gene and the Euglena $\Delta 8$ desaturase gene and triple transgenic plants expressing the IgASE1, the Euglena $\Delta 8$ and the Mortierella $\Delta 5$ desaturase gene. Analysis is of leaf tissue from rosette stage *Arabidopsis* plants. Each value represents the average of 2 measurements.

Fatty agid (mall)/ of			Plant source	
Fatty acid (mol% of total)	Wt	IgASE1 transgenic	lgASE1+Eu∆8 transgenic	IgASE1+Eu∆8+Mort∆5 transgenic
16:0	19.9	19.2	14.7	14.2
16:1	2.8	3.3	1.8	2.3
16:3	13.1	12.2	19.9	15.4
18:0	1.7	2.4	0.8	1.5
18:1n-9	1.7	5.1	1.6	3.4
18:2n-6	11.2	9.0	4.2	6.6
18:3n-3	50.1	31.0	36.0	31.2
20:2n-6	-	7.9	0.9	3.2
20:3,∆5,11,14	-			1.5
20:3n-6	-	-	9.1	1.5
20:4n-6 (ARA)	-	-		6.6
20:3n-3	-	9.9	4.0	4.8
20,4∆5,11,14,17	-	-	-	1.6
20:4n-3	-	-	7.2	2.9
20:5n-3 (EPA)	-	-	-	3.3
Total C20 PUFAs	-	17.8	21.2	22.2

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[0152] All transgenes are under the control of the 35S-CaMV viral promoter. Isochrysis $\Delta 9$ elongase (IgASE1) with SSU Rubisco transit sequence [T-DNA Basta-r] were retransformed with Euglena $\Delta 8$ -desaturase^{mut175+313} [T-DNA Kanamycin-r]. The double transformed line, which is homozygous for both Basta-r and Kanamycin-r, were transformed again with Mortierella $\Delta 5$ desaturase (T-DNA Hygromycin-r). The resulting triple transformed line is homozygous for both Basta-r and Kanamycin-r, but heterozygous for Hygromycin-r.

SEQUENCE LISTING

₅₅ [0153]

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45	His 145	Asp	Ile	Cys	His	His 150	Gln	Thr	Phe	Lys	Asn 155	Arg	Asn	Trp	Asn	Asn 160
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55	Cys	Trp	Lys	Asp		His	Asn	Ala	His	His	Ser	Ala	Thr	Asn		Gln

		180	185	190
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25	Gln Phe Tyr	Arg Ser Gln Typ 260	: Lys Lys Glu Ala Ile 265	Gly Leu Ala Leu 270
30	His Trp Thr 275	Leu Lys Ala Leu	1 Phe His Leu Phe Phe 280	Met Pro Ser Ile 285
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		His	Asn 370	Leu	Thr	Ala	Val	Ser 375	Tyr	Gln	Val	Glu	Gln 380	Leu	Cys	Gln	Lys	
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10		Leu	Leu	Arg	Tyr	Leu	Ala	Val	Phe	Ala	Arg	Met	Ala	Glu	Lys	Gln	Pro	
						405					410					415		
15		۲ ۱ -	C 111	Tura	7 - 7	Ton												
		Ala	GTĀ	цур	420	Бец												
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	-	•			5										10			
40	gao	ccg	gaa	ato	ctc	att	ggc	acc	ttc	tcg	tac	: ttg	cta	cto	aaa	ccg	g .9	6
	Asr	Pro	Glu			Ile	Gly	• Thr			Туг	: Leu	Leu			Pro)	
				20					25					30	I			
45	cto	, ctc	cgc	: aat	tcc	ggg	ctg	gtg	r gat	gag	aag	ı aag	ggc	gca	. tac	ago	y 1	.44
	Lei	ı Lev			Ser	Gly	Leu			Glu	Lys	: Lys			. Tyr	Arg	J	
50			35	5				40)				45					

	acg	tcc	atg	atc	"tgg	tac	aac	gtt	ctg	ctg	gcg	ctc	ttc	tct	gcg	ctg	192
	Thr	Ser	Met	Ile	Trp	Tyr	Asn	Val	Leu	Leu	Ala	Leu	Phe	Ser	Ala	Leu	
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10	Ser	Phe	Tyr	Val	Thr	Ala	Thr	Ala	Leu	Gly	Trp	Asp	Tyr	Gly	Thr	Gly	
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	Ala	Trp	Leu	Arg		Gln	Thr	Gly	Asp	Thr	Pro	Gln	Pro	Leu	Phe	Gln	
					85					90					95		
20										,							
		ccg															336
	Cys	Pro	Ser		Va⊥	Trp	Asp	Ser		Leu	Phe	Thr	Trp		Ala	Lys	
25				100					105					110			
	~~~	++-	tat	t > 0	+	220	t	~+~~	~~~~	tag	ata	<b>a</b> 2 <b>a</b>		~~~	+ ~ ~	ata	201
	-	ttc				_						_	-	-		-	384
30	ALA	Phe	115	τΥτ	Ser	цур	IYL	120	Gru	TÄT	цец		125	ATG	пр	Deu	
			440					120					<u></u>		· -· .		
	agg	gtc	tcc	ttt	ctc	caq	acc	ttc	cac	сас	ttt	aac	aca	cca	taa	gat	432
35		Val											-	_	-	-	
		130					135					- 140			-	2	
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45																	
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55	Thr	Ala	Ala	Gly	Tyr	Lys	Phe	Lys	Ala	Lys	Pro	Leu	Ile	Thr	Ala	Met	
											•						

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	1	.80	185	190
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	din fie Cys G	200 200	y Phe Leu Leu Val Trp ) 205	
10	aac gtc ccc t	gc ttc aac tcg gad	c aaa ggg aag ttg ttc	agc tgg gct 672
15	Asn Val Pro C 210	ys Phe Asn Ser Asp 215	o Lys Gly Lys Leu Phe 220	Ser Trp Ala
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20	Phe Asn Tyr A 225	la Tyr Val Gly Sei 230	r Val Phe Leu Leu Phe 235	Cys His Phe 240
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50	Leu Leu Ar	g Asn Ser Gly Leu	Val Asp Glu Lys Lys	Gly Ala Tyr Arg

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5	Thr Ser Met 50	Ile Trp Tyr	Asn Val Leu Leu Ala 55	Leu Phe Ser Ala Leu 60
10	Ser Phe Tyr 65	Val Thr Ala 70	Thr Ala Leu Gly Trp 75	Asp Tyr Gly Thr Gly 80
15	Ala Trp Leu	Arg Arg Gln	Thr Gly Asp Thr Pro 90	Gln Pro Leu Phé Gln 95
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45	Thr Ala Ala	Gly Tyr Lys 180	Phe Lys Ala Lys Pro 185	Leu Ile Thr Ala Met 190
50	Gln Ile Cys 195		Gly Gly Phe Leu Leu 200	Val Trp Asp Tyr Ile
55	Asn Val Pro 210	Cys Phe Asn	Ser Asp Lys Gly Lys 215	Leu Phe Ser Trp Ala 220

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	:	225					230					235					240	
5																		
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			1 Pro	o As <u></u>		a Ası	o Lys	s Lei	1 Arg			g GLI	n Th	r Th			.1	
30	1					5				10	J				1	5		
		220			- 00	t aat		• ata				<b>a</b> aa		a at	+ + ~	a	·+-	96
						t get									1.1			50
35	AId	Бλя	, 111;	20		a Ala	1 111		2!			1 911		у це 3:		5 56	T.	
				20		-			4.					J	0			
	cta	tet	t.c	g cto	c aa	a aa	gaa	a daa	a ato	ta	c ato	c da	c aa	a at	c at	c ta	t	144
40	_			-		s Gly												
			3				•	4(					. 4					
		-							-									
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	Asp	Let	ı Gli	n Sei	r Ph	e Asr	o His	s Pro	5 G1	y Gl	y Glı	ı Th	r Il	e Ly	s Me	t Ph	e	
		50	)				55	5				6	0					
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	Arg	Glu	Val	Phe	Lys	Ile	Val	Arg	Arg	Gly	Lys	Asp	Phe	Gly	Thr	Leu	۰.
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30		130					135	,				140					
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05													ctg			-	480
35		Tyr	His	Trp	Val		Thr	Gly	Thr	Ser		Leu	Leu	Ala	Val		
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10	* ~ ~	~~~~				~~~~	~+~				+			~ ~ ~		~~~	500
40							-					-	cag		-	-	528
	тÀт	GTÀ	TTE	Ser	165	AIG	Met	TIE	GTÀ	170	ASII	Val	Gln	UTS	175 ASD	AId	
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				180				5	185	₽				190		1	
50																	
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													Trp				
55	-	-		~			· · · · ·	-4		<b>_</b>	- 27		-				

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	His	Trp 210	Thr	His	His	Ala	Tyr 215	Thr	Asn	His	Ala	Glu 220	Met	Asp	Pro	Asp	
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	-			-	Glu		-					-			-	-	720
	225	1110	OT J	ML Q	Gru	230	1100	Dea	Dea		235	1155		110	Deu	240	
15	227					230	*				235					240	<i>z</i> .
	cat	ccc	gct	cgt	acc	tgg	cta	cat	cgc	ttt	caa	gca	ttc	ttt	tac	atg	768
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							•									. مت ر	
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	-	he Leu	Ser G		Phe	Thr	Gly	Gly		Asn	Phe	Gln			
	385			390					395	,			· .	400	
25	-		*** ~		-+			~~*	+ ~ ~	+		***	~++	~~~	1740
		ac ttg is Leu													1248
	n15 n	ITS Ded		105	Met	Der	Det	410	тр	ιγι	FIO	TÀT	415	ALA	
30			-	100				710					470		
	ccc a	ag gtc	cgc g	gaa att	tgc	gcc	aaa	cac	ggc	gtc	cac	tac	gcc	tac	1296
		ys Val				-				_			-		
35		• *	420				425					430			
	tac c	cg tgg	atc c	cac caa	aac	ttt	ctc	tcc	acc	gtc	cgc	tac	atg	cac	1344
40	Tyr P	ro Trp	Ile H	His Gln	Asn	Phe	Leu	Ser	Thr	Val	Arg	Tyr	Met	His	
		435				440			•		445				
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	Ala A	la Gly	Thr C	Gly Ala	Asn	Trp	Arg	Gln	Met	Ala	Arg	Glu	Asn	Pro	•
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<213> Phaeodactylum tricornutum

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10	Al	a Lys	His	Asn 20	Ala	Ala	Thr	Ile	Ser 25	Thr	Gln	Glu	Arg	Leu 30	Cys	Ser
15	Le	ı Ser	Ser	Leu	Lys	Gly	Glu	Glu	Val	Cvs	Ile	Asp	Glv	Ile	Ile	Tvr
			35		-	-		40		-		<b>-</b> .	45			-
20	۵s	) Leu	Gln	Ser	Phe	Asp	His	Pro	Glv	Glv	Glu	ጥከዮ	Tle	Lvs	Met	Phe
		50					55		1	1		60		-1-		
25	Gl	7 Gly	Asn	Asp	Val	Thr	Val	Gln	Tyr	Lys	Met	Ile	His	Pro	Tyr	His
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30	Th	r Glu	Lys	His	Leu 85	Glu	Lys	Met	Lys	Arg 90	Val	Gly	Lys	Val	Thr 95	Asp
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40	Ar	g Glu	Val 115	Phe	Lys	Ile	Val	Arg 120	Arg	Gly	Lys	Asp	Phe 125	Gly	Thr	Leu
45	Gl	y Trp 130	Phe	Phe	Arg	Ala	Phe 135	Cys	Tyr	Ile	Ala	Ile 140	Phe	Phe	Tyr	Leu
50	Gl	n Tyr	His	Trp	Val	Thr	Thr	Gly	Thr	Ser	Trp	Leu	Leu	Ala	Val	Ala

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5	Tyr-Gly Ile Ser Gl		Asn Val Gln His Asp Ala 175
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55	Leu Glu Trp Ser Tr 32		Ile Met Leu Met Gly Val 335

	Ala	Glu	Ser	Leu	Ala	Leu	Ala	Val	Leu	Phe	Ser	Leu	Ser	His	Asn	Phe
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	010	DCI	355	nsp	мгу	чэр	FIU	360		110	Deu	173	365	1111	Grà	Gru
10																
	Pro	Val	Ásp	Trp	Phe	Lys	Thr	Gln	Val	Glu	Thr	Ser	Cys	Thr	Tyr	Gly
		370					375					380				
15	Cly	Phe	T.011-	Cor	Cly	Cure	Phe	шрж	Gly	Gly	T.OU	<b>A</b> e <b>D</b>	Dho	Glm	vəl	Glu
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20																
	His	His	Leu	Phe	Pro	Arg	Met	Ser	Ser	Ala	Trp.	Tyr	Pro	Tyr	Ile	Ala
			-		405					410					415	
25	Pro	Lys	Val	Ara	Glu	Ile	Cvs	Ala	Lvs	His	Glv	Val	His	Tvr	Ala	Tvr
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			435					440					445	۰.		
35	Ala	Ala	Gly	Thr	Gly	Ala	Asn	Trp	Arg	Gln	Met	Ala	Arg	Glu	Asn	Pro
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	Gly	Ser	Ala	Ile	Thr	Thr	Tyr	Lys	Asn	Met	Asp	Ala	Thr	Thr	Val	Phe	
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	+~~		**~~		***		***		- المرجم		-	****	a++	a++	<b>a a</b>	+	<b>8</b> 1 <i>5</i>
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5																					
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					35					4	0			-		45			. ~		
40			:_ mi		<b>b</b> - 1	·	<b>1711a an</b>	<b>0</b> ]	0	<b>.</b>		٦ ٦	<b>1</b>	<b>m</b>	<b>a</b> 1	<b>—</b>	τ.		1	<u></u>	
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45		Le	eu L	vs L	ys (	Jlu	Cys	Pro	Thr	Gl	n G	lu F	ro	Glu	Ile	Pro	As	I az	le	Lvs	
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	Val	AId	τp	GIII	165	Ded	GIY	лтр	Leu	170	лıs	Gru	File	AIG	His 175	птр
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	21011	210	1120		, interest of the second secon	niu	215	ASH	Var	Vur	OTÀ	220	70P	CTÀ	105	Leu
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20	Val	Lys	Asp	Pro	Asp	Val	Arg	Arg	Val	Ala	Thr	Thr	Gln	Pro	Arg	Gln
				260					265					270		
25	Trp	Tyr	His	Ala	Tyr	Gln	His	Ile	Tyr	Leu	Ala	Val	Leu	Tyr	Gly	Thr
			275					280					285			
30	Leu	Ala	Leu	Lys	Ser	Ile	Phe	Leu	Asp	Asp	Phe	Leu	Ala	Tyr	Phe	Thr
		290					295					300				
	Gly	Ser	Ile	Gly	Pro	Val	Lys	Val	Ala	Lys	Met	Thr	Pro	Leu	Glu	Phe
35	- 305			-		310	-			-	315					320
	Acn	Tle	Phe	Phe	Gln	Gly	Lare	LOU	ī.eu	መነም	<b>دا</b> ۸	Dhe	መንም	Mot	Pho	Val
40	ASII	TTC	ETIC	r116	325	GIY	цуз	Deu	Deu	330	AIG	FIIe	ιyı	Mec	335	Var
	_	-	~		_	~ 7				- 1	-1					_
45	Leu	Pro	Ser	Val 340	lyr	GΙΥ	Vai	Hls	Ser 345	GIY	GIΫ	Thr	Phe	Leu 350	Ala	Leu
50	Tyr	Val	Ala 355	Ser	Gln	Leu	Ile	Thr 360	Gly	Trp	Met	Leu	Ala 365	Phe	Leu	Phe
			-										~ ~			
55	Gln	Val 370	Ala	His	Val	Val		Asp	Val	Ala	Phe		Thr	Pro	Glu	Gly
55	· .	ن ب _					375					380				

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	Gly	Lys	Val	Lys	Gly	Gly	Trp	Ala	Ala	Met	Gln	Val	Ala	Thr	Thr	Thr
5	385					390					395					400
	Asp	Phe	Ser	Pro	Arg 405	Ser	Trp	Phe	Trp	Gly 410	His	Val	Ser	Gly	Gly 415	Leu
10										410					410	
	Asn	Asn	Gln	Ile	Glu	His	His	Leu	Phe	Pro	Gly	Val	Cys	His	Val	His
15				420					425					430		
	Tyr	Pro	Ala	Ile	Gln	Pro	Ile	Val	Glu	Lys	Thr	Cys	Lys	Glu	Phe	Asp
20			435					440					445			
	Val	Pro	Tyr	Val	Ala	Tyr	Pro	Thr	Phe	Trp	Thr	Ala	Leu	Arg	Ala	His
25		450					455					460				
25											÷					
	Phe	Ala	His	Leu	Lys	Lys	Val	Gly	Leu	Thr	Glu	Phe	Arg	Leu	Asp	Gly
	465					470					475					480
30																

#### Claims

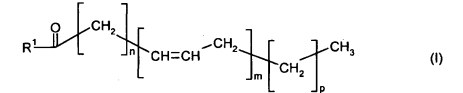
1. A process for the production of compounds of the following general formula I

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in transgenic oil producing plant with a content of at least 1 % by weight of said compounds referred to the total lipid content of said oil producing plant which comprises the following steps:

a) introduction of at least one nucleic acid in a transgenic oil producing plant, which encodes a  $\Delta$ -9-elongase having the sequence SEQ ID NO:3, and

b) introduction of at least one second nucleic acid which encodes a  $\Delta$ -8-desaturase having the sequence SEQ ID NO:1, and

c) if necessary introduction of at least a third nucleic acid, which encodes a  $\Delta$ -5-desaturase having the sequence SEQ ID NO:5, 7 or 9, and

d) cultivating and harvesting of said oil producing plant; and

⁵⁵ where the variables and substituents in formula I have the following meanings:

R¹ = hydroxyl-, Coenzyme A-(Thioester), phosphatidylcholine-, phosphatidyl-ethanolamine-, phosphatidylglycerol-, diphosphatidylglycerol-, phosphatidylserine-, phosphatidylinositol-, sphingolipid-, glycoshingolipid- or a

#### residue of the general formula II:

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$$H_{2}C-O-R^{2}$$

$$H_{2}C-O-R^{3}$$

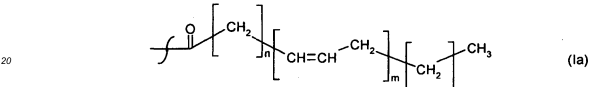
$$H_{2}C-O-f^{-}$$
(II)

 $R^2$  = hydrogen-, phosphatidylcholine-, phosphatidylethanolamine-, phosphatidylglycerol-, diphosphatidylglycerol-, phosphatidylserine-, phosphatidylinositol-, shingolipid-, glycoshingolipid-, glycoshingolipid- or saturated or unsaturated C₂-C₂₄-alkylcarbonyl-,

 $R^3$  = hydrogen-, saturated or unsaturated  $C_2$ - $C_{24}$ -alkylcarbonyl-, or  $R^2$  and  $R^3$  independent of each other a residue of the formula Ia:



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- 2. The process as claimed in claim 1, wherein the substituents R² and R³ are independent of each other saturated or unsaturated C₁₀-C₂₂-alkylcarbonyl-.
- The process as claimed in any of the claims 1 to 2, wherein the substituents R² and R³ are independent of each other saturated or unsaturated C₁₆-, C₁₈-, C₂₀- or C₂₂-alkyl-carbonyl-.
  - 4. The process as claimed in any of the claims 1 to 3, wherein the substituents R² and R³ are independent of each other unsaturated C₁₆-, C₁₈-, C₂₀- or C₂₂-alkylcarbonyl- with at least three double bonds.
- **5.** The process as claimed in any of the claims 1 to 4, wherein the transgenic plant is selected from the group consisting of rapeseed, poppy, mustard, hemp, castor bean, sesame, olive, calendula, punica, hazel nut, almond, macadamia, avocado, pumpkin, walnut, laurel, pistachio, primrose, canola, peanut, linseed, soybean, safflower, sunflower and borage.
- 40 **6.** The process as claimed in any of the claims 1 to 5, wherein the compounds of the general formula I are isolated in the form of their oils, lipids of free fatty acids.
  - 7. The process as claimed in any of the claims 1 to 6, wherein the compounds of the general formula I are isolated in a concentration of at least 5 % by weight referred to the total lipid content.

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- An isolated nucleic acid comprising a nucleotide sequence which encodes a ∆-8-desaturase having a nucleic acid sequence depicted in SEQ ID NO: 1.
- 9. polypeptide encoded by an isolated nucleic acid sequence as claimed in claims 8.
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- **10.** A gene construct comprising an isolated nucleic acid having the sequence SEQ ID NO: 1, where the nucleic acid is functionally linked to one or more regulatory signals.
- **11.** An oil producing plant comprising at least one nucleic acid as claimed in claim 8 or a gene construct as claimed in claim 10.

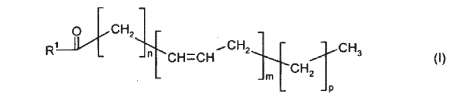
#### Patentansprüche

- 1. Verfahren zum Herstellen von Verbindungen der folgenden allgemeinen Formel I
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in einer transgenen ölproduzierenden Pflanze mit einem Gehalt von wenigstens 1 Gew.-% an den Verbindungen
 bezogen auf den Gesamtlipidgehalt der ölproduzierenden Pflanze, umfassend folgende Schritte:

a) Einführen wenigstens einer Nukleinsäure, die eine ∆-9-Elongase kodiert, mit der Sequenz SEQ ID NO:3 in eine transgene ölproduzierende Pflanze und

b) Einführen wenigstens einer zweiten Nukleinsäure, die eine ∆-8-Desaturase kodiert, mit der Sequenz SEQ ID NO:1 und

c) nötigenfalls Einführen wenigstens einer dritten Nukleinsäure, die eine ∆-5-Desaturase kodiert, mit der Sequenz SEQ ID NO:5, 7 oder 9 und

d) Kultivieren und Ernten der ölproduzierenden

Pflanze; und

wobei die Variablen und Substituenten in Formel I folgende Bedeutungen haben:

R¹ = Hydroxy-, Coenzym A- (Thioester),

Phosphatidylcholin-, Phosphatidyl-, Ethanolamin-, Phosphatidylglycerol-, Diphosphatidylglycerol-, Phosphati ³⁰ dylserin-, Phosphatidylinositol-, Sphingolipid-, Glycosphingolipid- oder ein Rest der allgemeinen Formel II:

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 $H_{2}C-O-R^{2}$   $HC-O-R^{3}$   $H_{2}C-O-f^{3}$  (II)

R² = Wasserstoff-, Phosphatidylcholin-,

Phosphatidylethanolamin-, Phosphatidylglycerol-, Diphosphatidylglycerol-, Phosphatidylserin-, Phosphatidylinositol-, Sphingolipid-, Glycosphingolipid-, Glycosphingolipid- oder gesättigtes oder nichtgesättigtes C₂-C₂₄-Alkylcarbonyl,

R³ = Wasserstoff-, gesättigtes oder

nichtgesättigtes C₂-C₂₄-Alkylcarbonyl-, oder R² und R³ sind unabhängig voneinander ein Rest der Formel Ia:

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n = 3, 4 oder 6, m = 3, 4 oder 5 und p = 0 oder 3.

- Verfahren gemäß Anspruch 1, wobei die Substituenten R² und R³ unabhängig voneinander gesättigtes oder nichtgesättigtes C₁₀-C₂₂-Alkylcarbonyl- sind.
  - 3. Verfahren gemäß einem der Ansprüche 1 bis 2, wobei die Substituenten R² und R³ unabhängig voneinander ge-

sättigtes oder nichtgesättigtes C16⁻, C18⁻, C20⁻oder C22⁻Alkylcarbonyl- sind.

- 4. Verfahren gemäß einem der Ansprüche 1 bis 3, wobei die Substituenten R² und R³ unabhängig voneinander nichtgesättigtes C₁₆-, C₁₈-, C₂₀- oder C₂₂-Alkylcarbonyl- mit wenigstens drei Doppelbindungen sind.
- 5. Verfahren gemäß einem der Ansprüche 1 bis 4, wobei die transgene Pflanze ausgewählt ist aus der Gruppe bestehend aus Raps, Mohn, Senf, Hanf, Castorbohne, Sesam, Olive, Calendula, Punica, Haselnuss, Mandel, Macadamia, Avocado, Kürbis, Walnuss, Lorbeer, Pistazie, Primel, Canola, Erdnuss, Leinsamen, Sojabohne, Distel, Sonnenblume und Borago.
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- 6. Verfahren gemäß einem der Ansprüche 1 bis 5, wobei die Verbindungen der allgemeinen Formel I in der Form ihrer Öle, Lipide von freien Fettsäuren isoliert werden.
- Verfahren gemäß einem der Ansprüche 1 bis 6, wobei die Verbindungen der allgemeinen Formel I in einer Konzentration von wenigstens 5 Gew.-% bezogen auf den Gesamtlipidgehalt isoliert werden.
  - 8. Isolierte Nukleinsäure, umfassend eine Nulkeotidsequenz, die eine ∆-8-Desaturase kodiert, mit der in SEQ ID NO:1 dargestellten Sequenz.
- **9.** Polypeptid, kodiert von einer isolierten Nukleinsäuresequenz gemäß Anspruch 8.
  - **10.** Genkonstrukt, umfassend eine isolierte Nukleinsäure mit der Sequenz SEQ ID NO:1, wobei die Nukleinsäure mit einem oder mehreren regulatorischen Signalen funktionell verbunden ist.
- ²⁵ **11.** Ölproduzierende Pflanze, umfassend wenigstens eine Nukleinsäure gemäß Anspruch 8 oder ein Genkonstrukt gemäß Anspruch 10.

#### Revendications

1. Procédé pour la production de composés de la formule générale suivante l :



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dans une plante oléagineuse transgénique ayant une teneur d'au moins 1 % en poids desdits composés par rapport à la teneur totale en lipides de ladite plante oléagineuse qui comprend les étapes suivantes :

a) introduction d'au moins un acide nucléique dans une plante oléagineuse transgénique qui code pour une ∆9-élongase ayant la séquence SEQ ID NO: 3, et

b) introduction d'au moins un deuxième activités qui code pour une ∆-8-désaturase ayant la séquence SEQ ID NO: 1, et

c) si nécessaire, introduction d'au moins un troisième acide nucléique, qui code pour une ∆-5-désaturase ayant la séquence SEQ ID NO: 5, 7 ou 9, et

d) culture et récolte de ladite plante oléagineuse ;

#### et

les variables et substituants dans la formule I ont les définitions suivantes :

⁵⁵ R¹ = hydroxyl-, coenzyme A-(thioester), phosphatidylcholine-, phosphatidyléthanolamine-, phosphatidylglycé rol-, diphosphatidylglycérol-, phosphatidylsérine-, phosphatidylinositol-, sphingolipide-, glycosphingolipide-, ou
 un résidu de formule générale II :

$$H_{2}C-O-R^{2}$$

$$HC-O-R^{3}$$

$$H_{2}C-O-f^{3}$$

$$(II)$$

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 $R^2$  = hydrogène-, phosphatidylcholine-, phosphatidyléthanolamine-, phosphatidylglycérol-, diphosphatidylglycérol-, phosphatidylsérine-, phosphatidylinositol-, sphingolipide-, glycosphingolipide- ou (alkyle en C₂-C₂₄)-carbonyle saturé ou insaturé,

 $R^3$  = hydrogène-, (alkyle en C₂-C₂₄)-carbonyle saturé ou insaturé, ou  $R^2$  et  $R^3$  indépendamment l'un de l'autre un résidu de formule la :

 $\int \left[ CH_{2} \right]_{n} \left[ CH=CH^{-CH_{2}} \right]_{m} \left[ CH_{2} \right]_{0} CH_{3}$ (ia)

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n = 3, 4 ou 6, m = 3, 4 ou 5 et p = 0 ou 3.

- Procédé selon la revendication 1, dans lequel les substituants R² et R³ sont indépendamment l'un de l'autre un (alkyle en C₁₀-C₂₂)-carbonyle saturé ou insaturé.
- Procédé selon l'une quelconque des revendications 1 à 2, dans lequel les substituants R² et R³ sont indépendamment l'un de l'autre un (alkyle en C₁₆, C₁₈, C₂₀ ou C₂₂)-carbonyle saturé ou insaturé.
- 4. Procédé selon l'une quelconque des revendications 1 à 3, dans lequel les substituants R² et R³ sont indépendamment l'un de l'autre un (alkyle en C₁₆, C₁₈, C₂₀ ou C₂₂)-carbonyle insaturé avec au moins trois doubles liaisons.

5. Procédé selon l'une quelconque des revendications 1 à 4, dans lequel la plante transgénique est choisie dans le groupe constitué du colza, du pavot, de la moutarde, du chanvre, du ricin, du sésame, de l'olive, du Calendula, du Punica, de la noisette, de la noix de macadamia, de l'avocat, de la citrouille, de la noix, du laurier, de la pistache, de l'onagre, du canola, de l'arachide, du lin, du soja, du carthame, du tournesol et de la bourrache.

- 6. Procédé selon l'une quelconque des revendications 1 à 5, dans lequel les composés de la formule générale I sont isolés sous la forme de leurs huiles, lipides ou acides gras libres.
- Procédé selon l'une quelconque des revendications 1 à 6, dans lequel les composés de la formule générale l sont isolés à une concentration d'au moins 5 % en poids par rapport à la teneur en lipide totale.
  - Acide nucléique isolé comprenant une séquence nucléotidique qui code pour une ∆-8-désaturase ayant une séquence d'acide nucléique décrite dans SEQ ID NO: 1.

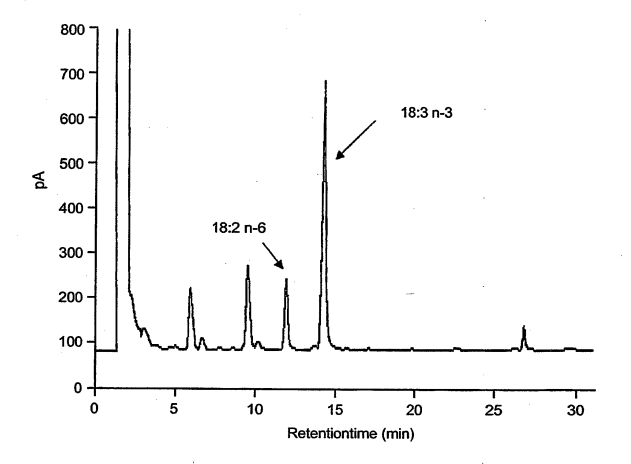
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- 9. Polypeptide codé par une séquence d'acide nucléique isolée selon la revendication 8.
- **10.** Construction génique comprenant un acide nucléique isolé ayant la séquence SEQ ID NO: 1, l'acide nucléique étant fonctionnellement lié à un ou plusieurs signaux régulateurs.

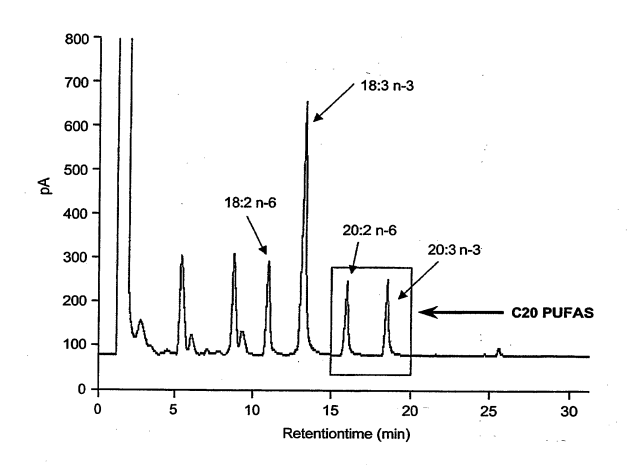
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**11.** Plante oléagineuse comprenant au moins un acide nucléique selon la revendication 8 ou une construction génique selon la revendication 10.

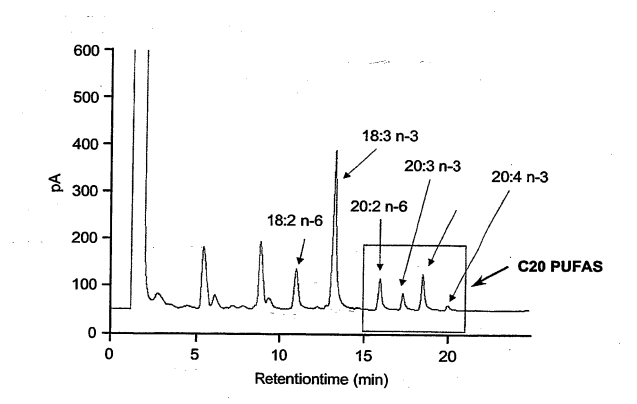
# FIG.1



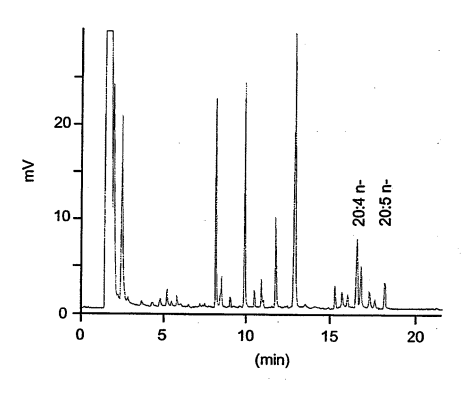
# FIG.2

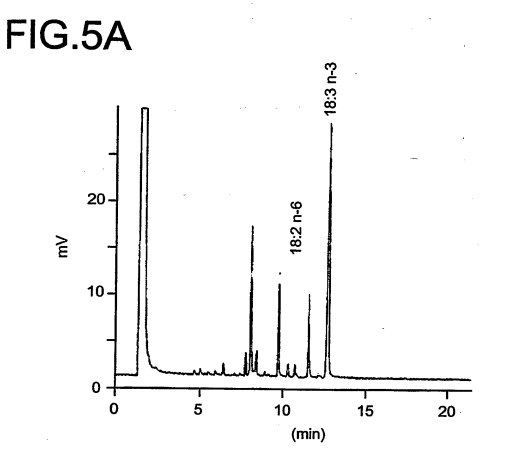


# FIG.3

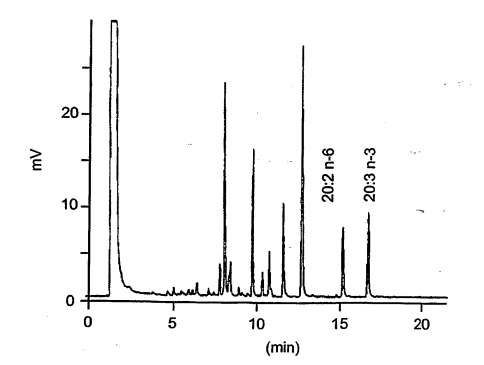


# FIG.4D











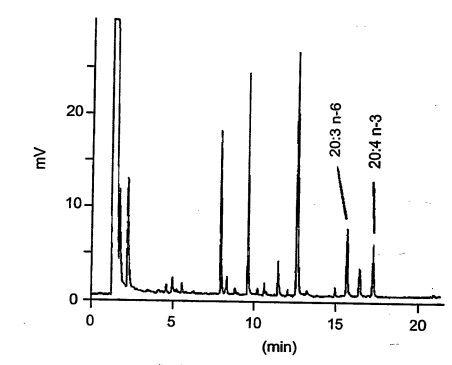
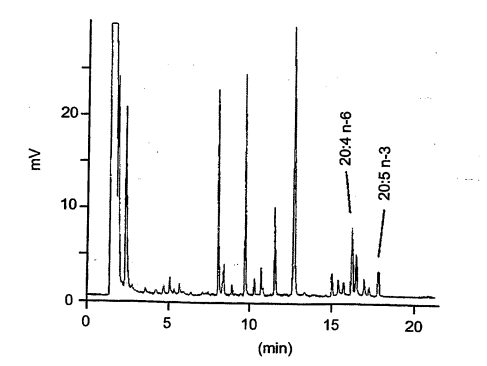


FIG.5D



## **REFERENCES CITED IN THE DESCRIPTION**

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