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(54) Title: MICROCAPILLARY FILMS AND FOAMS SUITABLE FOR CAPILLARY ACTION FLUID TRANSPORT

(57) Abstract: The instant invention provides microcapiUary films and /or foams suitable for capillary action fluid transport. The inventive microcapiUary film and/or foam containing suitable for capillary action fluid transport according to the present invention has a first end and a second end, and comprises:(a) a matrix comprising a thermoplastic material, and (b) at least one or more channels disposed in parallel in said matrix from the first end to the second end of said microcapiUary film and/or foam, wherein said one or more channels are at least 1 µm apart from each other, wherein each said one or more channels have a diameter in the range of at least 1 μm; wherein said microcapiUary film and/or foam has a thickness in the range of from 2 μm to 2000 μm.

MICROCAPILLARY FILMS AND FOAMS SUITABLE FOR CAPILLARY ACTION FLUID TRANSPORT

Field of Invention

The instant invention relates to microcapiUary films and/or foams suitable for capillary action fluid transport.

Background of the Invention

Capillary action, or capillarity, is the ability of a liquid to flow against gravity where liquid spontaneously rises in a narrow space such as between the hairs of a paint-brush, in a thin tube, in porous material such as paper, in some non-porous materials such as liquified carbon fiber, or in a cell. This effect can cause liquids to flow against the force of gravity, sun or any electro magnetic field affecting fluid flow. It occurs because of inter-molecular attractive forces between the transporting liquid and surrounding surface having a different surface energy. For the case of a tube, if the diameter of the tube is sufficiently small, then the combination of surface energy (which is caused by cohesion within the liquid) and force of adhesion between the liquid and tube wall act to lift the liquid.

While there are numerous methods for liquid transport based on capillary action, there is still a need for improved microcapiUary films and/or foams suitable for capillary action fluid transport.

Summary of the Invention

The instant invention provides microcapiUary films and/or foams suitable for capillary action fluid transport. The inventive microcapiUary film and/or foam containing suitable for capillary action fluid transport according to the present invention has a first end and a second end, and comprises: (a) a matrix comprising a thermoplastic material, and (b) at least one or more channels disposed in parallel in said matrix from the first end to the second end of said microcapiUary film and/or foam, wherein said one or more channels are at least 1 μιη apart from each other, wherein each said one or more channels have a diameter in the range of at least 1 μιη; wherein said microcapiUary film and/or foam has a thickness in the range of from 2 μιη to 2000 μιη.

In an alternative embodiment, the instant invention provides microcapiUary films and /or foams suitable for capillary action fluid transport, in accordance with any of the preceding

embodiments, except that the thermoplastic material is selected from the group consisting of polyolefin, e.g. polyethylene and polypropylene; polyamide, e.g. nylon 6; polyvinylidene chloride; polyvinylidene fluoride; polycarbonate; polystyrene; polyethylene terephthalate; polyurethane and/or polyester.

In an alternative embodiment, the instant invention provides microcapiUary films and /or foams suitable for capillary action fluid transport, in accordance with any of the preceding embodiments, except that the one or more channels have a cross sectional shape selected from the group consisting of circular, rectangular, oval, star, diamond, triangular, square, the like, and combinations thereof.

In an alternative embodiment, the instant invention provides microcapiUary films and /or foams suitable for capillary action fluid transport, in accordance with any of the preceding embodiments, except that the microcapiUary channels and optionally the microcapiUary film and/or foam surfaces are treated, e.g. surface treated, using a dielectric barrier discharge atmospheric or low pressure plasma including aerosoled functional molecules such as amines, hydroxyls, allyls, acrylics, fluorines, silicones, and the like to modify surface energy.

Brief Description of the Drawings

For the purpose of illustrating the invention, there is shown in the drawings a form that is exemplary; it being understood, however, that this invention is not limited to the precise arrangements and instrumentalities shown.

- **Fig. 1** is a top view of an inventive microcapiUary film or foam suitable for capillary action fluid transport;
- **Fig. 2** is a longitudinal-sectional view of an inventive microcapiUary film or foam suitable for capillary action fluid transport;
- **Fig. 3a-e** are various cross-sectional views of an inventive microcapiUary film or foam suitable for capillary action fluid transport;
- **Fig. 4** is an elevated view of an inventive microcapiUary film or foam suitable for capillary action fluid transport;
- **Fig. 5** is a segment of a longitudinal sectional view of the inventive microcapiUary film or foam suitable for capillary action fluid transport, as shown in figure 2;
- **Fig. 6** is an exploded view of an inventive microcapiUary film or foam suitable for capillary action fluid transport;

Figs. 7a-b are schematic illustration of a microcapiUary die;

Fig. 8 is a graph illustrating the results of tensiometer evaluation of the samples DOW101-DOW112.

Fig. 9 is a photograph of samples after tensiometer evaluation; and

Fig. 8 is another photograph of a sample after tensiometer evaluation.

Detailed Description of the Invention

Referring to the drawings wherein like numerals indicate like elements, there is shown, in figures 1-6, a first embodiment of a microcapiUary film or foam (10) suitable for capillary action fluid transport.

The inventive microcapiUary film or foam (10) suitable for capillary action fluid transport according to the present invention has a first end (14) and a second end (16), and comprises: (a) a matrix (18) comprising a thermoplastic material; (b) at least one or more channels (20) disposed in parallel in said matrix (18) from the first end (14) to the second end (16) of said microcapiUary film or foam (10), wherein said one or more channels (20) are at least 1 μιη apart from each other, and wherein each said one or more channels (20) have a diameter in the range of at least 1 μιη; wherein said microcapiUary film (10) has a thickness in the range of from 2 μιη to 2000 μιη.

The microcapiUary film or foam (10) suitable for capillary action fluid transport may have a thickness in the range of from 2 μ m to 2000 μ m; for example, microcapiUary film or foam (10) suitable for capillary action fluid transport may have a thickness in the range of from 10 to 2000 μ m; or in the alternative, from 200 to 800 μ m; or in the alternative, from 200 to 800 μ m; or in the alternative, from 300 to 1000 μ m; or in the alternative, from 300 to 1000 μ m; or in the alternative, from 300 to 900 μ m; or in the alternative, from 300 to 700 μ m. The film thickness to microcapiUary diameter ratio is in the range of from 2:1 to 2000:1. The term "microcapiUary film," as used herein refers to films as well as tapes, and may be formed into the shape of a pipe, rod or profile.

The microcapiUary film or foam (10) suitable for capillary action fluid transport may comprise at least 10 percent by volume of the matrix (18), based on the total volume of the microcapiUary film or foam (10) suitable for capillary action fluid transport; for example, the microcapiUary film or foam (10) suitable for capillary action fluid transport may comprise from 10 to 80 percent by volume of the matrix (18), based on the total volume of the microcapiUary film or foam (10) suitable for capillary action fluid transport; or in the alternative, from 20 to 80 percent by volume of the matrix (18), based on the total volume of the microcapiUary film or foam (10) suitable

for capillary action fluid transport; or in the alternative, from 30 to 80 percent by volume of the matrix (18), based on the total volume of the microcapillary film or foam (10) suitable for capillary action fluid transport.

The microcapillary film or foam (10) suitable for capillary action fluid transport may comprise from 20 to 90 percent by volume of voidage, based on the total volume of the microcapillary film or foam (10) suitable for capillary action fluid transport); for example, the microcapillary film or foam (10) suitable for capillary action fluid transport may comprise from 20 to 80 percent by volume of voidage, based on the total volume of the microcapillary film or foam (10) suitable for capillary action fluid transport; or in the alternative, from 20 to 70 percent by volume of voidage, based on the total volume of the microcapillary film or foam (10) suitable for capillary action fluid transport; or in the alternative, from 30 to 60 percent by volume of voidage, based on the total volume of the microcapillary film or foam (10) suitable for capillary action fluid transport.

The inventive microcapillary film or foam (10) has a first end (14) and a second end (16). At least one or more channels (20) are disposed in parallel in the matrix (18) from the first end (14) to the second end (16). The one or more channels (20) are at least $1 \mu \eta$, apart from each other. The one or more channels (20) have a diameter in the range of at least $1 \mu \eta$; for example, from $1 \mu \eta$ to 1998 $\mu \eta$; or in the alternative, from 5 to 990 $\mu \eta$; or in the alternative, from 5 to 890 $\mu \eta$; or in the alternative, from 5 to 790 $\mu \eta$; or in the alternative, from 5 to 690 $\mu \eta$ or in the alternative, from 5 to 590 $\mu \eta$. The one or more channels (20) may have a cross-sectional shape selected from the group consisting of circular, rectangular, oval, star, diamond, triangular, square, the like, and combinations thereof. The one or more channels (20) may further include one or more seals at the first end (14), the second end (16), therebetween the first point (14) and the second end (16), and/or combinations thereof.

The microcapillary channels, i.e. (inside of the microcapillaries), and optionally the microcapillary film or foam (10) surface may further be modified by surface treatment of the microcapillary channels, i.e. (inside of the microcapillaries) or the surface treatment of the film and /or foam surface, or combinations thereof via, for example, plasma surface treatment, and/or chemical grafting surface treatment. In one embodiment, the microcapillary channels, i.e. (inside of the microcapillaries), and optionally the microcapillary film or foam (10) surface are treated in a dielectric barrier discharge atmospheric or low pressure plasma including aerosoled functional

molecules such as amines, hydroxyls, allyls, acrylics, fluorines, silicones, and the like to modify surface energy. The surface treatment may be for a period in the range of from 1 second to one hour, for example, from 1 to 60 seconds. Surface treatment can be achieved via any known methods.

The matrix (18) comprises one or more thermoplastic materials. Such thermoplastic materials include, but are not limited to, polyolefin, e.g. polyethylene and polypropylene; polyamide, e.g. nylon 6; polyvinylidene chloride; polyvinylidene fluoride; polycarbonate; polystyrene; polyethylene terephthalate; polyester, and polyurethanes. The matrix (18) may be reinforced via, for example, glass or carbon fibers and/or any other mineral fillers such talc or calcium carbonate. Exemplary fillers include, but are not limited to, natural calcium carbonates, including chalks, calcites and marbles, synthetic carbonates, salts of magnesium and calcium, dolomites, magnesium carbonate, zinc carbonate, lime, magnesia, barium sulphate, barite, calcium sulphate, silica, magnesium silicates, talc, wollastonite, clays and aluminum silicates, kaolins, mica, oxides or hydroxides of metals or alkaline earths, magnesium hydroxide, iron oxides, zinc oxide, glass or carbon fiber or powder, wood fiber or powder or mixtures of these compounds.

Examples of thermoplastic materials include, but are not limited to, homopolymers and copolymers (including elastomers) of one or more alpha-olefins such as ethylene, propylene, 1butene, 3-methyl-l-butene, 4-methyl-l-pentene, 3-methyl-l-pentene, 1-hexene, 1-octene, 1-decene, and 1-dodecene, as typically represented by polyethylene, polypropylene, poly-1-butene, poly-3-methyl-l-butene, poly-3-methyl-l-pentene, poly-4-methyl-l-pentene, ethylene-propylene copolymer, ethylene-l-butene copolymer, and propylene- l-butene copolymer; copolymers (including elastomers) of an alpha-olefin with a conjugated or non-conjugated diene, as typically represented by ethylene-butadiene copolymer and ethylene-ethylidene norbornene copolymer; and polyolefins (including elastomers) such as copolymers of two or more alpha-olefins with a conjugated or nonconjugated diene, as typically represented by ethylene-propylene-butadiene copolymer, ethylenepropylene- dicyclopentadiene copolymer, ethylene-propylene- 1,5-hexadiene copolymer, and ethylene-propylene-ethylidene norbornene copolymer; ethylene-vinyl compound copolymers such as ethylene-vinyl acetate copolymer, ethylene-vinyl alcohol copolymer, ethylene-vinyl chloride copolymer, ethylene acrylic acid or ethylene-(meth)acrylic acid copolymers, and ethylene-(meth)acrylate copolymer; styrenic copolymers (including elastomers) such as polystyrene, ABS, acrylonitrile-styrene copolymer, cc-methylstyrene-styrene copolymer, styrene vinyl alcohol, styrene acrylates such as styrene methylacrylate, styrene butyl acrylate, styrene butyl methacrylate, and

styrene butadienes and crosslinked styrene polymers; and styrene block copolymers (including elastomers) such as styrene-butadiene copolymer and hydrate thereof, and styrene-isoprene-styrene triblock copolymer; polyvinyl compounds such as polyvinyl chloride, polyvinylidene chloride, vinyl chloride-vinylidene chloride copolymer, polymethyl acrylate, and polymethyl methacrylate; polyamides such as nylon 6, nylon 6,6, and nylon 12; thermoplastic polyesters such as polyethylene terephthalate and polybutylene terephthalate; polyurethane; polycarbonate, polyphenylene oxide, and the like; and glassy hydrocarbon-based resins, including poly-dicyclopentadiene polymers and related polymers (copolymers, terpolymers); saturated mono-olefins such as vinyl acetate, vinyl propionate, vinyl versatate, and vinyl butyrate and the like; vinyl esters such as esters of monocarboxylic acids, including methyl acrylate, ethyl acrylate, n-butyl acrylate, isobutyl acrylate, 2-ethylhexyl acrylate, dodecyl acrylate, n-octyl acrylate, phenyl acrylate, methyl methacrylate, ethyl methacrylate, and butyl methacrylate and the like; acrylonitrile, methacrylonitrile, acrylamide, mixtures thereof; resins produced by ring opening metathesis and cross metathesis polymerization and the like. These resins may be used either alone or in combinations of two or more.

In selected embodiments, thermoplastic material may, for example, comprise one or more polyolefins selected from the group consisting of ethylene-alpha olefin copolymers, propylene-alpha olefin copolymers, and olefin block copolymers. In particular, in select embodiments, the thermoplastic material may comprise one or more non-polar polyolefins.

In specific embodiments, polyolefins such as polypropylene, polyethylene, copolymers thereof, and blends thereof, as well as ethylene-propylene-diene terpolymers, may be used. In some embodiments, exemplary olefinic polymers include homogeneous polymers; high density polyethylene (HDPE); heterogeneously branched linear low density polyethylene (LLDPE); heterogeneously branched ultra low linear density polyethylene (ULDPE); homogeneously branched, linear ethylene/alpha-olefin copolymers; homogeneously branched, substantially linear ethylene/alpha-olefin polymers; and high pressure, free radical polymerized ethylene polymers and copolymers such as low density polyethylene (LDPE) or ethylene vinyl acetate polymers (EVA).

In one embodiment, the ethylene-alpha olefin copolymer may, for example, be ethylene-butene, ethylene-hexene, or ethylene-octene copolymers or interpolymers. In other particular embodiments, the propylene-alpha olefin copolymer may, for example, be a propylene-ethylene or a propylene-ethylene-butene copolymer or interpolymer.

In certain other embodiments, the thermoplastic material may, for example, be a semi-crystalline polymer and may have a melting point of less than 110°C. In another embodiment, the melting point may be from 25 to 100°C. In another embodiment, the melting point may be between 40 and 85°C.

In one particular embodiment, the thermoplastic material is a propylene/a-olefin interpolymer composition comprising a propylene/alpha-olefin copolymer, and optionally one or more polymers, e.g. a random copolymer polypropylene (RCP). In one particular embodiment, the propylene/alpha-olefin copolymer is characterized as having substantially isotactic propylene sequences. "Substantially isotactic propylene sequences" means that the sequences have an isotactic triad (mm) measured by ¹³C NMR of greater than about 0.85; in the alternative, greater than about 0.90; in another alternative, greater than about 0.92; and in another alternative, greater than about 0.93. Isotactic triads are well-known in the art and are described in, for example, U.S. Patent No. 5,504,172 and International Publication No. WO 00/01745, which refer to the isotactic sequence in terms of a triad unit in the copolymer molecular chain determined by ¹³C NMR spectra.

The propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 0.1 to 500 g/10 minutes, measured in accordance with ASTM D-1238 (at 230° C / 2.16 Kg). All individual values and subranges from 0.1 to 500 g/10 minutes are included herein and disclosed herein; for example, the melt flow rate can be from a lower limit of 0.1 g/10 minutes, 0.2 g/10 minutes, or 0.5 g/10 minutes to an upper limit of 500 g/10 minutes, 200 g/10 minutes, 100 g/10 minutes, or 25 g/10 minutes. For example, the propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 0.1 to 200 g/10 minutes; or in the alternative, the propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 0.2 to 100 g/10 minutes; or in the alternative, the propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 0.5 to 50 g/10 minutes; or in the alternative, the propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 1 to 50 g/10 minutes; or in the alternative, the propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 1 to 40 g/10 minutes; or in the alternative, the propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 1 to 40 g/10 minutes; or in the alternative, the propylene/alpha-olefin copolymer may have a melt flow rate in the range of from 1 to 30 g/10 minutes.

The propylene/alpha-olefin copolymer has a crystallinity in the range of from at least 1 percent by weight (a heat of fusion of at least 2 Joules/gram) to 30 percent by weight (a heat of

fusion of less than 50 Joules/gram). All individual values and subranges from 1 percent by weight (a heat of fusion of at least 2 Joules/gram) to 30 percent by weight (a heat of fusion of less than 50 Joules/gram) are included herein and disclosed herein; for example, the crystallinity can be from a lower limit of 1 percent by weight (a heat of fusion of at least 2 Joules/gram), 2.5 percent (a heat of fusion of at least 4 Joules/gram), or 3 percent (a heat of fusion of at least 5 Joules/gram) to an upper limit of 30 percent by weight (a heat of fusion of less than 50 Joules/gram), 24 percent by weight (a heat of fusion of less than 40 Joules/gram), 15 percent by weight (a heat of fusion of less than 24.8 Joules/gram) or 7 percent by weight (a heat of fusion of less than 11 Joules/gram). For example, the propylene/alpha-olefin copolymer may have a crystallinity in the range of from at least 1 percent by weight (a heat of fusion of at least 2 Joules/gram) to 24 percent by weight (a heat of fusion of less than 40 Joules/gram); or in the alternative, the propylene/alpha-olefin copolymer may have a crystallinity in the range of from at least 1 percent by weight (a heat of fusion of at least 2 Joules/gram) to 15 percent by weight (a heat of fusion of less than 24.8 Joules/gram); or in the alternative, the propylene/alpha-olefin copolymer may have a crystallinity in the range of from at least 1 percent by weight (a heat of fusion of at least 2 Joules/gram) to 7 percent by weight (a heat of fusion of less than 11 Joules/gram); or in the alternative, the propylene/alpha-olefin copolymer may have a crystallinity in the range of from at least 1 percent by weight (a heat of fusion of at least 2 Joules/gram) to 5 percent by weight (a heat of fusion of less than 8.3 Joules/gram). The crystallinity is measured via DSC method. The propylene/alpha-olefin copolymer comprises units derived from propylene and polymeric units derived from one or more alpha-olefin comonomers. Exemplary comonomers utilized to manufacture the propylene/alpha-olefin copolymer are C_2 , and C_4 to C_{10} alpha-olefins; for example, C_2 , C_4 , C_6 and C_8 alpha-olefins.

The propylene/alpha-olefin copolymer comprises from 1 to 40 percent by weight of one or more alpha-olefin comonomers. All individual values and subranges from 1 to 40 weight percent are included herein and disclosed herein; for example, the comonomer content can be from a lower limit of 1 weight percent, 3 weight percent, 4 weight percent, 5 weight percent, 7 weight percent, or 9 weight percent to an upper limit of 40 weight percent, 35 weight percent, 30 weight percent, 27 weight percent, 20 weight percent, 15 weight percent, 12 weight percent, or 9 weight percent. For example, the propylene/alpha-olefin copolymer comprises from 1 to 35 percent by weight of one or more alpha-olefin comonomers; or in the alternative, the propylene/alpha-olefin copolymer comprises from 1 to 30 percent by weight of one or more alpha-olefin comonomers; or in the

alternative, the propylene/alpha-olefin copolymer comprises from 3 to 27 percent by weight of one or more alpha-olefin comonomers; or in the alternative, the propylene/alpha-olefin copolymer comprises from 3 to 20 percent by weight of one or more alpha-olefin comonomers; or in the alternative, the propylene/alpha-olefin copolymer comprises from 3 to 15 percent by weight of one or more alpha-olefin comonomers.

The propylene/alpha-olefin copolymer has a molecular weight distribution (MWD), defined as weight average molecular weight divided by number average molecular weight (M_w/M_n) of 3.5 or less; in the alternative 3.0 or less; or in another alternative from 1.8 to 3.0.

Such propylene/alpha-olefin copolymers are further described in details in the U.S. Patent Nos. 6,960,635 and 6,525,157, incorporated herein by reference. Such propylene/alpha-olefin copolymers are commercially available from The Dow Chemical Company, under the tradename VERSIFYTM, or from ExxonMobil Chemical Company, under the tradename VISTAMAXXTM.

In one embodiment, the propylene/alpha-olefin copolymers are further characterized as comprising (A) between 60 and less than 100, preferably between 80 and 99 and more preferably between 85 and 99, weight percent units derived from propylene, and (B) between greater than zero and 40, preferably between 1 and 20, more preferably between 4 and 16 and even more preferably between 4 and 15, weight percent units derived from at least one of ethylene and/or a $C_{4\rightarrow}$ occ-olefin; and containing an average of at least 0.001, preferably an average of at least 0.005 and more preferably an average of at least 0.01, long chain branches/1000 total carbons. The maximum number of long chain branches in the propylene/alpha-olefin copolymer is not critical, but typically it does not exceed 3 long chain branches/1000 total carbons. The term long chain branch, as used herein with regard to propylene/alpha-olefin copolymers, refers to a chain length of at least one (1) carbon more than a short chain branch, and short chain branch, as used herein with regard to propylene/alpha-olefin copolymers, refers to a chain length of two (2) carbons less than the number of carbons in the comonomer. For example, a propylene/ 1-octene interpolymer has backbones with long chain branches of at least seven (7) carbons in length, but these backbones also have short chain branches of only six (6) carbons in length. Such propylene/alpha-olefin copolymers are further described in details in the U.S. Provisional Patent Application No. 60/988,999 and International Patent Application No. PCT/US08/082599, each of which is incorporated herein by reference.

In certain other embodiments, the thermoplastic material, e.g. propylene/alpha-olefin copolymer, may, for example, be a semi-crystalline polymer and may have a melting point of less

than 110°C. In preferred embodiments, the melting point may be from 25 to 100°C. In more preferred embodiments, the melting point may be between 40 and 85°C.

In other selected embodiments, olefin block copolymers, e.g., ethylene multi-block copolymer, such as those described in the International Publication No. WO2005/090427 and U.S. Patent Application Publication No. US 2006/0199930, incorporated herein by reference to the extent describing such olefin block copolymers and the test methods for measuring those properties listed below for such polymers, may be used as the thermoplastic material. Such olefin block copolymer may be an ethylene/a-olefin interpolymer:

(a) having a M_w/M_n from about 1.7 to about 3.5, at least one melting point, T_m , in degrees Celsius, and a density, d, in grams/cubic centimeter, wherein the numerical values of T_m and d corresponding to the relationship:

$$T_m > -2002.9 + 4538.5(d) - 2422.2(d)^2$$
; or

(b) having a M_w/M_n from about 1.7 to about 3.5, and being characterized by a heat of fusion, ΔH in J/g, and a delta quantity, ΔT , in degrees Celsius defined as the temperature difference between the tallest DSC peak and the tallest CRYSTAF peak, wherein the numerical values of ΔT and ΔH having the following relationships:

 $\Delta T >$ -0.1299($\Delta H)$ + 62.81 for ΔH greater than zero and up to 130 J/g,

 $\Delta T \ge 48^{\circ}C$ for ΔH greater than 130 J/g,

wherein the CRYSTAF peak being determined using at least 5 percent of the cumulative polymer, and if less than 5 percent of the polymer having an identifiable CRYSTAF peak, then the CRYSTAF temperature being 30 °C; or

(c) being characterized by an elastic recovery, Re, in percent at 300 percent strain and 1 cycle measured with a compression-molded film of the ethylene/a-olefin interpolymer, and having a density, d, in grams/cubic centimeter, wherein the numerical values of Re and d satisfying the following relationship when ethylene/a-olefin interpolymer being substantially free of a cross-linked phase:

Re > 1481-1629(d); or

(d) having a molecular fraction which elutes between 40 °C and 130 °C when fractionated using TREF, characterized in that the fraction having a molar comonomer content of at least 5 percent higher than that of a comparable random ethylene interpolymer fraction eluting between the same temperatures, wherein said comparable random ethylene interpolymer having the same

comonomer(s) and having a melt index, density, and molar comonomer content (based on the whole polymer) within 10 percent of that of the ethylene/a-olefin interpolymer; or

- (e) having a storage modulus at 25 °C, G′ (25 °C), and a storage modulus at 100 °C, G′ (100 °C), wherein the ratio of G′ (25 °C) to G′ (100 °C) being in the range of about 1:1 to about 9:1. Such olefin block copolymer, e.g. ethylene/a-olefin interpolymer may also:
- (a) have a molecular fraction which elutes between 40 °C and 130 °C when fractionated using TREF, characterized in that the fraction having a block index of at least 0.5 and up to about 1 and a molecular weight distribution, M_w/M_n , greater than about 1.3; or
- (b) have an average block index greater than zero and up to about 1.0 and a molecular weight distribution, M_n/M_n , greater than about 1.3.

In one embodiment, matrix (18) may further comprise a blowing agent thereby facilitating the formation a foam material. In one embodiment, the matrix may be a foam, for example a closed cell foam. In another embodiment, matrix (18) may further comprise one or more fillers thereby facilitating the formation a microporous matrix, for example, via orientation, e.g. biaxial orientation, or cavitation, e.g. uniaxial or biaxial orientation, or leaching, i.e. dissolving the fillers. Such fillers include, but are not limited to, natural calcium carbonates, including chalks, calcites and marbles, synthetic carbonates, salts of magnesium and calcium, dolomites, magnesium carbonate, zinc carbonate, lime, magnesia, barium sulphate, barite, calcium sulphate, silica, magnesium silicates, talc, wollastonite, clays and aluminum silicates, kaolins, mica, oxides or hydroxides of metals or alkaline earths, magnesium hydroxide, iron oxides, zinc oxide, glass or carbon fiber or powder, wood fiber or powder or mixtures of these compounds.

In production, the extrusion apparatus comprises a screw extruder driven by a motor. Thermoplastic material is melted and conveyed to a die (24), as shown in Figures 7a and 7b. The molten/semi-molten thermoplastic material passes through die (24), as shown in Figures 7a and 7b, and is formed into the desired shape and cross section. Referring to Figures 7a and 7b, die (24) includes an entry portion (26), a convergent portion (28), and an orifice (30), which has a predetermined shape. The molten/semi-molten thermoplastic polymer enters entry portion (26) of the die (24), and is gradually shaped by the convergent portion (28) until the melt exits the orifice (30). The die (24) further includes injectors (32). Each injector (32) has a body portion (34) having a conduit (36) therein which is fluidly connected to one or more gases, e.g. air or nitrogen, source (38) by means of second conduit (40) passing through the walls of die (24) around which the

molten/semi-molten thermoplastic material must flow to pass the orifice (30). The injector (30) further includes an outlet (42). The injector (32) is arranged such that the outlet (42) is located within the orifice (30). As the molten/semi-molten thermoplastic polymer exits the die orifice (30), one or more gases, e.g. air or nitrogen (12) is injected into the molten/semi-molten thermoplastic material thereby forming microcapillaries films and/or foams suitable for capillary action fluid transport.

The microcapiUary films or foams suitable for capillary action fluid transport according to the present invention may be formed, and then subjected to plasma treatment to modify the surface, e.g. forming a plasma coating (12). The plasma coating (12), as used herein, can be continuous, discontinuous, and/or integrated part of the surface. In the alternative, plasma gas such as a nobel gas and/or nitrogen may be injected into the molten/semi-molten thermoplastic material as part of the process for making the microcapiUary films or foams suitable for capillary action fluid transport and subsequently modifying microcapillaries surface by inducing a plasma form the inserted gas or gas mixture in the microcapillaries via for example a dielectric barrier discharge technology and. forming in the process a plasma coating and/or surface energy modification(12).

The microcapiUary films or foams suitable for capillary action fluid transport according to the present invention may be used in fluid transportation such as micro-irrigation. The microcapiUary films or foams suitable for capillary action fluid transport according to the present invention may be formed into capillary tubes. Capillary tubes are small irrigation distribution tubes (drip tubes) that are assembled into drip stakes to make a drip irrigation system for use in modern glasshouse horticulture. These capillary tubes are available in various sizes and lengths, and with the adjustment in working pressure, the water supply delivery volume can be achieved.

One or more inventive microcapiUary films and/forms suitable for capillary action fluid transport may be combined to form one or more layers in a multilayer structure, for example, a laminated multilayer structure or a coextraded multilayer structure. The microcapiUary films or foams suitable for capillary action fluid transport may comprise one or more parallel rows of microcapillaries (channels as shown in Fig. 3). Channels (20) (microcapillaries) may be disposed any where in matrix (10), as shown in Figs. 3.

Examples

The following examples illustrate the present invention but are not intended to limit the scope of the invention.

PlasmaZone® developed by VITO - Flemish Institute for Technological Research in Belgium was used as plasma equipment. ThePlasmaZone® (1 electrode of 40 cm width) operates according to the following specifications:

Table 1

Power	10-1000 W
Voltage	1-100 kV
Frequency	1-100 kHz
Plasma gas	nitrogen, air, oxygen, carbon dioxide, hydrogen, helium, argon, and any mixtures thereof
Gas consumption	5-50 l/min
Chemical precursors (liquid or gaseous)	optional: hydrocarbons, organic acids, siloxanes
Precursor consumption	1-20 ml/min
Homogeneous plasma area	variable, typical 8 to 20 cm
Inter electrode distance	0.5-25 mm
Working temperature	25-250 °C
Typical deposition rate for coatings	1 - 100 nm/sec
Inter electrode distance	0.5-25 mm

It consists of two electrodes from which the upper one is connected to high voltage and the lower one is grounded. A dielectric barrier discharge is generated between both electrodes. In order to surface treat the inner side of the capillary tapes, a specific gas mixture was blown in the capillaries. The capillary tape was then disposed in the plasma zone for a fixed time, and thereby forming a plasma chemically modified surface or coating on the surfaces of the inner side of the capillaries.

12 experimental conditions designated as DOW101-DOW112 in Table 2 were evaluated, and results are shown in Fig. 8. DOW101-DOW110 were samples with the surface, inside of the microcapillaries, treated via gaseous precursors. DOW111-DOW112 were samples with the surface inside of the microcapillaries treated via a liquid precursor that was atomized to a very fine aerosol.

Table 2

Sample No.	Functionality	Plasma Gas Type	Treatment Time
			(s)
DOW101	none	Не	5
DOW102	Amine	He/N ₂	5
DOW103	Amine	He/N ₂ /H ₂	5
DOW104	Amine	He/NH ₃	5
DOW105	Amine	He/N ₂ /H ₂ /NH ₃	20
DOW106	Amine	He/N ₂ /H ₂ /NH ₃	5
DOW107	Amine	He/N ₂ /H ₂ /NH ₃	20
DOW108	Amine	He/N ₂ /H ₂ /NH ₃	10
DOW109	Amine	He/N ₂ /H ₂ /NH ₃	20
DOW110	Hydroxyl	He/N ₂ /H ₂	20
DOW111	Hydroxyl	Не	20
DOW112		Не	5

The gas mixture's main component was nitrogen with small amounts of He (< 1% vol) as plasma initiating gas and depending on the desired functionality up to 5% vol of Hydrogen and/or, Ammonia.

A tensiometer test was used to evaluate the effect of the different experimental conditions on the hydrophilic behavior of the inner side of the capillaries. A LLDPE DOWLEXTM 2045G sample of about 7 cm height, 32.5 cm wide and a thickness of 790 μιη with 42 microcapillaries with each an average diameter of 400 μιη was put into a bath filled with de-ionized water at room temperature. The samples were immersed for 2 cm and the uptake of water in weight was measured continuously. After 2 minutes immersion, the samples were removed from the water and the uptake of water was inspected visually, and the results are shown in Figure 9.

Referring to Fig. 10, hydrocarbon, non polar fluids having surface tensions similar that of polyolefins resins capillary effects are achieved with or without treatment with options to induce capillary rise between 3-4 m provided the diameter is about 5-10 μιη and the surface tension of the tubes equals that of the liquid. LLDPE DOWLEXTM 2045G untreated prototype samples show capillary rise with n-hexane (18.43 mN/m @ 20°C). For a 400 μιη capillary a rise of about 35 mm is observed, while for a 145 μιη capillary a rise of about 80 mm is noted.

The present invention may be embodied in other forms without departing from the spirit and the essential attributes thereof, and, accordingly, reference should be made to the appended claims, rather than to the foregoing specification, as indicating the scope of the invention.

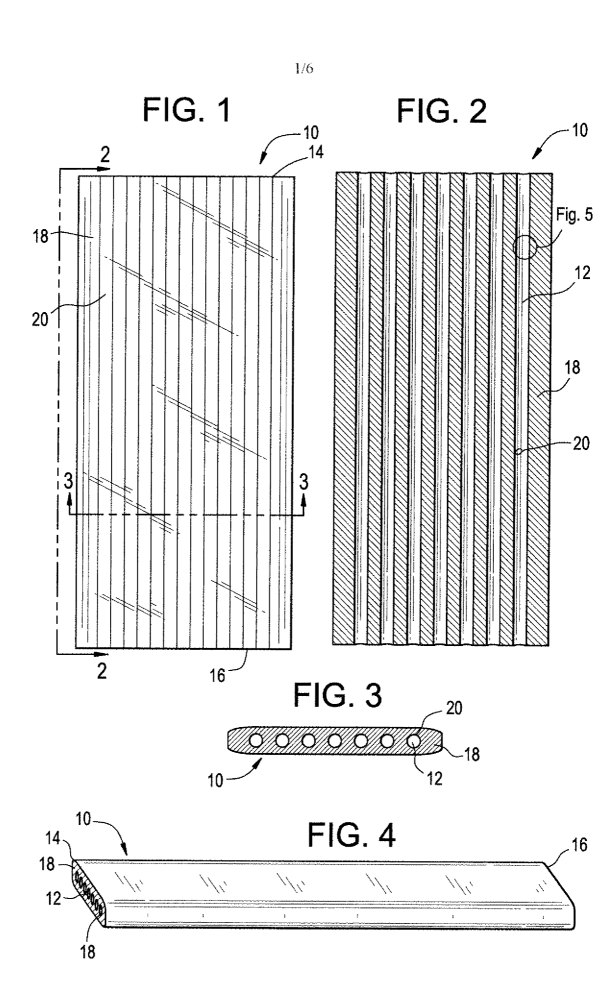
We Claim:

1. A microcapiUary film or foam having a first end and a second end, wherein said film comprises:

- (a) a matrix comprising a thermoplastic material,
- (b) at least one or more channels disposed in parallel in said matrix from the first end to the second end of said film, wherein said one or more channels are at least $1 \mu \eta$ apart from each other, and wherein each said one or more channels have a diameter in the range of at least $1 \mu \eta$;

wherein said film has a thickness in the range of from 2 μιη to 2000 μιη.

- 2. The microcapiUary film or foam of Claim 1, wherein said thermoplastic material is selected from the group consisting of polyolefin; polyamide; polyvinylidene chloride; polyvinylidene fluoride; polyurethane; polycarbonate; polystyrene; polyethylene vinylalcohol (PVOH), polyvinyl chloride, polylactic acid (PLA) and polyethylene terephthalate.
- 3. The microcapiUary film or foam of Claim 2, wherein said polyolefin is polyethylene or polypropylene.
- 4. The microcapiUary film or foam of Claim 2, wherein said polyamide is nylon 6.
- 5. The microcapiUary film or foam of Claim 1, wherein said one or more channels have a cross-sectional shape selected from the group consisting of circular, rectangular, oval, star, dimond, triangular, square, the like, and combinations thereof.
- 6. The microcapiUary film or foam of Claim 1, where said one or more channels and optionally the microcapiUary film or foam surface are surface treated.
- 7. A multilayer structure comprising the microcapiUary film or foam of Claim 1.
- 8. A pipe or profile comprising the microcapiUary film or foam of Claim 1.
- 9. An article comprising the microcapiUary film or foam of Claim 1.



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FIG. 5

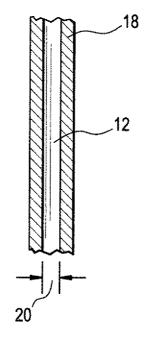
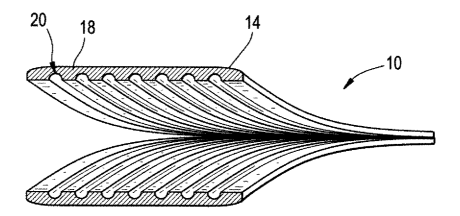


FIG. 6



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FIG. 7A

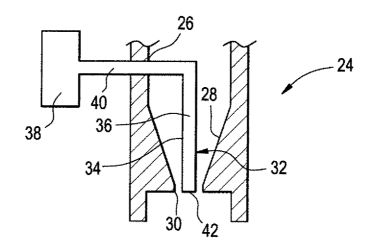
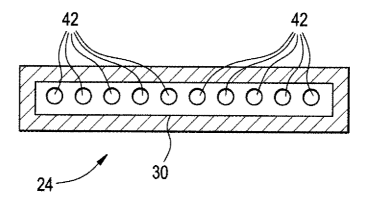
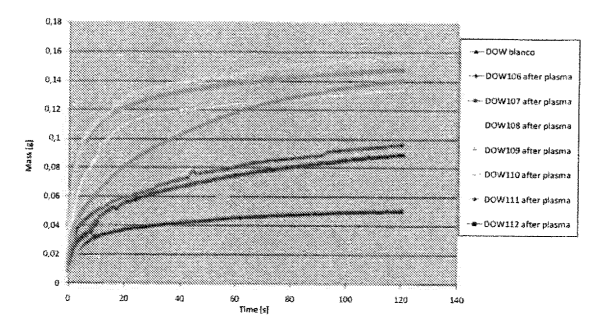


FIG. 7B



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Tensiometer evaluation of microcapillary tapes.



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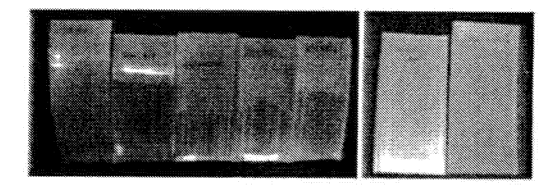


Fig. 9

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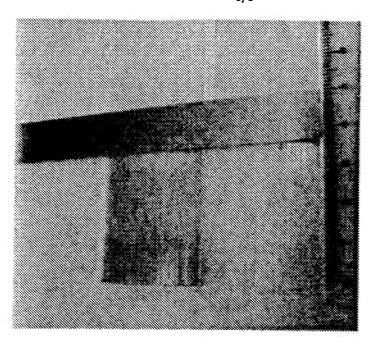


Fig. 10

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2012/07 1116

A. CLASSIFICATION OF SUBJECT MATTER C08J9/35 INV. B29D7/01 B32B3/20 C08J5/18 ADD. According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) B29D B32B C08J Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) EPO-Internal , WPI Data C. DOCUMENTS CONSIDERED TO BE RELEVANT Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X, P wo 2012/0943 15 AI (DOW GLOBAL TECHNOLOGIES 1-9 LLC [US]; K00 PMANS RUDO LF J [CH]; ZALAMEA BUST) 12 Jul y 2Q12 (2012 -07 - 12) claims; figures; examples Ε wo 2013/009538 A2 (DOW GLOBAL TECHNOLOGIES 1-9 LLC [US] ; K00 PMANS RUDO LF J [CH] ; ZALAMEA BUST) 17 January 2013 (2013 - 01 - 17) claims; figures; examples X, P W0 2012/0943 17 AI (DOW GLOBAL TECHNOLOGIES 1-9 LLC [US]; ZALAMEA BUSTILLO LUIS G [CH]; BONGAR) 12 July 2012 (20 12-07 - 12) claims; figures; exampl es -/--X Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier application or patent but published on or after the international filing date $\,$ "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other document of particular relevance; the claimed invention cannot be special reason (as specified) considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "O" document referring to an oral disclosure, use, exhibition or other "P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 18 March 2013 04/04/2013 Name and mailing address of the ISA/ Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016 Frison, Celine

INTERNATIONAL SEARCH REPORT

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PCT/US2012/07 1116

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVA Category* Citation of document, with indication, where appropria X DE 198 42 956 AI (BP CHEMI		Relevant to olaim No.
	te, of the relevant passages	Relevant to olaim No.
X DE 198 42 956 AI (BP CHEMI		
GMBH [DE]) 30 March 2000 (col umn 1, lines 1-9, 38-42) figures; examples col umn 2, lines 24-35 col umn 3, lines 3-50 col umn 3, line 67 - col umn	(2000-03 -30) ; claims;	1-9
DD 153 580 AI (HICKE HANS KONRAD; SCHWARZ HANS HARTN BRIGHT) 20 January 1982 claims; figures; examples	GEORG; FRIGGE MUT; TIERSCH	1-9

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2012/071116

Patent document cited in search report		Publication date		Patent family member(s)		Publication date
WO 2012094315	Al	12-07-2012	NONE			
Wo 2013009538	A2	17- -01- -2013	NONE			
wo 2012094317	Al	12- -07- -2012	NONE			
DE 19842956	Al	30032000	DE	19842956	Al	3 0 - 03 20 00
			ΕP	1131026	Al	1209- -2001
			WO	0016726	Al	30032000
DD 153580	Al	20011982	NONE			