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- (54) **Hideg folyási tulajdonságot javító adalék állati vagy növényi eredetű tüzelőolajokhoz**

Az európai szabadalom ellen, megadásának az Európai Szabadalmi Közlönyben való meghirdetésétől számított kilenc hónapon belül, felszólalást lehet benyújtani az Európai Szabadalmi Hivatalnál. (Európai Szabadalmi Egyezmény 99. cikk(1))

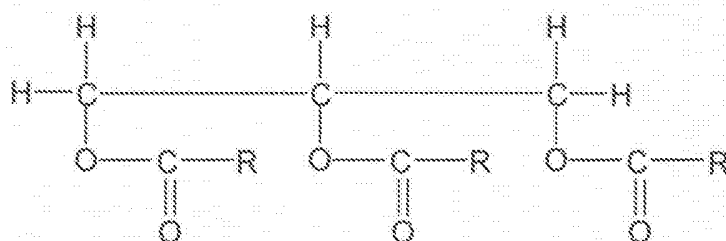
A fordítást a szabadalmas az 1995. évi XXXIII. törvény 84/H. §-a szerint nyújtotta be. A fordítás tartalmi helyességét a Szellemi Tulajdon Nemzeti Hivatala nem vizsgálta.

Cold flow improver for fuel oils of animal or vegetable origin

The present invention relates to an additive, to its use as a cold flow improver for vegetable or animal fuel oils and to correspondingly additized fuel oils.

In view of decreasing world crude oil reserves and the discussion about the environmentally damaging consequences of the use of fossil and mineral fuels, there is increasing interest in alternative energy sources based on renewable raw materials. These include in particular natural oils and fats of vegetable or animal origin. These are generally triglycerides of fatty acids having from 10 to 24 carbon atoms and a calorific value comparable to conventional fuels, but are at the same time regarded as being less harmful to the environment. Biofuels, i.e. fuels derived from animal or vegetable material, are obtained from renewable sources and, when they are combusted, generate only as much CO₂ as had previously been converted to biomass. It has been reported that less carbon dioxide is formed in the course of combustion than by the equivalent amount of crude oil distillate fuel, for example diesel fuel, and that very little sulfur dioxide is formed. In addition, they are biodegradable.

Oils obtained from animal or vegetable material are mainly metabolism products which include triglycerides of monocarboxylic acids, for example acids having from 10 to 25 carbon atoms, and corresponding to the formula



where R is an aliphatic radical which has from 10 to 25 carbon atoms and may be saturated or unsaturated.

In general, such oils contain glycerides from a series of acids whose number and type vary with the source of the oil, and they may additionally contain phosphoglycerides. Such oils can be obtained by processes known from the prior art.

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As a consequence of the sometimes unsatisfactory physical properties of the triglycerides, the industry has applied itself to converting the naturally occurring triglycerides to fatty acid esters of low alcohols such as methanol or ethanol.

10 A hindrance to the use of triglycerides and also of fatty acid esters of lower monohydric alcohols as a replacement for diesel fuel alone or in a mixture with diesel fuel has proven to be the flow behavior at low temperatures. The cause of this is the high uniformity of these oils in comparison to mineral oil middle distillates. For example, the rapeseed oil methyl ester (RME) has a Cold Filter Plugging Point (CFPP) of -14°C . It has hitherto been impossible using the prior art additives to
15 reliably obtain a CFPP value of -20°C required for use as a winter diesel in Central Europe, or of -22°C or lower for special applications. This problem is increased when oils are used which comprise relatively large amounts of the likewise readily available oils of sunflowers and soya.

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EP-B-0 665 873 discloses a fuel oil composition which comprises a biofuel, a fuel oil based on crude oil and an additive which comprises (a) an oil-soluble ethylene copolymer or (b) a comb polymer or (c) a polar nitrogen compound or (d) a compound in which at least one substantially linear alkyl group having from 10 to 30
25 carbon atoms is bonded to a nonpolymeric organic radical, in order to provide at least one linear chain of atoms which includes the carbon atoms of the alkyl groups and one or more nonterminal oxygen atoms, or (e) one or more of the components (a), (b), (c) and (d).

30 EP-B-0 629 231 discloses a composition which comprises a relatively large proportion of oil which consists substantially of alkyl esters of fatty acids which are derived from vegetable or animal oils or both, mixed with a small proportion of mineral oil cold flow improvers which comprises one or more of the following:

- (I) comb polymer, the copolymer (which may be esterified) of maleic anhydride or fumaric acid and another ethylenically unsaturated monomer, or polymer or copolymer of α -olefin, or fumarate or itaconate polymer or copolymer,
- 5 (II) polyoxyalkylene ester, ester/ether or a mixture thereof,
- (III) ethylene/unsaturated ester copolymer,
- (IV) polar, organic, nitrogen-containing paraffin crystal growth inhibitor,
- (V) hydrocarbon polymer,
- (VI) sulfur-carboxyl compounds and
- 10 (VII) aromatic pour point depressant modified with hydrocarbon radicals,

with the proviso that the composition comprises no mixtures of polymeric esters or copolymers of esters of acrylic and/or methacrylic acid which are derived from alcohols having from 1 to 22 carbon atoms.

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EP-B-0 543 356 discloses a process for preparing compositions having improved low temperature behavior for use as fuels or lubricants, starting from the esters of naturally occurring long-chain fatty acids with monohydric C₁-C₆-alcohols (FAE), which comprises

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- a) adding PPD additives (pour point depressants) known per se and used for improving the low temperature behavior of mineral oils in amounts of from 0.0001 to 10% by weight, based on the long-chain fatty acid esters FAE and
- b) cooling the nonadditized long-chain fatty acid esters FAE to a temperature
- 25 below the Cold Filter Plugging Point and
- c) removing the resulting precipitates (FAN).

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DE-A-40 40 317 discloses mixtures of fatty acid lower alkyl esters having improved cold stability comprising

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- a) from 58 to 95% by weight of at least one ester within the iodine number range from 50 to 150 and being derived from fatty acids having from 12 to 22 carbon atoms and lower aliphatic alcohols having from 1 to 4 carbon atoms,

- b) from 4 to 40% by weight of at least one ester of fatty acids having from 6 to 14 carbon atoms and lower aliphatic alcohols having from 1 to 4 carbon atoms and
- c) from 0.1 to 2% by weight of at least one polymeric ester.

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EP-B-0 153 176 discloses the use of polymers based on unsaturated dialkyl C₄-C₈-dicarboxylates having an average alkyl chain length of from 12 to 14 as cold flow improvers for certain crude oil distillate fuel oils. Mentioned as suitable comonomers are unsaturated esters, in particular vinyl acetate, but also α -olefins.

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EP-B-0 153 177 discloses an additive concentrate which comprises a combination of
 I) a copolymer having at least 25% by weight of an n-alkyl ester of a monoethylenically unsaturated C₄-C₈-mono- or -dicarboxylic acid, the average number of carbon atoms in the n-alkyl radicals being 12 – 14, and another
 15 unsaturated ester or an olefin, with
 II) another low temperature flow improver for distillate fuel oils.

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EP-A-1146108 teaches: the invention relates to additives for improving the cold flow properties of middle distillates, containing from 10 to 95% by weight of copolymers
 20 A), from 5 to 90% by weight of copolymers B) and, if required, from 0 to 70% by weight of copolymers C), which correspond to the following formulae:

A) copolymers of lower olefins and vinyl esters, containing

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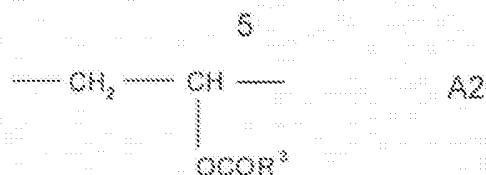
A1) from 85 to 97 mol% of bivalent structural units of the formula



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in which R¹ and R², independently of one another, are hydrogen or methyl, and

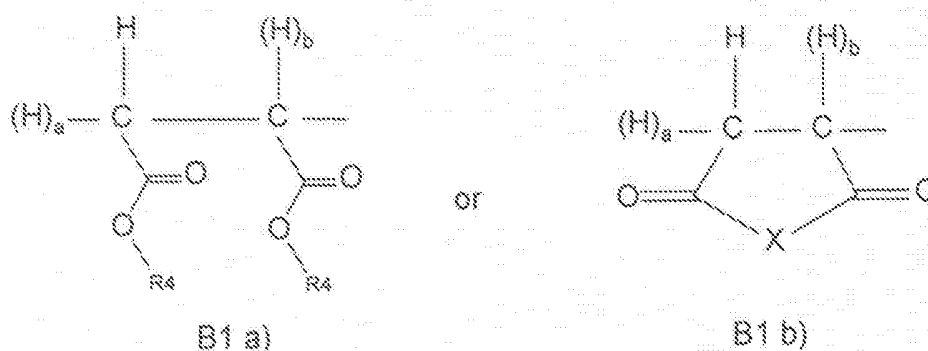
A2) at least 3 mol% of bivalent structural units of the formula



in which R^3 is saturated, branched C_6 - C_{15} -alkyl which has a tertiary carbon atom, wherein R^3 is bonded with its tertiary carbon atom to the carboxyl function,

B) copolymers comprising

B1) from 40 to 60 mol% of bivalent structural units of the formula



where X is O or N - R^4 and

in which a and b are 0 or 1 and $a + b = 1$, and

B2) from 60 to 40 mol% of bivalent structural units of the formula



and, if required,

B3) from 0 to 20 mol% of bivalent structural units which are derived from polyolefins, the polyolefins being derivable from monoolefins having 3 to 5 carbon atoms and wherein

- a) R^4 is an alkyl or alkenyl radical having 10 to 40 carbon atoms or an alkoxyalkyl radical having 1 to 100 alkoxy units and 1 to 30 carbon atoms in the alkyl radical and
- b) R^5 is an alkyl radical having 10 to 50 carbon atoms and
- c) the number of carbon atoms of the polyolefin molecules on which the structural units B3) are based is from 35 to 350, and, if required,

C) a further copolymer differing from A) and B) and comprising ethylene and one or more vinyl esters or acrylates, which by itself is effective as a cold flow improver for middle distillates.

US-2003/0163951 teaches multifunctional cold temperature additives comprising copolymers of dicarboxylic acid derivatives and olefins, onto which have been grafted nitrogen-containing compounds or esters.

US-5391632 teaches terpolymers based on α,β -unsaturated dicarboxylic anhydrides, α,β -unsaturated compounds and polyoxyalkylene ethers as claimed in claim 1 of US-5391632, and a process for preparation thereof as claimed in claim 5 of US-5391632.

The invention likewise provides a mixture of at least one terpolymer as claimed in claim 1 and at least one ethylene-vinyl ester copolymer. The terpolymers according to the invention, and mixtures thereof, are used as paraffin inhibitors in crude oils and mineral oil products.

US-4985048 teaches polymer mixtures of a copolymer (A_1) composed of 10-60% by weight of vinyl acetate and 40-90% by weight of ethylene or a copolymer (A_2) composed of 15-50% by weight of vinyl acetate, 0.5-20% by weight of C_5 - C_{24} - α -olefin and 30-70% by weight of ethylene and a copolymer (B) composed of 10-90% by weight of C_6 - C_{24} - α -olefin and 10-90% by weight of N- C_5 - C_{12} -alkylmaleimide, where the mixing ratio of the copolymers (A_1) or (A_2) to (B) is 100:1 to 1:1. These polymer mixtures are used as flow improvers in mineral oil distillates.

WO 95/22300 (= EP 0 746 598) discloses comb polymers in which the alkyl radicals have an average of less than 12 carbon atoms. These additives are especially suitable for oils having cloud points of less than -10°C , although the oils may also be native hydrocarbon oils (page 21, line 16 ff.). However, native oils have cloud points of about -2°C upward.

It has hitherto often been impossible using the existing additives to reliably adjust fatty acid esters to a CFPP value of -20°C required for use as a winter diesel in Central Europe or of -22°C and lower for special applications. An additional problem with the existing additives is the lacking cold temperature change stability of the additized oils, i.e. the CFPP value of the oils attained rises gradually when the oil is stored for a prolonged period at changing temperatures in the region of the cloud point or below.

It is therefore an object of the invention to provide additives for improving the cold flow behavior of fatty acid esters which are derived, for example, from rapeseed oil, sunflower oil and/or soya oil and attain CFPP values of -20°C and below which remain constant even when the oil is stored for a prolonged period in the region of its cloud point or below.

It has now been found that, surprisingly, an additive comprising ethylene copolymers and comb polymers is an excellent flow improver for such fatty acid esters.

The invention therefore provides an additive comprising the constituents A and B in a weight ratio of from 10 : 1 to 1 : 10

- A) a copolymer of ethylene and 8 - 21 mol% of at least one acrylic or vinyl ester having a $\text{C}_1\text{-C}_{18}$ -alkyl radical and
- B) a comb polymer containing structural units of

- B1) at least one olefin as monomer 1, which bears at least one $\text{C}_8\text{-C}_{18}$ -alkyl radical on the olefinic double bond, and
- B2) at least one ethylenically unsaturated dicarboxylic acid as monomer 2,

which bears at least one C₈-C₁₆-alkyl radical bonded via an amide and/or imide moiety,

wherein the sum Q

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$$Q = \sum_i w_{1i} \cdot n_{1i} + \sum_j w_{2j} \cdot n_{2j}$$

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of the molar averages of the carbon chain length distributions in the alkyl radicals of monomer 1 on the one hand and the alkyl radicals of the amide and/or imide groups of monomer 2 on the other hand is from 23 to 27, in a molar B1):B2) ratio of from 1.5 : 1 to 1: 1.5 where

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w_1 is the molar proportion of the individual chain lengths in the alkyl radicals of monomer 1,

w_2 is the molar proportion of the individual chain lengths in the alkyl radicals of the amide and/or imide groups of monomer 2,

n_1 are the individual chain lengths in the alkyl radicals of monomer 1,

n_2 are the individual chain lengths in the alkyl radicals of the amide and/or imide groups of monomer 2,

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i is the serial variable for the individual chain lengths in the alkyl radicals of monomer 1, and

j is the serial variable for the individual chain lengths in the alkyl radicals of the amide and/or imide groups of monomer 2.

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The invention further provides a fuel oil composition comprising a fuel oil of animal or vegetable origin and from 0.001 to 5% by weight of the above-defined additive.

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The invention further provides the use of from 0.001 to 5% by weight of the above-defined additive for improving the cold flow properties of fuel oils of animal or vegetable origin.

The invention further provides a process for improving the cold flow properties of fuel oils of animal or vegetable origin by adding the above-defined additive in an amount of from 0.001 to 5% by weight to fuel oils of animal or vegetable origin.

In a preferred embodiment of the invention, Q has values of from 24 to 26.

Here, side chain length of olefins refers to the alkyl radical branching from the polymer backbone, i.e. the chain length of the monomeric olefin minus the two olefinically bonded carbon atoms. In the case of olefins having nonterminal double bonds, for example olefins having vinylidene moiety, it is correspondingly the total chain length of the olefin minus the double bond merging into the polymer backbone that has to be taken into account.

Useful ethylene copolymers A) are those which contain from 8 to 21 mol% of one or more vinyl and/or (meth)acrylic ester and from 79 to 92 mol% of ethylene. Particular preference is given to ethylene copolymers having from 10 to 18 mol% and especially from 12 to 16 mol%, of at least one vinyl ester. Suitable vinyl esters are derived from fatty acids having linear or branched alkyl groups having from 1 to 18 carbon atoms and preferably from 1 to 12 carbon atoms. Examples include vinyl acetate, vinyl propionate, vinyl butyrate, vinyl hexanoate, vinyl heptanoate, vinyl octanoate, vinyl laurate and vinyl stearate, and also esters of vinyl alcohol based on branched fatty acids, such as vinyl isobutyrate, vinyl pivalate, vinyl 2-ethylhexanoate, vinyl isononanoate, vinyl neononanoate, vinyl neodecanoate and vinyl neoundecanoate. Particular preference is given to vinyl acetate. Likewise suitable as comonomers are esters of acrylic and methacrylic acids having from 1 to 20 carbon atoms in the alkyl radical, such as methyl (meth)acrylate, ethyl (meth)acrylate, propyl (meth)acrylate, n- and isobutyl (meth)acrylate, and hexyl, octyl, 2-ethylhexyl, decyl, dodecyl, tetradecyl, hexadecyl and octadecyl (meth)acrylate, and also mixtures of two, three, four or else more of these comonomers.

Apart from ethylene, particularly preferred terpolymers of vinyl 2-ethylhexanoate, of vinyl neononanoate or of vinyl neodecanoate contain preferably from 3.5 to 20 mol%, in particular from 8 to 15 mol%, of vinyl acetate, and from 0.1 to 12 mol%, in particular from 0.2 to 5 mol%, of at least one long-chain in addition to ethylene and from 8 to 18 mol% of vinyl esters, additionally contain from 0.5 to 10 mol% of olefins such as propene, butene, isobutylene, hexene, 4-methylpentene, octene,

diisobutylene and/or norbornene.

The copolymers A preferably have molecular weights which correspond to melt viscosities at 140°C of from 20 to 10 000 mPas, in particular from 30 to 5000 mPas, and especially from 50 to 1000 mPas. The degrees of branching determined by means of ¹H NMR spectroscopy are preferably between 1 and 9 CH₃/100 CH₂ groups, in particular between 2 and 6 CH₃/100 CH₂ groups, for example from 2.5 to 5 CH₃/100 CH₂ groups, which do not stem from the comonomers.

- 10 The copolymers (A) can be prepared by the customary copolymerization processes, for example suspension polymerization, solution polymerization, gas phase polymerization or high pressure bulk polymerization. Preference is given to carrying out the high pressure bulk polymerization at pressures of from 50 to 400 MPa, preferably from 100 to 300 MPa, and temperatures from 100 to 300°C, preferably from 150 to 220°C. In a particularly preferred preparation variant, the polymerization is effected in a multizone reactor in which the temperature difference between the peroxide feeds along the tubular reactor is kept very low, i.e. < 50°C, preferably < 30°C, in particular < 15°C. The temperature maxima in the individual reaction zones preferably differ by less than 30°C, more preferably by less than 20°C and especially by less than 10°C.

- The reaction of the monomers is initiated by radical-forming initiators (radical chain initiators). This substance class includes, for example, oxygen, hydroperoxides, peroxides and azo compounds, such as cumene hydroperoxide, t-butyl hydroperoxide, dilauroyl peroxide, dibenzoyl peroxide, bis(2-ethylhexyl) peroxydicarbonate, t-butyl perpivalate, t-butyl permaleate, t-butyl perbenzoate, dicumyl peroxide, t-butyl cumyl peroxide, di(t-butyl) peroxide, 2,2'-azobis(2-methylpropanonitrile), 2,2'-azobis(2-methylbutyronitrile). The initiators are used individually or as a mixture of two or more substances in amounts of from 0.01 to 20% by weight, preferably from 0.05 to 10% by weight, based on the monomer mixture.

The high pressure bulk polymerization is carried out in known high pressure

reactors, for example autoclaves or tubular reactors, batchwise or continuously, and tubular reactors have proven particularly useful. Solvents such as aliphatic and/or aromatic hydrocarbons or hydrocarbon mixtures, benzene or toluene may be present in the reaction mixture. Preference is given to the substantially solvent-free procedure. In a preferred embodiment of the polymerization, the mixture of the monomers, the initiator and, if used, the moderator, are fed to a tubular reactor via the reactor entrance and also via one or more side branches. Preferred moderators are, for example, hydrogen, saturated and unsaturated hydrocarbons, for example propane or propene, aldehydes, for example propionaldehyde, n-butyraldehyde or isobutyraldehyde, ketones, for example acetone, methyl ethyl ketone, methyl isobutyl ketone, cyclohexanone, and alcohols, for example butanol. The comonomers and also the moderators may be metered into the reactor either together with ethylene or else separately via sidestreams. The monomer streams may have different compositions (EP-A-0 271 738 and EP-A-0 922 716).

Examples of suitable co- or terpolymers include:

ethylene-vinyl acetate copolymers having 10 - 40% by weight of vinyl acetate and 60 - 90% by weight of ethylene;

the ethylene-vinyl acetate-hexene terpolymers known from DE-A-34 43 475;

the ethylene-vinyl acetate-diisobutylene terpolymers described in EP-B-0 203 554;

the mixture of an ethylene-vinyl acetate-diisobutylene terpolymer and an

ethylene/vinyl acetate copolymer known from EP-B-0 254 284;

the mixtures of an ethylene-vinyl acetate copolymer and an ethylene-vinyl acetate-N-vinylpyrrolidone terpolymer disclosed in EP-B-0 405 270;

the ethylene/vinyl acetate/isobutyl vinyl ether terpolymers described in EP-B-0 463 518;

the ethylene/vinyl acetate/neononanoate or -vinyl neodecanoate terpolymers which,

apart from ethylene, contain 10 - 35% by weight of vinyl acetate and 1 - 25% by weight of the particular neo compound, known from EP-B-0 493 769;

the terpolymers of ethylene, a first vinyl ester having up to 4 carbon atoms and a
5 second vinyl ester which is derived from a branched carboxylic acid having up to 7 carbon atoms or a branched but nontertiary carboxylic acid having from 8 to 15 carbon atoms, described in EP 0778875;

the terpolymers of ethylene, the vinyl ester of one or more aliphatic C₂- to C₂₀-
10 monocarboxylic acids and 4-methylpentene-1, described in DE-A-196 20 118;

the terpolymers of ethylene, the vinyl ester of one or more aliphatic C₂- to C₂₀-
monocarboxylic acids and bicyclo[2.2.1]hept-2-ene, disclosed in DE-A-196 20 119;

the terpolymers of ethylene and at least one olefinically unsaturated comonomer
15 which contains one or more hydroxyl groups, described in EP-A-0 926 168.

Preference is given to using mixtures of the same or different ethylene copolymers. The polymers on which the mixtures are based more preferably differ in at least one
20 characteristic. For example, they may contain different comonomers, different comonomer contents, molecular weights and/or degrees of branching. The mixing ratio of the different ethylene copolymers is preferably between 20:1 and 1:20, preferably from 10:1 to 1:10, in particular from 5:1 to 1:5.

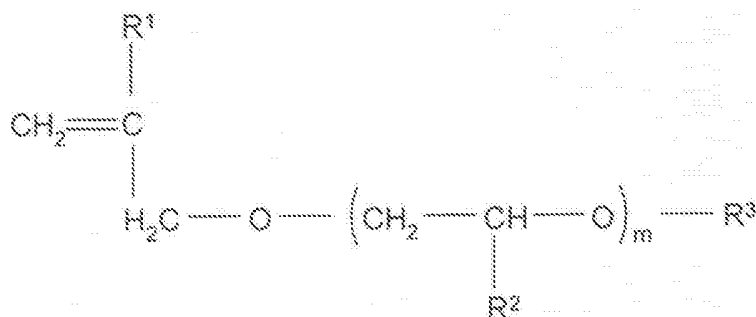
The copolymers B are preferably derived from ethylenically unsaturated dicarboxylic
25 acids and their derivatives such as esters and anhydrides. Preference is given to maleic acid, fumaric acid, itaconic acid and the esters thereof with lower alcohols having from 1 to 6 carbon atoms and also anhydrides thereof, for example maleic anhydride. Particularly suitable comonomers are monoolefins having from 10 to 20, in particular having from 12 to 18, carbon atoms. These are preferably linear and the
30 double bond is dicarboxylic acids and their derivatives such as esters and anhydrides. Preference is given to maleic acid, fumaric acid, itaconic acid and the esters thereof with lower alcohols having from 1 to 6 carbon atoms and also anhydrides thereof, for example maleic anhydride. Particularly suitable comonomers

are monoolefins having from 10 to 20, in particular having from 12 to 18, carbon atoms. These are preferably linear and the double bond is preferably terminal as, for example, in dodecene, tridecene, tetradecene, pentadecene, hexadecene, heptadecene and octadecene. The ratio of dicarboxylic acid or dicarboxylic acid derivative to olefin or olefins in the polymer is in the range from 1: 1.5 to 1.5:1, and it is especially equimolar.

Also present in copolymer B may be minor amounts of up to 20 mol%, preferably <10 mol%, especially <5 mol%, of further comonomers which are copolymerizable with ethylenically unsaturated dicarboxylic acids and the olefins specified, for example relatively short- and relatively long-chain olefins, allyl polyglycol ethers, C₁-C₃₀-alkyl (meth)acrylates, vinylaromatics or C₁-C₂₀-alkyl vinyl ethers.

Poly(isobutylene) having a molecular weight up to 5000 g/mol are likewise used in minor amounts, and preference is given to highly reactive variants having a high proportion of terminal vinylidene groups. These further comonomers are not taken into account in the calculation of the factor Q determining the effectiveness.

Alkyl polyglycol ethers correspond to the general formula



where

R¹ is hydrogen or methyl,

R² is hydrogen or C₁-C₄-alkyl,

m is a number from 1 to 100,

170°C. The preferred preparative process is the solvent-free bulk polymerization, although it is also possible to carry out the polymerization in the presence of aprotic solvents such as benzene, toluene, xylene or of relatively high-boiling aromatic, aliphatic or isoaliphatic solvents or solvent mixtures, such as kerosene or Solvent Naphtha. Particular preference is given to the polymerization in aliphatic or isoaliphatic solvents having little moderating influence. The proportion of solvent in the polymerization mixture is generally between 10 and 90% by weight, preferably between 35 and 60% by weight. In the case of the solution polymerization, the reaction temperature can be set in a particularly simple manner via the boiling point of the solvent or by working under reduced or elevated pressure.

The average molecular mass of the inventive copolymers B is generally between 1200 and 200 000 g/mol, in particular between 2000 and 100 000 g/mol, measured by means of gel permeation chromatography (GPC) against polystyrene standards in THF. Inventive copolymers B have to be oil-soluble in the dosages relevant to the practice, i.e. they have to dissolve without residue at 50°C in the oil to be additized.

The reaction of the monomers is initiated by radical-forming initiators (radical chain initiators). This substance class includes, for example, oxygen, hydroperoxides and peroxides such as cumene hydroperoxide, t-butyl hydroperoxide, dilauroyl peroxide, dibenzoyl peroxide, bis(2-ethylhexyl) peroxydicarbonate, t-butyl perpivalate, t-butyl permaleate, t-butyl perbenzoate, dicumyl peroxide, t-butyl cumyl peroxide, di(t-butyl) peroxide, and azo compounds such as 2,2'-azobis(2-methylpropanonitrile) or 2,2'-azobis(2-methylbutyronitrile). The initiators are used individually or as a mixture of two or more substances in amounts of from 0.01 to 20% by weight, preferably from 0.05 to 10% by weight, based on the monomer mixture.

The copolymers can be prepared either by reaction of maleic acid, fumaric acid and/or itaconic acid or the derivatives thereof with the appropriate amine and subsequent copolymerization or by copolymerization of olefin or olefins with at least one unsaturated dicarboxylic acid or a derivative thereof, for example itaconic anhydride and/or maleic anhydride, and subsequent reaction with amines.

Preference is given to carrying out a copolymerization with anhydrides and converting the resultant copolymer after the preparation to an amide and/or an imide.

5 In both cases, the reaction with amines is effected, for example, by reacting with from 0.8 to 2.5 mol of amine per mole of anhydride, preferably with from 1.0 to 2.0 mol of amine per mole of anhydride, at from 50 to 300°C. When approx. 1 mol of amine is used per mole of anhydride, monoamides which additionally bear one carboxyl group per amide group are formed preferentially at reaction temperatures of
10 from approx. 50 to 100°C. At higher reaction temperatures of from approx. 100 to 250°C, amides are formed preferentially from primary amines with elimination of water. When relatively large amounts of amine are used, preferably 2 mol of amine per mole of anhydride, amide ammonium salts are formed at from approx. 50 to 200°C and diamides are formed at higher temperatures of, for example, 100-300°C,
15 preferably 120-250°C. The water of reaction can be distilled off by means of an inert gas stream or removed by means of azeotropic distillation in the presence of an organic solvent. For this purpose, preference is given to using 20-80% by weight, in particular 30-70% by weight, especially 35-55% by weight, of at least one organic solvent. Useful monoamides are copolymers (50% dilution in solvent) having acid
20 numbers of 30 - 70 mg of KOH/g, preferably 40 - 60 mg of KOH/g. Corresponding copolymers having acid numbers of less than 40 mg of KOH/g, especially less than 30 mg of KOH/g, are considered as diamides or imides. Particular preference is given to monoamides and imides.

25 Suitable amines are primary and secondary amines having one or two C₈-C₁₆-alkyl radicals. They may bear one, two or three amino groups which are bonded via alkylene radicals having two or three carbon atoms. Preference is given to monoamines. In particular, they bear linear alkyl radicals, but they may also contain minor amounts, for example up to 30% by weight, preferably up to 20% by weight
30 and especially up to 10% by weight of (1- or 2-)branched amines. Either shorter- or longer-chain amines may be used, but their proportion is preferably below 20 mol% and especially below 10 mol%, for example between 1 and 5 mol%, based on the total amount of amines used.

Particularly preferred primary amines are octylamine, 2-ethylhexylamine, decylamine, undecylamine, dodecylamine, n-tridecylamine, isotridecylamine, tetradecylamine, pentadecylamine, hexadecylamine and mixtures thereof.

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Preferred secondary amines are dioctylamine, dinonylamine, didecylamine, didodecylamine, ditetradecylamine, dihexadecylamine, and also amines having different alkyl chain lengths, for example N-octyl-N-decylamine, N-decyl-N-dodecylamine, N-decyl-N-tetradecylamine, N-decyl-N-hexadecylamine, N-dodecyl-N-tetradecylamine, N-dodecyl-N-hexadecylamine, N-tetradecyl-N-hexadecylamine.

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Also suitable in accordance with the invention are secondary amines which, in addition to a C_8 - C_{16} -alkyl radical, bear shorter side chains having from 1 to 5 carbon atoms, for example methyl or ethyl groups. In the case of secondary amines, it is the average of the alkyl chain lengths of from C_8 to C_{16} that is taken into account as the alkyl chain length n for the calculation of the Q factor. Neither shorter nor longer alkyl radicals, where present, are taken into account in the calculation, since they do not contribute to the effectiveness of the additives.

15

Particularly preferred copolymers B are monoamides and imides of primary monoamines.

20

The use of mixtures of different olefins in the polymerization and mixtures of different amines in the amidation or imidation allows the effectiveness to be further adapted to specific fatty acid ester compositions.

25

In a preferred embodiment, the additives, in addition to constituents A and B, may also comprise polymers and copolymers based on C_{10} - C_{24} -alkyl acrylates or methacrylates (constituent C). These poly(alkyl acrylates) and methacrylates have molecular weights of from 800 to 1 000 000 g/mol and are preferably derived from caprylic alcohol, caproic alcohol, undecyl alcohol, lauryl alcohol, myristyl alcohol, cetyl alcohol, palmitoleyl alcohol, stearyl alcohol or mixtures thereof, for example coconut alcohol, palm alcohol, tallow fatty alcohol or behenyl alcohol.

30

In a preferred embodiment, mixtures of the copolymers B according to the invention are used, with the proviso that the mean of the Q values of the mixing components in turn assumes values of from 23 to 27 and preferably values from 24 to 26.

- 5 The mixing ratio of the additives A and B according to the invention is (in parts by weight) from 20:1 to 1:20, preferably from 10:1 to 1:10, in particular from 5:1 to 1:2. The proportion of component C in the formulations of A, B and C may be up to 40% by weight; it is preferably less than 20% by weight, in particular between 1 and 10% by weight.

10

The additives according to the invention are added to oils in amounts of from 0.001 to 5% by weight, preferably from 0.005 to 1% by weight and especially from 0.01 to 0.5% by weight. They may be used as such or else dissolved or dispersed in solvents, for example aliphatic and/or aromatic hydrocarbons or hydrocarbon

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mixtures, for example toluene, xylene, ethylbenzene, decane, pentadecane, petroleum fractions, kerosene, naphtha, diesel, heating oil, isoparaffins or commercial solvent mixtures such as Solvent Naphtha, ®Shellsol AB, ®Solvesso 150, ®Solvesso 200, ®Exxsol, ®Isopar and ®Shellsol D types. They are preferably dissolved in fuel oil of animal or vegetable origin based on fatty acid alkyl esters. The additives according to the invention preferably comprise 1 - 80%, especially 10 - 70%, in particular 25 - 60%, of solvent.

20

In a preferred embodiment, the fuel oil, which is frequently also referred to as biodiesel or biofuel, is a fatty acid alkyl ester made from fatty acids having from 12 to 24 carbon atoms and alcohols having from 1 to 4 carbon atoms. Typically, a relatively large portion of the fatty acids contains one, two or three double bonds.

25

Examples of oils which are derived from animal or vegetable material and in which the additive according to the invention can be used are rapeseed oil, coriander oil, soya oil, cottonseed oil, sunflower oil, castor oil, olive oil, peanut oil, maize oil, almond oil, palmseed oil, coconut oil, mustardseed oil, bovine tallow, bone oil, fish oils and used cooking oils. Further examples include oils which are derived from wheat, jute, sesame, shea tree nut, arachis oil and linseed oil. The fatty acid alkyl

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esters also referred to as biodiesel can be derived from these oils by processes known from the prior art. Rapeseed oil, which is a mixture of fatty acids partially esterified with glycerol, is preferred, since it is obtainable in large amounts and is obtainable in a simple manner by extractive pressing of rapeseeds. In addition,
5 preference is given to the likewise widely available oils of sunflowers and soya, and also to their mixtures with rapeseed oil.

Particularly suitable biofuels are low alkyl esters of fatty acids. These include, for example, commercially available mixtures of the ethyl, propyl, butyl and in particular
10 methyl esters of fatty acids having from 14 to 22 carbon atoms, for example of lauric acid, myristic acid, palmific acid, palmitolic acid, stearic acid, oleic acid, elaidic acid, petroselic acid, ricinolic acid, elaeostearic acid, linolic acid, linolenic acid, eicosanoic acid, gadoleic acid, docosanoic acid or erucic acid, each of which preferably has an iodine number of from 50 to 150, in particular from 90 to 125. Mixtures having
15 particularly advantageous properties are those which comprise mainly, i.e. comprise at least 50% by weight, methyl esters of fatty acids having from 16 to 22 carbon atoms, and 1, 2 or 3 double bonds. The preferred lower alkyl esters of fatty acids are the methyl esters of oleic acid, linoleic acid, linolenic acid and erucic acid.

Commercial mixtures of the type mentioned are obtained, for example, by
20 hydrolyzing and esterifying or by transesterifying animal and vegetable fats and oils with lower aliphatic alcohols. Equally suitable as starting materials are used cooking oils. To prepare lower alkyl esters of fatty acids, it is advantageous to start from fats and oils having a high iodine number, for example sunflower oil, rapeseed oil,
25 coriander oil, castor oil, soya oil, cottonseed oil, peanut oil or bovine tallow. Preference is given to lower alkyl esters of fatty acids based on a novel type of rapeseed oil, more than 80% by weight of whose fatty acid component is derived from unsaturated fatty acids having 18 carbon atoms.

A biofuel is therefore an oil which is obtained from vegetable or animal material or
30 both or a derivative thereof which can be used as a fuel and in particular as a diesel or heating oil. Although many of the above oils can be used as biofuels, preference is given to vegetable oil derivatives, and particularly preferred biofuels are alkyl ester

derivatives of rapeseed oil, cottonseed oil, soya oil, sunflower oil, olive oil or palm oil, and very particular preference is given to rapeseed oil methyl ester, sunflower oil methyl ester and soya oil methyl ester. Particularly preferred biofuels or components in the biofuel are additionally also used fatty acid esters, for example used fatty acid methyl esters.

The additive can be introduced into the oil to be additized in accordance with prior art processes. When more than one additive component or coadditive component is to be used, such components can be introduced into the oil together or separately in any desired combination.

The additives according to the invention allow the CFPP value of biodiesel to be adjusted to values of below -20°C and sometimes to values of below -25°C , as required for provision on the market for use in winter in particular. Equally, the pour point of biodiesel is reduced by the addition of the inventive additives. The inventive additives are particularly advantageous in problematic oils which contain a high proportion of esters of saturated fatty acids of more than 4%, in particular of more than 5% and especially having from 7 to 25%, for example having from 8 to 20%, as present, for example, in oils from sunflowers and soya. Such oils are characterized by cloud points of above -5°C and especially of above -3°C . It is thus also possible using the inventive additives to adjust mixtures of rapeseed oil methyl ester and sunflower and/or soya oil fatty acid methyl ester to CFPP values of -20°C and below. In addition, the oils additized in this way have a good cold temperature change stability, i.e. the CFPP value remains constant even on storage under winter conditions.

To prepare additive packages for specific solutions to problems, the additives according to the invention can also be used together with one or more oil-soluble coadditives which alone improve the cold flow properties of crude oils, lubricant oils or fuel oils. Examples of such coadditives are polar compounds which effect paraffin dispersion (paraffin dispersants) and also oil-soluble amphiphiles with the proviso that they differ from the comb polymers B.

The additives according to the invention can be used in a mixture with paraffin dispersants. Paraffin dispersants reduce the size of the paraffin crystals and have the effect that the paraffin particles do not separate but remain dispersed colloiddally with a distinctly reduced tendency to sedimentation. Useful paraffin dispersants have proven to be both low molecular weight and polymeric oil-soluble compounds having ionic or polar groups, for example amine salts and/or amides. Particularly preferred paraffin dispersants comprise reaction products of secondary fatty amines having from 20 to 44 carbon atoms, in particular dicocoamine, ditallow fat amine, distearylamine and dibehenylamine with carboxylic acids and derivatives thereof.

Paraffin dispersants which are obtained by reacting aliphatic or aromatic amines, preferably long-chain aliphatic amines, with aliphatic or aromatic mono-, di-, tri- or tetracarboxylic acids or their anhydrides (cf. US 4 211 534) have proven particularly useful. Equally suitable as paraffin dispersants are amides and ammonium salts of aminoalkylene polycarboxylic acids such as nitrilotriacetic acid or ethylenediamine-tetraacetic acid with secondary amines (cf. EP 0 398 101). Other paraffin dispersants are copolymers of maleic anhydride and α,β -unsaturated compounds which may optionally be reacted with primary monoalkylamines and/or aliphatic alcohols (cf. EP 0 154 177) and the reaction products of alkenyl-spiro-bislactones with amines (cf. EP 0 413 279 B1) and, according to EP 0 606 055 A2, reaction products of terpolymers based on α,β -unsaturated dicarboxylic anhydrides, α,β -unsaturated compounds and polyoxyalkylene ethers of lower unsaturated alcohols.

The mixing ratio (in parts by weight) of the additives according to the invention with paraffin dispersants is from 1:10 to 20:1, preferably from 1:1 bis 10:1.

The additives can be used alone or else together with other additives, for example with other pour point depressants or dewaxing assistants, with antioxidants, cetane number improvers, dehazers, demulsifiers, detergents, dispersants, defoamers, dyes, corrosion inhibitors, conductivity improvers, sludge inhibitors, odorants and/or additives for reducing the cloud point.

Examples

Characterization of the test oils:

The CFPP value is determined to EN 116 and the cloud point is determined to ISO 3015.

Table 1: Characterization of the test oils used

Oil No.		CP	CFPP
E 1	Rapeseed oil methyl ester	-2.3	-14°C
E 2	80% of rapeseed oil methyl ester + 20% of sunflower oil methyl ester	-1.6	-10°C
E 3	90% of rapeseed oil methyl ester + 10% of soya oil methyl ester	-2.0	-8°C

Table 2: Carbon chain distribution of the fatty acid methyl esters used to prepare the test oils (main constituents; area% by GC):

	C ₁₆	C ₁₆ '	C ₁₈	C ₁₈ '	C ₁₈ "	C ₁₈ '''	C ₂₀	C ₂₀ '	C ₂₂	Σ saturated
RME	4.4	0.4	1.6	57.8	21.6	8.8	1.5	0.7	0.2	7.7
SFME	6.0	0.1	3.8	28.7	58.7	0.1	0.3	0.3	0.7	10.8
SoyaME	10.4	0.1	4.1	24.8	51.3	6.9	0.5	0.4	0.4	15.4

RME = rapeseed oil methyl ester; SFME = sunflower oil methyl ester

SoyaME = soya oil methyl ester

The following additives were used:

Ethylene copolymers A

The ethylene copolymers used are commercial products having the characteristics specified in Table 2. The products were used as 65% or 50% (A3) dilutions in kerosene.

Table 3: Characterization of the ethylene copolymers used

Example	Comonomer(s)	V140	CH ₃ /100 CH ₂
A1	13.6 mol% of vinyl acetate	130 mPas	3.7
A2	13.7 mol% of vinyl acetate and 1.4 mol% of vinyl neodecanoate	105 mPas	5.3
A3	9.4 mol% of vinyl acetate	220 mPas	6.2
A4	Mixture of EVA copolymer having 16 mol% of vinyl acetate with EVA having 5 mol% of vinyl acetate in a 13:1 ratio	95 mPas 350 mPas	3.2 5.7

Comb polymers B

- 5 Maleic anhydride (MA) is polymerized with α -olefins in a relatively high-boiling aromatic hydrocarbon mixture at 160°C in the presence of a mixture of equal parts of tert-butyl peroxybenzoate and tert-butyl peroxy-2-ethylhexanoate as a radical chain initiator. Table 3 lists by way of example, various copolymers and the molar proportions of the monomers used to prepare them, and also chain length (R) and
- 10 molar amount (based on MA) of the amine used for derivatization and the factor Q calculated therefrom. Unless stated otherwise, the amines used are monoalkylamines.

- The reactions with amines are effective in the presence of Solvent Naphtha (50% by
- 15 weight) at from 50 to 100°C to give the monoamide or to give the amide ammonium salt, and at from 160 to 200°C with azeotropic separation of water of reaction to give the imide or diamide. The degree of amidation is inversely proportional to the acid number.

- 20 Table 4: Characterization of the comb polymers used

Example	Comonomers	Amine		Q	Acid number [mg KOH/g]
		R	mol		
B1	MA-co-C _{14/16} - α -olefin (1 : 0.5 : 0.5)	C ₁₀	1	23.0	60
B2	MA-co-C _{14/16} - α -olefin (1 : 0.5 : 0.5)	C ₁₂	1	25.0	58
B3	MA-co-C _{14/16} - α -olefin (1 : 0.5 : 0.5)	C ₁₄	1	27.0	56
B4 (C)	MA-co-C _{14/16} - α -olefin (1 : 0.5 : 0.5)	C ₁₆	1	29.0	55
B5	MA-co-C _{12/14} - α -olefin (1 : 0.5 : 0.5)	C ₁₄	1	25.0	57
B6	MA-co-C _{12/14} - α -olefin (1 : 0.5 : 0.5)	C ₁₂	1	23.0	55
B7	MA-co-C ₁₆ - α -olefin (1 : 1)	C ₁₂	1	26.0	56
B8	MA-co-C ₁₄ - α -olefin (1 : 1)	C ₁₄	1	26.0	58
B9	MA-co-C ₁₀ - α -olefin (1 : 1)	C ₁₆	0.5	25.0	59
		C ₁₈	0.5		
B10	MA-co-C _{14/16} - α -olefin-co-(allyl methyl polyglycol) (1 : 0.45 : 0.45 : 0.1)	C ₁₂	1	25.0	56
B11 (C)	MA-co-C ₁₀ - α -olefin (1 : 1)	C ₁₂	1	20.0	57
B12	MA-co-C _{14/16} - α -olefin (1 : 0.5 : 0.5)	C ₁₂	2	25.0	0.32
B13	MA-co-C _{14/16} - α -olefin (1 : 0.5 : 0.5)	C ₁₂	1	25.0	1.5
B14	MA-co-C _{14/16} - α -olefin (1 : 0.5 : 0.5)	di-C ₁₂	1	25.0	50
B15 (C)	Fumarate-vinyl acetate (1 : 1)	C ₁₄	2	n. a.	0.4

n.a. = not applicable

(C) = comparative example

Poly(alkyl (meth)acrylates) C

- 5 The poly(alkyl (meth)acrylates) used were the compounds listed in the table as 50% dilutions in relatively high-boiling solvent. The K values were determined according to Ubbelohde at 25°C in 5% toluenic solution.

Table 5: Characterization of the poly(acrylates) used

C1	Poly(octadecyl acrylate), K value 32
C2	Poly(dodecyl acrylate), K value 35.6
C3	Poly(beheryl acrylate), K value 22.4

Effectiveness of the terpolymers

5 The CFPP value (to EN 116, in °C) of different biofuels according to the above table was determined after the addition of 1200 ppm, 1500 ppm and also 2000 ppm, of additive mixture. Percentages relate to parts by weight in the particular mixtures. The results reported in Tables 5 to 7 show that comb polymers having the factor Q according to the invention achieve excellent CFPP reductions even at low dosages and offer additional potential at higher dosages.

Table 6: CFPP testing in test oil E1

Ex.	Comb polymer	Ethylene copolymer	Poly- acrylate	CFPP in test oil 1		
				1200 ppm	1500 ppm	2000 ppm
1	20% B1	80% A2	—	-19	-21	-23
2	20% B2	80% A2	—	-24	-24	-24
3	20% B3	80% A2	—	-19	-20	-23
4 (C)	20% B4	80% A2	—	-15	-16	-17
5	20% B7	80% A2	—	-21	-22	-24
6	20% B9	80% A2	—	-23	-23	-24
7	20% B10	80% A2	—	-23	-23	-25
8	20% B10	80% A3	—	-18	-20	-21
9	10% B10	90% A1	—	-21	-22	-23
10 (C)	20% B11	80% A2	—	-17	-19	-18
11	20% B12	80% A2	—	-19	-22	-21
12	20% B13	80% A2	—	-23	-24	-24
13	20% B14	80% A2	—	-21	-23	-24
14 (C)	20% B15	80% A2	—	-18	-17	-17
15	19% B2	76% A2	5% C1	-20	-24	-26
16	19% B2	76% A2	5% C2	-21	-23	-25
17	19% B2	76% A2	5% C3	-22	-23	-24
18 (C)	—	A2	—	-15	-18	-17
19 (C)	—	—	C1	-9	-11	-12
20 (C)	—	—	C3	-18	—	-17

Table 7: CFPP testing in test oil E2

Ex.	Comb polymer	Ethylene copolymer	CFPP in test oil 2	
			1500 ppm	2000 ppm
22	25% B2	75% A4	-20	-24
23	25% B3	75% A4	-21	-22
24 (C)	25% B4	75% A4	-13	-15
25	25% B5	75% A4	-21	-23
21	25% B6	75% A4	-19	-22
26	25% B7	75% A4	-21	-24
27	25% B8	75% A4	-20	-23
28	30% B9	70% A2	-21	-24
29	20% B10	80% A3	-20	-22
30 (C)	25% B11	75% A2	-15	-16
31	25% B12	75% A4	-20	-22
32	25% B13	75% A4	-22	-25
33	25% B14	75% A4	-18	-20
34 (C)	25% B15	75% A4	-15	-17
37 (C)	—	100% A4	-12	-12

Table 8: CFPP testing in test oil E3

Ex.	Comb polymer	Ethylene copolymer	Poly- acrylate	CFPP in test oil E3		
				1200 ppm	1500 ppm	2000 ppm
21	20% B1	80% A1	—	-18	-20	-21
22	20% B2	80% A1	—	-20	-21	-23
23	20% B3	80% A1	—	-20	-22	-21
24 (C)	20% B4	80% A1	—	-11	-15	-16
25	20% B5	80% A1	—	-20	-20	-22
26	20% B7	80% A1	—	-20	-21	-23
27	20% B8	80% A1	—	-19	-21	-22
28	25% B9	75% A2	—	-20	-21	-23
29	15% B10	85% A3	—	-18	-18	-20
30 (C)	20% B11	80% A2	—	-15	-17	-17
31	20% B12	80% A1	—	-19	-20	-20
32	20% B13	80% A1	—	-21	-22	-23
33	20% B14	80% A1	—	-19	-20	-20
34 (C)	20% B15	80% A1	—	-15	-17	-18
35	19% B2	76% A1	5% C1	-19	-21	-22
36	19% B2	76% A1	5% C3	-20	-21	-23
37 (C)	—	A1	—	-13	-13	-11
38 (C)	—	—	C3	-14	-16	-16

Cold temperature change stability of fatty acid methyl esters

To determine the cold temperature change stability of an oil, the CFPP value to DIN EN 116 before and after a standardized cold temperature change treatment are compared.

500 ml of biodiesel (test oil E1) are treated with the appropriate cold temperature additive, introduced into a measuring cylinder and stored in a programmable cold chamber for a week. Within this time, a program is run through which repeatedly cools to -13°C and then heats back to -3°C . 6 of these cycles are run through in succession (Table 8).

Table 9: Cooling program for determining the cold temperature change stability:

Section	Time	End	Duration	Description
A → B	$+5^{\circ}\text{C}$	-3°C	8 h	Precooling to cycle start temperature
B → C	-3°C	-3°C	2 h	Constant temperature, beginning of cycle
C → D	-3°C	-13°C	14 h	Temperature reduction, commencement of crystal formation
D → E	-13°C	-13°C	2 h	Constant temperature, crystal growth
E → F	-13°C	-3°C	6 h	Temperature increase, melting of the crystals
F → B				6 further B → F cycles are carried out.

Subsequently, the additized oil sample is heated to room temperature without

agitation. A sample of 50 ml is taken for CFPP measurements from each of the upper, middle and lower sections of the measuring cylinder.

A deviation between the mean values of the CFPP values after storage and the CFPP value before storage and also between the individual phases of less than 3 K

5 shows a good cold temperature change stability.

Table 10: Cold temperature change stability of the additized oil:

Example	Additive		CFPP before storage	CFPP after storage					
	Comb polymer	Ethylene copolymer		Dosage	lower	Δ CFPP (lower)	middle	Δ CFPP (middle)	upper
39	20% B2	80% A2	-24°C	-23°C	1 K	-24°C	0 K	-25°C	-1 K
40	19% B2 5% C1	76% A2	-24°C	-22°C	2 K	-23°C	1 K	-24°C	0 K
41	20% B14	80% A4	-23°C	-22°C	1 K	-21°C	2 K	-22°C	1 K
42	25% B13	75% A4	-23°C	-22°C	1 K	-23°C	0 K	-23°C	0 K
43 (C)	—	A4	-20°C	-12°C	8 K	-12.5°C	7.5 K	-14°C	6 K

The CFPP values reported are mean values of a double determination

Hideg folyási tulajdonságot javító adalék állati vagy növényi eredetű tüzelőolajokhoz

Szabadalmi igénypontok

1. Adalék, amely 10:1-től 1:10-ig terjedő tömegarányban tartalmaz A) és B) összetevőt, amely

A) összetevő egy kopolimer, amely etilénből és legalább egy akril- vagy vinilészter 8-21 mol%-ából áll, amely 1-18 szénatomos alkilgyököt hordoz, és a

B) összetevő egy fésűs polimer, amely a következő szerkezeti egységeket tartalmazza:

B1) 1. monomerként legalább egy olefint, amely az olefines kettős kötésen legalább egy 8-18 szénatomos alkilgyököt hordoz, és

B2) 2. monomerként legalább egy etilénesen telítetlen dikarbonsavat, amely amid- és/vagy imid-molekularészen keresztül legalább egy 8-16 szénatomos alkilgyököt hordoz, ahol egyrészt az 1. monomer alkilgyökeiben, másrészt a 2. monomer amid- és/vagy imidcsoportjainak alkilgyökeiben a szénlánc-hosszúság-eloszlások moláris átlagának Q összegének

$$Q = \sum_i w_{1i} \cdot n_{1i} + \sum_j w_{2j} \cdot n_{2j}$$

értéke 23-27, 1:1,5-től 1,5:1-ig terjedő B1):B2) molarány mellett, ahol

w_1 jelentése az 1. monomer alkilgyökeiben az egyes lánc-hosszúságok moláris hányada,

w_2 jelentése a 2. monomer amid- és/vagy imidcsoportjainak alkilgyökeiben az egyes lánc-hosszúságok moláris hányada,

n_1 jelentése az 1. monomer alkilgyökeiben az egyes lánc-hosszúságok,

n_2 jelentése a 2. monomer amid- és/vagy imidcsoportjainak alkilgyökeiben az egyes lánc-hosszúságok,

i jelentése az 1. monomer alkilgyökeiben az egyes lánc-hosszúságokhoz tartozó kontroll változó,

j jelentése a 2. monomer amid- és/vagy imidcsoportjainak alkilgyökeiben az egyes lánc-hosszúságokhoz tartozó kontroll változó.

2. Az 1. igénypont szerinti adalék, ahol Q értéke 24-26.

3. Az 1. és/vagy 2. igénypont szerinti adalék, ahol az A összetevő, az etilént nem számítva, tartalmaz 3,5-20 mol% vinilacetátot és 0,1-12 mol% vinil-2-etilhexanoátot, vinil-neononanoátot és/vagy vinil-neodekanoátot, és az összes komonomer-tartalom 8 és 21 mol% közötti.

4. Az 1-3. igénypontok bármelyike szerinti adalék, ahol az A összetevő, az etilénen és a 8-18 mol% vinilésztereken kívül 0,5 - 10 mol% mennyiségben olefineket is tartalmaz, amelyek a következők közül vannak kiválasztva: propén, butén, izobutilén, hexén, 4-metilpentén, oktén, diizobutilén és norbornén.

5. Az 1-4. igénypontok bármelyike szerinti adalék, ahol az A összetevőt alkotó kopolimerek olvadákvizkozitása 20 és 10 000 mPas közötti.

6. Az 1-5. igénypontok bármelyike szerinti adalék, ahol az A összetevőt alkotó kopolimerek elágazási foka 1-9 $\text{CH}_3/100 \text{ CH}_2$ csoport, amely nem a komonomerekről származik.

7. Az 1-6. igénypontok bármelyike szerinti adalék, ahol a B összetevőt alkotó kopolimerek tartalmaznak komonomereket, amelyek a maleinsav, fumársav vagy itakonsav amidjaiból és/vagy imidjeiből származnak.

8. Az 1-7. igénypontok bármelyike szerinti adalék, ahol a B összetevő amidjai és/vagy imidjei primer aminosokból származnak.

9. Az 1-8. igénypontok bármelyike szerinti adalék, ahol a B összetevő amidjai és/vagy imidjei lineáris alkilgyökökkel rendelkező aminosokból származnak.

10. Az 1-9. igénypontok bármelyike szerinti adalék, ahol a B összetevő amidjai és/vagy imidjei monoaminokból származnak.

11. Az 1-10. igénypontok bármelyike szerinti adalék, ahol a B összetevőt alkotó kopolimerek tartalmaznak α -olefinekből származó komonomereket.

12. Az 1-11. igénypontok bármelyike szerinti adalék, ahol az A és B összetevőn felül egy C összetevő is jelen van, az A, B és C összetevő együttes tömegére vonatkoztatva legfeljebb 40 tömeg% mennyiségben, amely C összetevő egy polimer vagy kopolimer, amely (C_{10} - C_{24} -

alkil)-akrilát egységeket vagy metakrilát egységeket foglal magában, és amelynek molekulatömege 800 - 1 000 000 g/mol.

13. Az 1-12. igénypontok bármelyike szerinti adalék, amely tartalmaz poláros nitrogéntartalmú paraffin diszpergenset.

14. Tüzelőolaj-készítmény, amely egy állati vagy növényi eredetű tüzelőolajat és 0,001-5 tömeg% mennyiségben 1-13. igénypontok bármelyike szerinti adalékot tartalmaz.

15. A 14. igénypont szerinti tüzelőolaj-készítmény, ahol az állati vagy növényi eredetű tüzelőolaj 14-24-szénatomos monokarbonsav és 1-4 szénatomos alkohol egy vagy több észterét tartalmazza.

16. A 15. igénypont szerinti tüzelőolaj-készítmény, ahol az alkohol metanol vagy etanol.

17. A 14-16. igénypontok bármelyike szerinti tüzelőolaj-készítmény, ahol az állati vagy növényi eredetű tüzelőolaj 5 tömeg%-nál nagyobb mennyiségben tartalmaz telített zsírsavakkal képzett észtereket.

18. Az 1-13. igénypontok bármelyike szerinti adalék alkalmazása 0,001-5 tömeg% mennyiségben, állati vagy növényi eredetű tüzelőolajok hideg folyási tulajdonságainak javítására.