Abstract:

A fiber produced by melt spinning a composition comprising an intimate admixture of (a) a thermoplastic starch; (b) a thermoplastic polymer; and (c) an oil, wax, or combination thereof present in an amount of 5 wt% to 40 wt%, based upon the total weight of the composition and where the oil, wax, or combination is dispersed through-out the thermoplastic polymer.
FIBER OF STARCH-POLYMER-OIL COMPOSITIONS

FIELD OF THE INVENTION

The present invention relates to fibers formed from compositions comprising intimate admixtures of thermoplastic starch, thermoplastic polymers and oils, waxes, or combinations thereof. The present invention also relates to methods of making these compositions.

BACKGROUND OF THE INVENTION

Thermoplastic polymers are used in a wide variety of applications. However, thermoplastic polymers, such as polypropylene and polyethylene, pose additional challenges compared to other polymer species, especially with respect to formation of, for example, fibers. This is because the material and processing requirements for production of fibers are much more stringent than for producing other forms, for example, films. For the production of fibers, polymer melt flow characteristics are more demanding on the material's physical and rheological properties vs other polymer processing methods. Also, the local shear/extensional rate and shear rate are much greater in fiber production than other processes and, for spinning very fine fibers, small defects, slight inconsistencies, or phase incompatibilities in the melt are not acceptable for a commercially viable process. Moreover, high molecular weight thermoplastic polymers cannot be easily or effectively spun into fine fibers. Given their availability and potential strength improvement, it would be desirable to provide a way to easily and effectively spin such high molecular weight polymers.

Most thermoplastic polymers, such as polyethylene, polypropylene, and polyethylene terephthalate, are derived from monomers (e.g., ethylene, propylene, and terephthalic acid, respectively) that are obtained from non-renewable, fossil-based resources (e.g., petroleum, natural gas, and coal). Thus, the price and availability of these resources ultimately have a significant impact on the price of these polymers. As the worldwide price of these resources escalates, so does the price of materials made from these polymers. Furthermore, many consumers display an aversion to purchasing products that are derived solely from petrochemicals, which are non-renewable fossil based resources. In some instances, consumers are hesitant to purchase products made from non-renewable fossil-based resources. Other consumers may have adverse perceptions about products derived from petrochemicals as being "unnatural" or not environmentally friendly.
Thermoplastic polymers and thermoplastic starches are often incompatible with, or have poor miscibility with additives (e.g., oils, pigments, organic dyes, perfumes, etc.) that might otherwise contribute to a reduced consumption of these polymers in the manufacture of downstream articles. Heretofore, the art has not effectively addressed how to reduce the amount of thermoplastic polymers derived from non-renewable, fossil-based resources in the manufacture of common articles employing these polymers. Accordingly, it would be desirable to address this deficiency. Existing art has combined polypropylene with additives, with polypropylene as the minor component to form cellular structures. These cellular structures are the purpose behind including renewable materials that are later removed or extracted after the structure is formed. U.S. Patent No. 3,093,612 describes the combination of polypropylene with various fatty acids where the fatty acid is removed. The scientific paper *J. Appl. Polym. Sci* 82 (1) pp. 169-177 (2001) discloses use of diluents on polypropylene for thermally induced phase separation to produce an open and large cellular structure but at low polymer ratio, where the diluent is subsequently removed from the final structure. The scientific paper *J. Appl. Polym. Sci* 105 (4) pp. 2000-2007 (2007) produces microporous membranes via thermally induced phase separation with dibutyl phthalate and soy bean oil mixtures, with a minor component of polypropylene. The diluent is removed in the final structure. The scientific paper *Journal of Membrane Science* 108 (1-2) pp. 25-36 (1995) produces hollow fiber microporous membranes using soy bean oil and polypropylene mixtures, with a minor component of polypropylene and using thermally induced phase separation to produce the desired membrane structure. The diluent is removed in the final structure. In all of these cases, the diluent as described is removed to produce the final structure. These structures before the diluent is removed are oily with excessive amounts of diluent to produce very open microporous structures with pore sizes > 10μm.

There have been many attempts to make nonwoven articles. However, because of costs, the difficulty in processing, and end-use properties, there are only a limited number of options. Useful fibers for nonwoven articles are difficult to produce and pose additional challenges compared to films and laminates. This is because the material and processing characteristics for fibers is much more stringent than for producing films, blow-molding articles, and injection-molding articles. For the production of fibers, the processing time during structure formation is typically much shorter and flow characteristics are more demanding on the material’s physical and rheological characteristics. The local strain rate and shear rate are much greater in fiber production than other processes. Additionally, a homogeneous composition is required for fiber spinning. For spinning very fine fibers, small defects, slight inconsistencies, or non-homogeneity
in the melt are not acceptable for a commercially viable process. The more attenuated the fibers, the more critical the processing conditions and selection of materials.

Thus, a need exists for fibers from compositions of thermoplastic polymers that allow for use of higher molecular weight and/or decreased non-renewable resource based materials, and/or incorporation of further additives, such as perfumes and dyes. A still further need is for fibers from compositions that leave the additive present to deliver renewable materials in the final product and that can also enable the addition of further additives into the final structure, such as dyes and perfumes, for example.

SUMMARY OF THE INVENTION

In one aspect, the invention is directed to fibers produced by melt spinning compositions comprising an intimate admixture of a thermoplastic starch (TPS), a thermoplastic polymer and an oil, wax, or combination thereof present in an amount of about 5 wt% to about 40 wt%, based upon the total weight of the composition. The composition can be in the form of pellets produced to be used as-is or for storage for future use, for example to make fibers. Optionally, the composition can be further processed into the final usable form, such as fibers, films and molded articles. The fibers can have a diameter of less than 200 μm. The fibers can be monocomponent or bicomponent, discrete and/or continuous, in addition to being round or shaped. The fiber can be thermally bondable.

The thermoplastic polymer can comprise a polyolefin, a polyester, a polyamide, copolymers thereof, or combinations thereof. The thermoplastic polymer can comprise polypropylene, and can have a melt flow index of greater than 0.5 g/10 min or of greater than 5 g/10 min. The thermoplastic polymer can be selected from the group consisting of polypropylene, polyethylene, polypropylene co-polymer, polyethylene co-polymer, polyethylene terephthalate, polybutylene terephthalate, polylactic acid, polyhydroxyalkanoates, polyamide-6, polyamide-6.6, and combinations thereof. The preferred thermoplastic polymer comprises polypropylene. The polypropylene can have a weight average molecular weight of about 20 kDa to about 400 kDa. The thermoplastic polymer can be present in the composition in an amount of about 20 wt% to about 90 wt%, about 30 wt% to about 70 wt%, based upon the total weight of the composition. The thermoplastic polymer can be derived from a renewable bio-based feed stock origin, such as bio polyethylene or bio polypropylene, and/or can be recycled source, such as post consumer use.
The oil, wax, or combination thereof can be present in the composition in an amount of about 5 wt% to about 40 wt%, about 8 wt% to about 30 wt%, or about 10 wt% to about 20 wt%, based upon the total weight of the composition. The oil, wax, or combination thereof can comprise a lipid, which can be selected from the group consisting of a monoglyceride, diglyceride, triglyceride, fatty acid, fatty alcohol, esterified fatty acid, epoxidized lipid, maleated lipid, hydrogenated lipid, alkyd resin derived from a lipid, sucrose polyester, or combinations thereof. The wax can be selected from the group consisting of a partially hydrogenated plant oil, a hydrogenated plant oil, an epoxidized plant oil, a maleated plant oil. Specific examples of such plant oils include soybean oil, corn oil, canola oil, and palm kernel oil. The oil, wax, or combination thereof can comprise a mineral oil or wax, such as a linear alkane, a branched alkane, or combinations thereof. The oil, wax, or combination thereof can be selected from the group consisting of soybean oil, epoxidized soybean oil, maleated soybean oil, corn oil, cottonseed oil, canola oil, beef tallow, castor oil, coconut oil, coconut seed oil, corn germ oil, fish oil, linseed oil, olive oil, oiticica oil, palm kernel oil, palm oil, palm seed oil, peanut oil, rapeseed oil, safflower oil, sperm oil, sunflower seed oil, tall oil, tung oil, whale oil, tristearin, triolein, tripalmitin, 1,2-dipalmitoolein, 1,3-dipalmitoolein, 1-palmito-3-stearo-2-olein, 1-palmito-2-stearo-3-olein, 2-palmito-1-stearo-3-olein, trilinolein, 1,2-dipalmitolinolein, 1-palmito-dilinolein, 1-stearo-dilinolein, 1,2-diacetopalmitin, 1,2-distearo-olein, 1,3-distearo-olein, trimyristin, trilaurin, capric acid, caproic acid, caprylic acid, lauric acid, lauroleic acid, linoleic acid, linolenic acid, myristic acid, myristoleic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, and combinations thereof.

The oil, wax, or combination thereof can be dispersed within the thermoplastic starch and thermoplastic polymer such that the oil, wax, or combination has a droplet size of less than 10 \( \mu \text{m} \), less than 5 \( \mu \text{m} \), less than 1 \( \mu \text{m} \), or less than 500 nm within the thermoplastic polymer. The oil, wax, or combination can be a renewable material.

The thermoplastic starch (TPS) can comprise a starch or a starch derivative and a plasticizer. The thermoplastic starch can be present in an amount about 10 wt% to about 80 wt% or about 20 wt% to about 40 wt%, based upon the total weight of the composition. The composition can be in the form of pellets produced to be used as-is or for storage for future use, for example to make fibers. Optionally, the composition can be further processed into the final usable form, such as fibers, films and molded articles.

The plasticizer can comprise a polyol. Specific polyols contemplated include mannitol, sorbitol, glycerin, and combinations thereof. The plasticizer can be selected from the group
consisting of glycerol, ethylene glycol, propylene glycol, ethylene diglycol, propylene diglycol, ethylene triglycol, propylene triglycol, polyethylene glycol, polypropylene glycol, 1,2-propanediol, 1,3-propanediol, 1,2-butanediol, 1,3-butanediol, 1,4-butanediol, 1,5-pentanediol, 1,6-hexanediol, 1,5-hexanediol, 1,2,6-hexanetriol, 1,3,5-hexanetriol, neopentyl glycol, trimethylolpropane, pentaerythritol, sorbitol, glycerol ethoxylate, tridecyl adipate, isodecyl benzoate, tributyl citrate, tributyl phosphate, dimethyl sebacate, urea, pentaerythritol ethoxylate, sorbitol acetate, pentaerythritol acetate, ethylenebisformamide, sorbitol diacetate, sorbitol monoethoxylate, sorbitol diethoxylate, sorbitol hexaethoxylate, sorbitol dipropoxylate, aminosorbitol, trihydroxymethylaminomethane, glucose/PEG, a reaction product of ethylene oxide with glucose, trimethylolpropane monoethoxylate, mannitol monoacetate, mannitol monoethoxylate, butyl glucoside, glucose monoethoxylate, a-methyl glucoside, carboxymethylsorbitol sodium salt, sodium lactate, polyglycerol monoethoxylate, erythriol, arabitol, adonitol, xylitol, mannitol, iditol, galactitol, allitol, malitol, formaide, N-methylformamide, dimethyl sulfoxide, an alkylamide, a polyglycerol having 2 to 10 repeating units, and combinations thereof.

The starch or starch derivative can be selected from the group consisting of starch, hydroxyethyl starch, hydroxypropyl starch, carboxymethylated starch, starch phosphate, starch acetate, a cationic starch, (2-hydroxy-3-trimethyl(aminomethyl)propyl) starch chloride, a starch modified by acid, base, or enzyme hydrolysis, a starch modified by oxidation, and combinations thereof.

The compositions disclosed herein can further comprise an additive. The additive can be oil soluble or oil dispersible. Examples of additives include perfume, dye, pigment, surfactant, nanoparticle, nucleating agent, clarifying agent, anti-microbial agent, antistatic agent, filler, or combination thereof.

In another aspect, provided is a method of making a composition as disclosed herein, the method comprising a) mixing the thermoplastic polymer, in a molten state, with the wax, also in the molten state, to form the admixture; and b) cooling the admixture to a temperature at or less than the solidification temperature of the thermoplastic polymer in 10 seconds or less to form the composition. The method of making a composition can comprise a) melting a thermoplastic polymer to form a molten thermoplastic polymer; b) mixing the molten thermoplastic polymer and a wax to form an admixture; and c) cooling the admixture to a temperature at or less than the solidification temperature of the thermoplastic polymer in 10 seconds or less. The mixing can be at a shear rate of greater than 10 s⁻¹, or about 30 to about 100 s⁻¹. The admixture can be cooled in
10 seconds or less to a temperature of 50°C or less. The composition can be pelletized. The
pelletizing can occur after cooling the admixture or before or simultaneous to cooling the
admixture. The composition can be made using an extruder, such as a single- or twin-screw extruder. Alternatively, the method of making a composition can comprise a) melting a
thermoplastic polymer to form a molten thermoplastic polymer; b) mixing the molten
thermoplastic polymer and a wax to form an admixture; and c) spinning the molten mixture to
form filaments or fibers which solidify upon cooling.

DETAILED DESCRIPTION OF THE INVENTION

Fibers described herein are made by melt spinning compositions disclosed herein
comprise an intimate admixture of a thermoplastic starch, thermoplastic polymer, and an oil, wax
or combination thereof. The term "intimate admixture" refers to the physical relationship of the
oil or wax, the thermoplastic starch, and thermoplastic polymer, wherein the oil or wax is
dispersed within the thermoplastic polymer and/or thermoplastic starch. The droplet size of the
oil or wax within in the thermoplastic polymer is a parameter that indicates the level of
dispersion of the oil or wax within the thermoplastic polymer and/or thermoplastic starch. The
smaller the droplet size, the higher the dispersion of the oil or wax within the thermoplastic
polymer and/or thermoplastic starch, the larger the droplet size the lower the dispersion of the oil
or wax within the thermoplastic polymer and/or thermoplastic starch. The oil, wax, or both
associate with the thermoplastic polymer, but are mixed into both the TPS and thermoplastic
polymer during formation of the compositions as disclosed herein. As used herein, the term
"admixture" refers to the intimate admixture of the present invention, and not an "admixture" in
the more general sense of a standard mixture of materials.

The droplet size of the oil or wax within the thermoplastic polymer and/or thermoplastic
starch is less than 10 µm, and can be less than 5 µm, less than 1 µm, or less than 500 nm. Other
contemplated droplet sizes of the oil and/or wax dispersed within the thermoplastic polymer
and/or thermoplastic starch include less than 9.5 µm, less than 9 µm, less than 8.5 µm, less than 8
µm, less than 7.5 µm, less than 7 µm, less than 6.5 µm, less than 6 µm, less than 5.5 µm, less
than 4.5 µm, less than 4 µm, less than 3.5 µm, less than 3 µm, less than 2.5 µm, less than 2 µm,
less than 1.5 µm, less than 900 nm, less than 800 nm, less than 700 nm, less than 600 nm, less
than 400 nm, less than 300 nm, and less than 200 nm.

The droplet size of the oil or wax can be measured by scanning electron microscopy
(SEM) indirectly by measuring a void size in the thermoplastic polymer and/or thermoplastic
starch, after removal of the oil and/or wax from the composition. Removal of the oil or wax is
typically performed prior to SEM imaging due to incompatibility of the oil or wax and the SEM imaging technique. Thus, the void measured by SEM imaging is correlated to the droplet size of the oil or wax in the composition.

One exemplary way to achieve the dispersion of the oil or wax within the thermoplastic polymer and/or thermoplastic starch is by admixing the thermoplastic polymer, in a molten state, the thermoplastic starch, in the molten state, and the oil and/or wax (which is also in the molten state). Each of the thermoplastic polymer and thermoplastic starch is melted (e.g., exposed to temperatures greater than the solidification temperature) to provide the molten thermoplastic polymer and molten thermoplastic starch, and mixed with the oil or wax. One or both of the thermoplastic polymer and thermoplastic starch can be melted prior to addition of the oil or wax or one or both can be melted in the presence of the oil or wax.

The thermoplastic polymer, thermoplastic starch, and oil or wax can be mixed, for example, at a shear rate of greater than 10s⁻¹. Other contemplated shear rates include greater than 10, about 15 to about 1000, or up to 500 s⁻¹. The higher the shear rate of the mixing, the greater the dispersion of the oil or wax in the composition as disclosed herein. Thus, the dispersion can be controlled by selecting a particular shear rate during formation of the composition.

The oil or wax and molten thermoplastic polymer and molten thermoplastic starch can be mixed using any mechanical means capable of providing the necessary shear rate to result in a composition as disclosed herein. Non-limiting examples of mechanical means include a mixer, such as a Haake batch mixer, and an extruder (e.g., a single- or twin-screw extruder).

The mixture of molten thermoplastic polymer, molten thermoplastic starch, and oil or wax is then rapidly (e.g., in less than 10 seconds) cooled to a temperature lower than the solidification temperature (either via traditional thermoplastic polymer crystallization or passing below the polymer glass transition temperature) of the thermoplastic polymer and/or thermoplastic starch. The admixture can be cooled to less than 200°C, less than 150°C, less than 100°C, less than 75°C, less than 50°C, less than 40°C, less than 30°C, less than 20°C, less than 15°C, less than 10°C, or to a temperature of about 0°C to about 30°C, about 0°C to about 20°C, or about 0°C to about 10°C. For example, the mixture can be placed in a low temperature liquid (e.g., the liquid is at or below the temperature to which the mixture is cooled) or gas. The liquid can be ambient or controlled temperature water. The gas can be ambient air or controlled temperature and humidity air. Any quenching media can be used so long as it cools the admixture rapidly. Additional liquids such as oils, alcohols and ketones can be used for quenching, along with mixtures comprising water (sodium chloride for example) depending on
the admixture composition. Additional gases can be used, such as carbon dioxide and nitrogen, or any other component naturally occurring in atmospheric temperature and pressure air.

Optionally, the composition is in the form of pellets. Pellets of the composition can be formed prior to, simultaneous to, or after cooling of the mixture. The pellets can be formed by strand cutting or underwater pelletizing. In strand cutting, the composition is rapidly quenched (generally in a time period much less than 10 seconds) then cut into small pieces. In underwater pelletizing, the mixture is cut into small pieces and simultaneously or immediately thereafter placed in the presence of a low temperature liquid that rapidly cools and solidifies the mixture to form the pelleted composition. Such pelletizing methods are well understood by the ordinarily skilled artisan. Pellet morphologies can be round or cylindrical, and can have no dimension larger than 10 mm, more preferably less than 5 mm, or no dimension larger than 2 mm.

**Thermoplastic starch**

As used herein, "thermoplastic starch" or "TPS" means a native starch or a starch derivative that has been rendered thermoplastic by treatment with one or more plasticizers. Thermoplastic starch compositions are well known and disclosed in several patents, for example: U.S. Patent Nos. 5,280,055; 5,314,934; 5,362,777; 5,844,023; 6,214,907; 6,242,102; 6,096,809; 6,218,321; 6,235,815; 6,235,816; and 6,231,970, each incorporated herein by reference.

**Starch:** The starch used in the disclosed compositions is destructurized starch. The term "thermoplastic starch" refers to destructured starch with a plasticizer.

Since natural starch generally has a granular structure, it needs to be destructurized before it can be melt processed like a thermoplastic material. For gelatinization, e.g., the process of destructuring the starch, the starch can be destructurized in the presence of a solvent which acts as a plasticizer. The solvent and starch mixture is heated, typically under pressurized conditions and shear to accelerate the gelatinization process. Chemical or enzymatic agents may also be used to destructurize, oxidize, or derivatize the starch. Commonly, starch is destructured by dissolving the starch in water. Fully destructured starch results when the particle size of any remaining undestructured starch does not impact the extrusion process, e.g., the fiber spinning process. Any remaining undestructured starch particle sizes are less than 30 μm, preferably less 20 μm, more preferably less than 10 μm, or less than 5 μm. The residual particle size can be determined by pressing the final formulation into a thin film (50 μm or less) and placing the film into a light microscope under cross polarized light. Under cross polarized light, the signature
maltese cross, indicative of undestructured starch, can be observed. If the average size of these particles is above the target range, the destructured starch has not been prepared properly.

Suitable naturally occurring starches can include, but are not limited to, corn starch, potato starch, sweet potato starch, wheat starch, sago palm starch, tapioca starch, rice starch, soybean starch, arrow root starch, bracken starch, lotus starch, cassava starch, waxy maize starch, high amylose corn starch, and commercial amylose powder. Blends of starch may also be used. Though all starches are useful herein, the present invention is most commonly practiced with natural starches derived from agricultural sources, which offer the advantages of being abundant in supply, easily replenishable and inexpensive in price. Naturally occurring starches, particularly corn starch, wheat starch, and waxy maize starch, are the preferred starch polymers of choice due to their economy and availability.

Modified starch may also be used. Modified starch is defined as non-substituted or substituted starch that has had its native molecular weight characteristics changed (i.e. the molecular weight is changed but no other changes are necessarily made to the starch). If modified starch is desired, chemical modifications of starch typically include acid or alkali hydrolysis and oxidative chain scission to reduce molecular weight and molecular weight distribution. Natural, unmodified starch generally has a very high average molecular weight and a broad molecular weight distribution (e.g. natural corn starch has an average molecular weight of up to about 60,000,000 grams/mole (g/mol)). The average molecular weight of starch can be reduced to the desirable range for the present invention by acid reduction, oxidation reduction, enzymatic reduction, hydrolysis (acid or alkaline catalyzed), physical/mechanical degradation (e.g., via the thermomechanical energy input of the processing equipment), or combinations thereof. The thermomechanical method and the oxidation method offer an additional advantage when carried out in situ. The exact chemical nature of the starch and molecular weight reduction method is not critical as long as the average molecular weight is in an acceptable range.

Ranges of number average molecular weight for starch or starch blends added to the melt can be from about 3,000 g/mol to about 20,000,000 g/mol, preferably from about 10,000 g/mol to about 10,000,000 g/mol, preferably from about 15,000 to about 5,000,000 g/mol, more preferably from about 20,000 g/mol to about 3,000,000 g/mol. In other embodiments, the average molecular weight is otherwise within the above ranges but about 1,000,000 or less, or about 700,000 or less.

Substituted starch can be used. If substituted starch is desired, chemical modifications of starch typically include etherification and esterification. Substituted starches may be desired for better compatibility or miscibility with the thermoplastic polymer and plasticizer. Alternatively,
modified and substituted starches can be used to aid in the destructuring process by increasing
the gelatinization process. However, this must be balanced with the reduction in the rate of
degradability. The degree of substitution of the chemically substituted starch is from about 0.01
to 3.0. A low degree of substitution, 0.01 to 0.06, may be preferred.

The weight of starch in the composition includes starch and its naturally occurring bound
water content. The term "bound water" means the water found naturally occurring in starch and
before mixing of starch with other components to make the composition of the present invention.
The term "free water" means the water that is added in making the composition of the present
invention. A person of ordinary skill in the art would recognize that once the components are
mixed in a composition, water can no longer be distinguished by its origin. The starch typically
has a bound water content of about 5% to 16% by weight of starch. It is known that additional
free water may be incorporated as the polar solvent or plasticizer, and not included in the weight
of the starch.

**Plasticizer:** A plasticizer can be used in the present invention to destructurize the starch
and enable the starch to flow, i.e. create a thermoplastic starch. The same plasticizer may be
used to increase melt processability or two separate plasticizers may be used. The plasticizers
may also improve the flexibility of the final products, which is believed to be due to the lowering
of the glass transition temperature of the composition by the plasticizer. The plasticizers should
preferably be substantially compatible with the polymeric components of the disclosed
compositions so that the plasticizers may effectively modify the properties of the composition.
As used herein, the term "substantially compatible" means when heated to a temperature above
the softening and/or the melting temperature of the composition, the plasticizer is capable of
forming a substantially homogeneous mixture with starch.

An additional plasticizer or diluent for the thermoplastic polymer may be present to lower
the polymer's melting temperature and improve overall compatibility with the thermoplastic
starch blend. Furthermore, thermoplastic polymers with higher melting temperatures may be used
if plasticizers or diluents are present which suppress the melting temperature of the polymer. The
plasticizer will typically have a molecular weight of less than about 100,000 g/mol and may
preferably be a block or random copolymer or terpolymer where one or more of the chemical
species is compatible with another plasticizer, starch, polymer, or combinations thereof.

Nonlimiting examples of useful hydroxyl plasticizers include sugars such as glucose,
sucrose, fructose, raffinose, maltodextrose, galactose, xylose, maltose, lactose, mannose
erythrose, glycerol, and pentaerythritol; sugar alcohols such as erythritol, xylitol, malitol,
mannitol and sorbitol; polyols such as ethylene glycol, propylene glycol, dipropylene glycol, butylene glycol, hexane triol, and the like, and polymers thereof; and mixtures thereof. Also useful herein as hydroxyl plasticizers are poloxomers and poloxamines. Also suitable for use herein are hydrogen bond forming organic compounds which do not have hydroxyl group, including urea and urea derivatives; anhydrides of sugar alcohols such as sorbitan; animal proteins such as gelatin; vegetable proteins such as sunflower protein, soybean proteins, cotton seed proteins; and mixtures thereof. Other suitable plasticizers are phthalate esters, dimethyl and diethylsuccinate and related esters, glycerol triacetate, glycerol mono and diacetates, glycerol mono, di, and tripropionates, and butanoates, which are biodegradable. Aliphatic acids such as ethylene acrylic acid, ethylene maleic acid, butadiene acrylic acid, butadiene maleic acid, propylene acrylic acid, propylene maleic acid, and other hydrocarbon based acids. All of the plasticizers may be use alone or in mixtures thereof.

Preferred plasticizers include glycerin, mannitol, and sorbitol, with sorbitol being the most preferred. The amount of plasticizer is dependent upon the molecular weight, amount of starch, and the affinity of the plasticizer for the starch. Generally, the amount of plasticizer increases with increasing molecular weight of starch.

The thermoplastic starch can be present in the compositions disclosed herein in a weight percent of about 10 wt% to about 80 wt%, about 10 wt% to about 60 wt%, or about 20 wt% to about 40 wt%, based upon the total weight of the composition. Specific contemplated amounts of thermoplastic starch include about 10 wt%, about 11 wt%, about 12 wt%, about 13 wt%, about 14 wt%, about 15 wt%, about 16 wt%, about 17 wt%, about 18 wt%, about 19 wt%, about 20 wt%, about 21 wt%, about 22 wt%, about 23 wt%, about 24 wt%, about 25 wt%, about 26 wt%, about 27 wt%, about 28 wt%, about 29 wt%, about 30 wt%, about 31 wt%, about 32 wt%, about 33 wt%, about 34 wt%, about 35 wt%, about 36 wt%, about 37 wt%, about 38 wt%, about 39 wt%, about 40 wt%, about 41 wt%, about 42 wt%, about 43 wt%, about 44 wt%, about 45 wt%, about 46 wt%, about 47 wt%, about 48 wt%, about 49 wt%, about 50 wt%, about 51 wt%, about 52 wt%, about 53 wt%, about 54 wt%, about 55 wt%, about 56 wt%, about 57 wt%, about 58 wt%, about 59 wt%, about 60 wt%, about 61 wt%, about 62 wt%, about 63 wt%, about 64 wt%, about 65 wt%, about 66 wt%, about 67 wt%, about 68 wt%, about 69 wt%, about 70 wt%, about 71 wt%, about 72 wt%, about 73 wt%, about 74 wt%, about 75 wt%, about 76 wt%, about 77 wt%, about 78 wt%, about 79 wt%, and about 80 wt%, based upon the total weight of the composition.
**Thermoplastic polymers**

Thermoplastic polymers, as used in the disclosed compositions, are polymers that melt and then, upon cooling, crystallize or harden, but can be re-melted upon further heating. Suitable thermoplastic polymers used herein have a melting temperature (also referred to as solidification temperature) from about 60°C to about 300°C, from about 80°C to about 250°C, or from 100°C to 215°C, with the preferred range from 100°C to 180°C.

The molecular weight of the thermoplastic polymer is sufficiently high to enable entanglement between polymer molecules and yet low enough to be melt spinnable. Addition of the oil into the composition allows for compositions containing higher molecular weight thermoplastic polymers to be processed, compared to compositions without an oil. Thus, suitable thermoplastic polymers can have weight average molecular weights of about 1000 kDa or less, about 5 kDa to about 800 kDa, about 10 kDa to about 700 kDa, or about 20 kDa to about 400 kDa.

The thermoplastic polymers can be derived from renewable resources or from fossil minerals and oils. The thermoplastic polymers derived from renewable resources are bio-based, for example such as bio produced ethylene and propylene monomers used in the production polypropylene and polyethylene. These material properties are essentially identical to fossil based product equivalents, except for the presence of carbon-14 in the thermoplastic polymer. Renewable and fossil based thermoplastic polymers can be combined together in the present invention in any ratio, depending on cost and availability. Recycled thermoplastic polymers can also be used, alone or in combination with renewable and/or fossil derived thermoplastic polymers. The recycled thermoplastic polymers can be pre-conditioned to remove any unwanted contaminants prior to compounding or they can be used during the compounding and extrusion process, as well as simply left in the admixture. These contaminants can include trace amounts of other polymers, pulp, pigments, inorganic compounds, organic compounds and other additives typically found in processed polymeric compositions. The contaminants should not negatively impact the final performance properties of the admixture, for example, causing spinning breaks during a fiber spinning process.

Suitable thermoplastic polymers generally include polyolefins, polyesters, polyamides, copolymers thereof, and combinations thereof. The thermoplastic polymer can be selected from the group consisting of polypropylene, polyethylene, polypropylene co-polymer, polyethylene co-polymer, polyethylene terephthalate, polybutylene terephthalate, polylactic acid, polyhydroxyalkanoates, polyamide-6, polyamide-6,6, and combinations thereof. The polymer
can be polypropylene based, polyethylene based, polyhydroxyalkanoate based polymer systems, copolymers and combinations thereof.

More specifically, however, the thermoplastic polymers preferably include polyolefins such as polyethylene or copolymers thereof, including low, high, linear low, or ultra low density polyethylenes, polypropylene or copolymers thereof, including atactic polypropylene; isotactic polypropylene, metallocene isotactic polypropylene, polybutylene or copolymers thereof; polyamides or copolymers thereof, such as Nylon 6, Nylon 11, Nylon 12, Nylon 46, Nylon 66; polyesters or copolymers thereof, such as maleic anhydride polypropylene copolymer, polyethylene terephthalate; olefin carboxylic acid copolymers such as ethylene/acrylic acid copolymer, ethylene/maleic acid copolymer, ethylene/methacrylic acid copolymer, ethylene/vinyl acetate copolymers or combinations thereof; polyacrylates, polymethacrylates, and their copolymers such as poly(methyl methacrylates). Other nonlimiting examples of polymers include polycarbonates, polyvinyl acetates, poly(oxymethylene), styrene copolymers, polycrylates, polymethacrylates, poly(methyl methacrylates), polystyrene/methyl methacrylate copolymers, polyetherimides, polysulfones, or combinations thereof. In some embodiments, thermoplastic polymers include polypropylene, polyethylene, polyamides, polyvinyl alcohol, ethylene acrylic acid, polyolefin carboxylic acid copolymers, polyesters, and combinations thereof.

More specifically, however, the thermoplastic polymers preferably include polyolefins such as polyethylene or copolymers thereof, including low density, high density, linear low density, or ultra low density polyethylenes such that the polyethylene density ranges between 0.90 grams per cubic centimeter to 0.97 grams per cubic centimeter, most preferred between 0.92 and 0.95 grams per cubic centimeter. The density of the polyethylene will be determined by the amount and type of branching and depends on the polymerization technology and comonomer type. Polypropylene and/or polypropylene copolymers, including atactic polypropylene; isotactic polypropylene, syndiotactic polypropylene, and combination thereof can also be used. Polypropylene copolymers, especially ethylene can be used to lower the melting temperature and improve properties. These polypropylene polymers can be produced using metallocene and Ziegler-Natta catalyst systems. These polypropylene and polyethylene compositions can be combined together to optimize end-use properties. Polybutylene is also a useful polyolefin.

Biodegradable thermoplastic polymers also are contemplated for use herein. Biodegradable materials are susceptible to being assimilated by microorganisms, such as molds, fungi, and bacteria when the biodegradable material is buried in the ground or otherwise contacts
the microorganisms (including contact under environmental conditions conducive to the growth of the microorganisms). Suitable biodegradable polymers also include those biodegradable materials which are environmentally-degradable using aerobic or anaerobic digestion procedures, or by virtue of being exposed to environmental elements such as sunlight, rain, moisture, wind, temperature, and the like. The biodegradable thermoplastic polymers can be used individually or as a combination of biodegradable or non-biodegradable polymers. Biodegradable polymers include polyesters containing aliphatic components. Among the polyesters are ester polycondensates containing aliphatic constituents and poly(hydroxycarboxylic) acid. The ester polycondensates include diacids/diol aliphatic polyesters such as polybutylene succinate, polybutylene succinate co-adipate, aliphatic/aromatic polyesters such as terpolymers made of butylene diol, adipic acid and terephthalic acid. The poly(hydroxycarboxylic) acids include lactic acid based homopolymers and copolymers, polyhydroxybutyrate (PHB), or other polyhydroxyalkanoate homopolymers and copolymers. Such polyhydroxyalkanoates include copolymers of PHB with higher chain length monomers, such as C₆-C₁₂ and higher, polyhydroxyalkanoates, such as those disclosed in U.S. Patent Nos. RE 36,548 and 5,990,271.

An example of a suitable commercially available polylactic acid is NATUREWORKS from Cargill Dow and LACEA from Mitsui Chemical. An example of a suitable commercially available diacid/diol aliphatic polyester is the polybutylene succinate/adipate copolymers sold as BIONOLLE 1000 and BIONOLLE 3000 from the Showa High Polymer Company, Ltd. (Tokyo, Japan). An example of a suitable commercially available aliphatic/aromatic copolyester is the poly(tetramethylene adipate-co-terephthalate) sold as EASTAR BIO Copolyester from Eastman Chemical or ECOFLEX from BASF.

Non-limiting examples of suitable commercially available polypropylene or polypropylene copolymers include Basell Profax PH-835 (a 35 melt flow rate Ziegler-Natta isotactic polypropylene from Lyondell-Basell), Basell Metocene MF-650W (a 500 melt flow rate metalloocene isotactic polypropylene from Lyondell-Basell), Polybond 3200 (a 250 melt flow rate maleic anhydride polypropylene copolymer from Crompton), Exxon Achieve 3854 (a 25 melt flow rate metalloocene isotactic polypropylene from Exxon-Mobil Chemical), Mosten NB425 (a 25 melt flow rate Ziegler-Natta isotactic polypropylene from Unipetrol), Danimer 27510 (a polyhydroxyalkanoate polypropylene from Danimer Scientific LLC), Dow Aspun 681 1A (a 27 melt index polyethylene polypropylene copolymer from Dow Chemical), and Eastman 9921 (a polyester terephthalic homopolymer with a nominally 0.81 intrinsic viscosity from Eastman Chemical).
The thermoplastic polymer component can be a single polymer species as described above or a blend of two or more thermoplastic polymers as described above.

If the polymer is polypropylene, the thermoplastic polymer can have a melt flow index of greater than 5 g/10 min, as measured by ASTM D-1238, used for measuring polypropylenes.

Other contemplated melt flow indices include greater than 10 g/10 min, greater than 20 g/10 min, or about 5 g/10 min to about 50 g/10 min.

**Oils and Waxes**

An oil or wax, as used in the disclosed composition, is a lipid, mineral oil (or wax), or combination thereof. An oil is used to refer to a compound that is liquid at room temperature (e.g., has a melting point of 25°C or less) while a wax is used to refer to a compound that is a solid at room temperature (e.g., has a melting point of greater than 25°C). The wax can also have a melting point lower than the melting temperature of the highest volumetric polymer component in the composition. The term wax hereafter can refer to the component either in the solid crystalline state or in the molten state, depending on the temperature. The wax can be solid at a temperature at which the thermoplastic polymer and/or thermoplastic starch are solid. For example, polypropylene is a semicrystalline solid at 90°C, which can be above melting temperature of the wax.

A wax, as used in the disclosed composition, is a lipid, mineral wax, or combination thereof, wherein the lipid, mineral wax, or combination thereof has a melting point of greater than 25°C. More preferred is a melting point above 35°C, still more preferred above 45°C and most preferred above 50°C. The wax can have a melting point that is lower than the melting temperature of the thermoplastic polymer in the composition. The terms "wax" and "oil" are differentiated by crystallinity of the component at or near 25°C. In all cases, the "wax" will have a maximum melting temperature less than the thermoplastic polymer, preferably less than 100°C and most preferably less than 80°C. The wax can be a lipid. The lipid can be a monoglyceride, diglyceride, triglyceride, fatty acid, fatty alcohol, esterified fatty acid, epoxidized lipid, maleated lipid, hydrogenated lipid, alkyd resin derived from a lipid, sucrose polyester, or combinations thereof. The mineral wax can be a linear alkane, a branched alkane, or combinations thereof. The waxes can be partially or fully hydrogenated materials, or combinations and mixtures thereof, that were formally liquids at room temperature in their unmodified forms. When the temperature is above the melting temperature of the wax, it is a liquid oil. When in the molten state, the wax can be referred to as an "oil". The terms "wax" and "oil" only have meaning when
measured at 25°C. The wax will be a solid at 25°C, while an oil is not a solid at 25°C. Otherwise they are used interchangeably above 25°C.

The lipid can be a monoglyceride, diglyceride, triglyceride, fatty acid, fatty alcohol, esterified fatty acid, epoxidized lipid, maleated lipid, hydrogenated lipid, alkyd resin derived from a lipid, sucrose polyester, or combinations thereof. The mineral oil or wax can be a linear alkane, a branched alkane, or combinations thereof. The waxes can be partially or fully hydrogenated materials, or combinations and mixtures thereof, that were formally liquids at room temperature in their unmodified forms.

Non-limiting examples of oils or waxes contemplated in the compositions disclosed herein include beef tallow, castor oil, coconut oil, coconut seed oil, corn germ oil, cottonseed oil, fish oil, linseed oil, olive oil, oiticica oil, palm kernel oil, palm oil, palm seed oil, peanut oil, rapeseed oil, safflower oil, soybean oil, sperm oil, sunflower seed oil, tall oil, tung oil, whale oil, and combinations thereof. Non-limiting examples of specific triglycerides include triglycerides such as, for example, tristearin, triolein, tripalmitin, 1,2-dipalmitoolein, 1,3-dipalmitoolein, 1-palmito-3-stearo-2-olein, 1-palmito-2- stearo-3-olein, 2-palmito-1-stearo-3-olein, trilinolein, 1,2-dipalmilitolnolein, 1-palmito-dilnolein, 1-stearo- dilnolein, 1,2-diacetopalmitin, 1,2-distearo- olein, 1,3-distearo-olein, trimyristin, trilaurin and combinations thereof. Non-limiting examples of specific fatty acids contemplated include capric acid, caprylic acid, lauric acid, lauroleic acid, linoleic acid, linolenic acid, myristic acid, myristoleic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, and mixtures thereof.

Because the wax may contain a distribution of melting temperatures to generate a peak melting temperature, the wax melting temperature is defined as having a peak melting temperature 25°C or above as defined as when > 50 weight percent of the wax component melts at or above 25°C. This measurement can be made using a differential scanning calorimeter (DSC), where the heat of fusion is equated to the weight percent fraction of the wax.

The wax number average molecular weight, as determined by gel permeation chromatography (GPC), should be less than 2kDa, preferably less than 1.5kDa, still more preferred less than 1.2kDa.

The amount of wax is determined via gravimetric weight loss method. The solidified mixture is placed, with the narrowest specimen dimension no greater than 1mm, into acetone at a ratio of 1g or mixture per 100g of acetone using a refluxing flask system. First the mixture is weighed before being placed into the reflux flask, and then the acetone and mixtures are heated
to 60°C for 20 hours. The sample is removed and air dried for 60 minutes and a final weight
determined. The equation for calculating the weight percent wax is

\[
\text{weight \% wax} = \frac{\text{[initial mass-final mass]}}{\text{[initial mass]}} \times 100\%
\]

Because the oil may contain a distribution of melting temperatures to generate a peak melting temperature, the oil melting temperature is defined as having a peak melting temperature 25°C or below as defined when > 50 weight percent of the oil component melts at or below 25°C. This measurement can be made using a differential scanning calorimeter (DSC), where the heat of fusion is equated to the weight percent fraction of the oil.

The oil number average molecular weight, as determined by gel permeation chromatography (GPC), should be less than 2kDa, preferably less than 1.5kDa, still more preferred less than 1.2kDa.

The oil or wax can be from a renewable material (e.g., derived from a renewable resource). As used herein, a "renewable resource" is one that is produced by a natural process at a rate comparable to its rate of consumption (e.g., within a 100 year time frame). The resource can be replenished naturally, or via agricultural techniques. Non-limiting examples of renewable resources include plants (e.g., sugar cane, beets, corn, potatoes, citrus fruit, woody plants, lignocellulosics, hemicellulosics, cellulosic waste), animals, fish, bacteria, fungi, and forestry products. These resources can be naturally occurring, hybrids, or genetically engineered organisms. Natural resources such as crude oil, coal, natural gas, and peat, which take longer than 100 years to form, are not considered renewable resources. Mineral oil, petroleum, and petroleum jelly are viewed as a by-product waste stream of coal, and while not renewable, it can be considered a by-product oil.

Specific examples of mineral wax include paraffin (including petrolatum), Montan wax, as well as polyolefin waxes produced from cracking processes, preferentially polyethylene derived waxes. Mineral waxes and plant derived waxes can be combined together. Plant based waxes can be differentiated by their carbon-14 content.

The oil or wax, as disclosed herein, is present in the composition at a weight percent of about 5 wt% to about 40 wt%, based upon the total weight of the composition. Other contemplated wt% ranges of the oil or wax include about 8 wt% to about 30 wt%, with a preferred range from about 10 wt% to about 30 wt%, about 10 wt% to about 20 wt%, or about 12 wt% to about 18 wt%, based upon the total weight of the composition. Specific oil or wax wt% contemplated include about 5 wt%, about 6 wt%, about 7 wt%, about 8 wt%, about 9 wt%, about
10 wt%, about 11 wt%, about 12 wt%, about 13 wt%, about 14 wt%, about 15 wt%, about 16 wt%, about 17 wt%, about 18 wt%, about 19 wt%, about 20 wt%, about 21 wt%, about 22 wt%, about 23 wt%, about 24 wt%, about 25 wt%, about 26 wt%, about 27 wt%, about 28 wt%, about 29 wt%, about 30 wt%, about 31 wt%, about 32 wt%, about 33 wt%, about 34 wt%, about 35 wt%, about 36 wt%, about 37 wt%, about 38 wt%, about 39 wt%, and about 40 wt%, based upon the total weight of the composition.

**Additives**

The compositions disclosed herein can further include an additive. The additive can be dispersed throughout the composition, or can be substantially in the thermoplastic polymer portion of the thermoplastic layer, substantially in the oil portion of the composition, or substantially in the TPS portion of the composition. In cases where the additive is in the oil portion of the composition, the additive can be oil soluble or oil dispersible. Alkyd resins can also be added to the composition. Alkyd resins comprise, for example, polyols, polyacids, and/or anhydrides.

Non-limiting examples of classes of additives contemplated in the compositions disclosed herein include perfumes, dyes, pigments, nanoparticles, antistatic agents, fillers, and combinations thereof. The compositions disclosed herein can contain a single additive or a mixture of additives. For example, both a perfume and a colorant (e.g., pigment and/or dye) can be present in the composition. The additive(s), when present, is/are present in a weight percent of about 0.05 wt% to about 20 wt%, or about 0.1 wt% to about 10 wt%. Specifically contemplated weight percentages include about 0.5 wt%, about 0.6 wt%, about 0.7 wt%, about 0.8 wt%, about 0.9 wt%, about 1 wt%, about 1.1 wt%, about 1.2 wt%, about 1.3 wt%, about 1.4 wt%, about 1.5 wt%, about 1.6 wt%, about 1.7 wt%, about 1.8 wt%, about 1.9 wt%, about 2 wt%, about 2.1 wt%, about 2.2 wt%, about 2.3 wt%, about 2.4 wt%, about 2.5 wt%, about 2.6 wt%, about 2.7 wt%, about 2.8 wt%, about 2.9 wt%, about 3 wt%, about 3.1 wt%, about 3.2 wt%, about 3.3 wt%, about 3.4 wt%, about 3.5 wt%, about 3.6 wt%, about 3.7 wt%, about 3.8 wt%, about 3.9 wt%, about 4 wt%, about 4.1 wt%, about 4.2 wt%, about 4.3 wt%, about 4.4 wt%, about 4.5 wt%, about 4.6 wt%, about 4.7 wt%, about 4.8 wt%, about 4.9 wt%, about 5 wt%, about 5.1 wt%, about 5.2 wt%, about 5.3 wt%, about 5.4 wt%, about 5.5 wt%, about 5.6 wt%, about 5.7 wt%, about 5.8 wt%, about 5.9 wt%, about 6 wt%, about 6.1 wt%, about 6.2 wt%, about 6.3 wt%, about 6.4 wt%, about 6.5 wt%, about 6.6 wt%, about 6.7 wt%, about 6.8 wt%, about 6.9 wt%, about 7 wt%, about 7.1 wt%, about 7.2 wt%, about 7.3 wt%, about 7.4 wt%, about 7.5 wt%, about 7.6 wt%, about 7.7 wt%, about 7.8 wt%, about 7.9 wt%, about 8
wt%, about 8.1 wt%, about 8.2 wt%, about 8.3 wt%, about 8.4 wt%, about 8.5 wt%, about 8.6 wt%, about 8.7 wt%, about 8.8 wt%, about 8.9 wt%, about 9 wt%, about 9.1 wt%, about 9.2 wt%, about 9.3 wt%, about 9.4 wt%, about 9.5 wt%, about 9.6 wt%, about 9.7 wt%, about 9.8 wt%, about 9.9 wt%, and about 10 wt%.

As used herein the term "perfume" is used to indicate any odoriferous material that is subsequently released from the composition as disclosed herein. A wide variety of chemicals are known for perfume uses, including materials such as aldehydes, ketones, alcohols, and esters. More commonly, naturally occurring plant and animal oils and exudates including complex mixtures of various chemical components are known for use as perfumes. The perfumes herein can be relatively simple in their compositions or can include highly sophisticated complex mixtures of natural and synthetic chemical components, all chosen to provide any desired odor. Typical perfumes can include, for example, woody/earthy bases containing exotic materials, such as sandalwood, civet and patchouli oil. The perfumes can be of a light floral fragrance (e.g. rose extract, violet extract, and lilac). The perfumes can also be formulated to provide desirable fruity odors, e.g. lime, lemon, and orange. The perfumes delivered in the compositions and articles of the present invention can be selected for an aromatherapy effect, such as providing a relaxing or invigorating mood. As such, any material that exudes a pleasant or otherwise desirable odor can be used as a perfume active in the compositions and articles of the present invention.


Contemplated fillers include, but are not limited to inorganic fillers such as, for example, the oxides of magnesium, aluminum, silicon, and titanium. These materials can be added as inexpensive fillers or processing aides. Other inorganic materials that can function as fillers include hydrous magnesium silicate, titanium dioxide, calcium carbonate, clay, chalk, boron
nitride, limestone, diatomaceous earth, mica glass quartz, and ceramics. Additionally, inorganic salts, including alkali metal salts, alkaline earth metal salts, phosphate salts, can be used.

Contemplated surfactants include anionic surfactants, amphoteric surfactants, or a combination of anionic and amphoteric surfactants, and combinations thereof, such as surfactants disclosed, for example, in U.S. Patent Nos. 3,929,678 and 4,259,217 and in EP 414 549, WO93/08876 and WO93/08874.

Contemplated nanoparticles include metals, metal oxides, allotropes of carbon, clays, organically modified clays, sulfates, nitrates, hydroxides, oxy/hydroxides, particulate water-insoluble polymers, silicates, phosphates and carbonates. Examples include silicon dioxide, carbon black, graphite, grapheme, fullerenes, expanded graphite, carbon nanotubes, talc, calcium carbonate, betonite, montmorillonite, kaolin, silica, aluminosilicates, boron nitride, aluminum nitride, barium sulfate, calcium sulfate, antimony oxide, feldspar, mica, nickel, copper, iron, cobalt, steel, gold, silver, platinum, aluminum, wollastonite, aluminum oxide, zirconium oxide, titanium dioxide, cerium oxide, zinc oxide, magnesium oxide, tin oxide, iron oxides (Fe₃O₄) and mixtures thereof. Nanoparticles can increase strength, thermal stability, and/or abrasion resistance of the compositions disclosed herein, and can give the compositions electric properties.

Additional contemplated additives include nucleating and clarifying agents for the thermoplastic polymer. Specific examples, suitable for polypropylene, for example, are benzoic acid and derivatives (e.g. sodium benzoate and lithium benzoate), as well as kaolin, talc and zinc glycerolate. Dibenzaldene sorbitol (DBS) is an example of a clarifying agent that can be used. Other nucleating agents that can be used are organocarboxylic acid salts, sodium phosphate and metal salts (for example aluminum dibenzoate) The nucleating or clarifying agents can be added in ranges from 20 parts per million (20ppm) to 20,000ppm, more preferred range of 200ppm to 2000ppm and the most preferred range from 1000ppm to 1500ppm. The addition of the nucleating agent can be used to improve the tensile and impact properties of the finished admixture composition.

Contemplated anti-static agents include fabric softeners which are known to provide antistatic benefits. For example those fabric softeners that have a fatty acyl group which has an iodine value of above 20, such as N,N-di(tallowoyl-oxy-ethyl)-N,N-dimethyl ammonium methylsulfate.
Fibers

The fibers in the present invention may be monocomponent or multicomponent. The term "fiber" is defined as a solidified polymer shape with a length to thickness ratio of greater than 1,000. The monocomponent fibers of the present invention may also be multiconstituent. Multiconstituent fiber, as used herein, is defined to mean a fiber containing more than one chemical species or material. Multiconstituent and alloyed polymers have the same meaning in the present invention and can be used interchangeably. Generally, fibers may be of monocomponent or multicomponent types. Component, as used herein, is defined as a separate part of the fiber that has a spatial relationship to another part of the fiber. The term multicomponent, as used herein, is defined as a fiber having more than one separate part in spatial relationship to one another. The term multicomponent includes bicomponent, which is defined as a fiber having two separate parts in a spatial relationship to one another. The different components of multicomponent fibers are arranged in substantially distinct regions across the cross-section of the fiber and extend continuously along the length of the fiber. Methods for making multicomponent fibers are well known in the art. Multicomponent fiber extrusion was well known in the 1960's. DuPont was a lead technology developer of multicomponent capability, with US 3,244,785 and US 3,704,971 providing a technology description of the technology used to make these fibers. "Bicomponent Fibers" by R. Jeffries from Merrow Publishing in 1971 laid a solid groundwork for bicomponent technology. More recent publications include "Taylor-Made Polypropylene and Bicomponent Fibers for the Nonwoven Industry," Tappi Journal December 1991 (pl03) and "Advanced Fiber Spinning Technology" edited by Nakajima from Woodhead Publishing.

The nonwoven fabric formed in the present invention may contain multiple types of monocomponent fibers that are delivered from different extrusion systems through the same spinneret. The extrusion system, in this example, is a multicomponent extrusion system that delivers different polymers to separate capillaries. For instance, one extrusion system would deliver polypropylene with wax and the other a polypropylene copolymer such that the copolymer composition melts at different temperatures. In a second example, one extrusion system might deliver a polyethylene resin and the other polypropylene with wax. In a third example, one extrusion system might deliver a polypropylene resin with 30weight percent wax and the other a polypropylene resin with 30weight percent wax that has a molecular weight different from the first polypropylene resin. The polymer ratios in this system can range from 95:5 to 5:95, preferably from 90:10 to 10:90 and 80:20 to 20:80.
Bicomponent and multicomponent fibers may be in a side-by-side, sheath-core (symmetric and eccentric), segmented pie, ribbon, islands-in-the-sea configuration, or any combination thereof. The sheath may be continuous or non-continuous around the core. Non-inclusive examples of exemplarily multicomponent fibers are disclosed in US Patent 6,746,766. The ratio of the weight of the sheath to the core is from about 5:95 to about 95:5. The fibers of the present invention may have different geometries that include, but are not limited to; round, elliptical, star shaped, trilobal, multilobal with 3-8 lobes, rectangular, H-shaped, C-shaped, I-shape, U-shaped and other various eccentricities. Hollow fibers can also be used. Preferred shapes are round, trilobal and H-shaped. The round and trilobal fiber shapes can also be hollow.

Sheath and core bicomponent fibers are preferred. In one preferred case, the component in the core may contain the thermoplastic polymer and wax, while the sheath does not. In this case the exposure to wax at the surface of the fiber is reduced or eliminated. In another preferred case, the sheath may contain the wax and the core does not. In this case the concentration of wax at the fiber surface is higher than in the core. Using sheath and core bicomponent fibers, the concentration of the wax can be selected to impart desired properties either in the sheath or core, or some concentration gradient. It should be understood that islands-in-a-sea bicomponent fibers are considered to be a type of sheath and core fiber, but with multiple cores. Segmented pie fibers (hollow and solid) are contemplated. For one example, to split regions that contain wax from regions that do not contain wax using segmented pie type of bicomponent fiber design.

Splitting may occur during mechanical deformation, application of hydrodynamic forces or other suitable processes.

Tricomponent fibers are also contemplated. One example of a useful tricomponent fiber would be a three layered sheath/sheath/core fiber, where each component contains a different amount of wax. Different amounts of wax in each layer may provide additional benefits. For example, the core can be a blend of 10 melt flow polypropylene with 30 weight percent wax. The middle layer sheath may be a blend of 25 melt flow polypropylene with 20 weight percent wax and the outer layer may be straight 35 melt flow rate polypropylene. It is preferred that the wax content between each layer is less than 40 wt%, more preferably less than 20wt%. Another type of useful tricomponent fiber contemplated is a segmented pie type bicomponent design that also has a sheath.

A "highly attenuated fiber" is defined as a fiber having a high draw down ratio. The total fiber draw down ratio is defined as the ratio of the fiber at its maximum diameter (which is
typically results immediately after exiting the capillary) to the final fiber diameter in its end use. The total fiber draw down ratio will be greater than 1.5, preferably greater than 5, more preferably greater than 10, and most preferably greater than 12. This is necessary to achieve the tactile properties and useful mechanical properties.

The fiber will have a diameter of less than 200 µm. The fiber diameter can be as low as 0.1 µm if the mixture is being used to produce fine fibers. The fibers can be either essentially continuous or essentially discontinuous. Fibers commonly used to make spunbond nonwovens will have a diameter of from about 5 µm to about 30 µm, more preferably from 10 µm to about 20 µm and most preferred from 12 µm to about 18 µm. Fine fiber diameter will have a diameter from 0.1 µm to about 5 µm, preferably from 0.2 µm to about 3 µm and most preferred from 0.3 µm to about 2 µm. Fiber diameter is controlled by die geometry, spinning speed or drawing speed, mass through-put, and blend composition and rheology. The fibers as described herein can be environmentally degradable.

The fibers described herein are typically used to make disposable nonwoven articles. The articles are commonly flushable. The term "flushable" as used herein refers to materials which are capable of dissolving, dispersing, disintegrating, and/or decomposing in a septic disposal system such as a toilet to provide clearance when flushed down the toilet without clogging the toilet or any other sewage drainage pipe. The fibers and resulting articles may also be aqueous responsive. The term aqueous responsive as used herein means that when placed in water or flushed, an observable and measurable change will result. Typical observations include noting that the article swells, pulls apart, dissolves, or observing a general weakened structure.

The hydrophilicity and hydrophobicity of the fibers can be adjusted in the present invention. The base resin properties can have hydrophilic properties via copolymerization (such as the case for certain polyesters (EASTONE from Eastman Chemical, the sulfopolyester family of polymers in general) or polyolefins such as polypropylene or polyethylene) or have materials added to the base resin to render it hydrophilic. Exemplarily examples of additives include CIBA Irgasurf® family of additives. The fibers in the present invention can also be treated or coated after they are made to render them hydrophilic. In the present invention, durable hydrophilicity is preferred. Durable hydrophilicity is defined as maintaining hydrophilic characteristics after more than one fluid interaction. For example, if the sample being evaluated is tested for durable hydrophilicity, water can be poured on the sample and wetting observed. If the sample wets out it is initially hydrophilic. The sample is then completely rinsed with water and dried. The
rinsing is best done by putting the sample in a large container and agitating for ten seconds and then drying. The sample after drying should also wet out when contacted again with water.

After the fiber is formed, the fiber may further be treated or the bonded fabric can be treated. A hydrophilic or hydrophobic finish can be added to adjust the surface energy and chemical nature of the fabric. For example, fibers that are hydrophobic may be treated with wetting agents to facilitate absorption of aqueous liquids. A bonded fabric can also be treated with a topical solution containing surfactants, pigments, slip agents, salt, or other materials to further adjust the surface properties of the fiber.

The fibers in the present invention can be crimped, although it is preferred that they are not crimped. Crimped fibers are generally produced in two methods. The first method is mechanical deformation of the fiber after it is already spun. Fibers are melt spun, drawn down to the final filament diameter and mechanically treated, generally through gears or a stuffer box that imparts either a two dimensional or three dimensional crimp. This method is used in producing most carded staple fibers. The second method for crimping fibers is to extrude multicomponent fibers that are capable of crimping in a spunlaid process. One of ordinary skill in the art would recognize that a number of methods of making bicomponent crimped spunbond fibers exists; however, for the present invention, three main techniques are considered for making crimped spunlaid nonwovens. The first is crimping that occurs in the spinline due to differential polymer crystallization in the spinline, a result of differences in polymer type, polymer molecular weight characteristics (e.g., molecular weight distribution) or additives content. A second method is differential shrinkage of the fibers after they have been spun into a spunlaid substrate. For instance, heating the spunlaid web can cause fibers to shrink due to differences in crystallinity in the as-spun fibers, for example during the thermal bonding process. A third method of causing crimping is to mechanically stretch the fibers or spunlaid web (generally for mechanical stretching the web has been bonded together). The mechanical stretching can expose differences in the stress-strain curve between the two polymer components, which can cause crimping.

The tensile strength of a fiber is approximately greater than 25Mega Pascal (MPa). The fibers as disclosed herein have a tensile strength of greater than about 50MPa, preferably greater than about 75MPa, and more preferably greater than about 100MPa. Tensile strength is measured using an Instron following a procedure described by ASTM standard D 3822-91 or an equivalent test.
The fibers as disclosed herein are not brittle and have a toughness of greater than 2 MPa, greater than 50 MPa, or greater than 100 MPa. Toughness is defined as the area under the stress-strain curve where the specimen gauge length is 25 mm with a strain rate of 50 mm per minute. Elasticity or extensibility of the fibers may also be desired.

The fibers as disclosed herein can be thermally bondable if enough thermoplastic polymer is present in the fiber or on the outside component of the fiber (i.e. sheath of a bicomponent). Thermally bondable fibers are best used in the pressurized heat and thru-air heat bonding methods. Thermally bondable is typically achieved when the composition is present at a level of greater than about 15%, preferably greater than about 30%, most preferably greater than about 40%, and most preferably greater than about 50% by weight of the fiber.

The fibers disclosed herein can be environmentally degradable depending upon the amount of the composition that is present and the specific configuration of the fiber. "Environmentally degradable" is defined as being biodegradable, disintegratable, dispersible, flushable, or compostable or a combination thereof. The fibers, nonwoven webs, and articles can be environmentally degradable. As a result, the fibers may be easily and safely disposed of either in existing composting facilities or may be flushable and can be safely flushed down the drain without detrimental consequences to existing sewage infrastructure systems. The flushability of the fibers when used in disposable products such as wipes and feminine hygiene items offer additional convenience and discretion to the consumer.

The term "biodegradable" refers to matter that, when exposed to an aerobic and/or anaerobic environment, is eventually reduced to monomelic components due to microbial, hydrolytic, and/or chemical actions. Under aerobic conditions, biodegradation leads to the transformation of the material into end products such as carbon dioxide and water. Under anaerobic conditions, biodegradation leads to the transformation of the materials into carbon dioxide, water, and methane. The biodegradability process is often described as mineralization. Biodegradability means that all organic constituents of the matter (e.g., fibers) are subject to decomposition eventually through biological activity.

There are a variety of different standardized biodegradability methods that have been established over time by various organizations and in different countries. Although the tests vary in the specific testing conditions, assessment methods, and criteria desired, there is reasonable convergence between different protocols so that they are likely to lead to similar conclusions for most materials. For aerobic biodegradability, the American Society for Testing and Materials (ASTM) has established ASTM D 5338-92: Test methods for Determining Aerobic
Biodegradation of Plastic Materials under Controlled Composting Conditions. The ASTM test measures the percent of test material that mineralizes as a function of time by monitoring the amount of carbon dioxide being released as a result of assimilation by microorganisms in the presence of active compost held at a thermophilic temperature of 58°C. Carbon dioxide production testing may be conducted via electrolytic respirometry. Other standard protocols, such as ASTM D 3011-94, may also be used. Standard biodegradation tests in the absence of oxygen are described in various protocols such as ASTM D 5511-94. These tests are used to simulate the biodegradability of materials in an anaerobic solid-waste treatment facility or sanitary landfill. However, these conditions are less relevant for the type of disposable applications that are described for the fibers and nonwovens as described herein.

Disintegration occurs when the fibrous substrate has the ability to rapidly fragment and break down into fractions small enough not to be distinguishable after screening when composted or to cause drainpipe clogging when flushed. A disintegratable material will also be flushable. Most protocols for disintegratability measure the weight loss of test materials over time when exposed to various matrices. Both aerobic and anaerobic disintegration tests are used. Weight loss is determined by the amount of fibrous test material that is no longer collected on an 18 mesh sieve with 1 millimeter openings after the materials is exposed to wastewater and sludge. For disintegration, the difference in the weight of the initial sample and the dried weight of the sample recovered on a screen will determine the rate and extent of disintegration. The testing for biodegradability and disintegration are very similar as a very similar environment, or the same environment, will be used for testing. To determine disintegration, the weight of the material remaining is measured while for biodegradability, the evolved gases are measured. The fibers disclosed herein can rapidly disintegrate.

The fibers as disclosed herein can also be compostable. ASTM has developed test methods and specifications for compostability. The test measures three characteristics: biodegradability, disintegration, and lack of ecotoxicity. Tests to measure biodegradability and disintegration are described above. To meet the biodegradability criteria for compostability, the material must achieve at least about 60% conversion to carbon dioxide within 40 days. For the disintegration criteria, the material must have less than 10% of the test material remain on a 2 millimeter screen in the actual shape and thickness that it would have in the disposed product. To determine the last criteria, lack of ecotoxicity, the biodegradation byproducts must not exhibit a negative impact on seed germination and plant growth. One test for this criteria is detailed in
OECD 208. The International Biodegradable Products Institute will issue a logo for compostability once a product is verified to meet ASTM 6400-99 specifications. The protocol follows Germany's DIN 54900 which determine the maximum thickness of any material that allows complete decomposition within one composting cycle.

The fibers described herein can be used to make disposable nonwoven articles. The articles are commonly flushable. The term "flushable" as used herein refers to materials which are capable of dissolving, dispersing, disintegrating, and/or decomposing in a septic disposal system such as a toilet to provide clearance when flushed down the toilet without clogging the toilet or any other sewage drainage pipe. The fibers and resulting articles may also be aqueous responsive. The term aqueous responsive as used herein means that when placed in water or flushed, an observable and measurable change will result. Typical observations include noting that the article swells, pulls apart, dissolves, or observing a general weakened structure.

The nonwoven products produced from the fibers exhibit certain mechanical properties, particularly, strength, flexibility, softness, and absorbency. Measures of strength include dry and/or wet tensile strength. Flexibility is related to stiffness and can attribute to softness. Softness is generally described as a physiologically perceived attribute which is related to both flexibility and texture. Absorbency relates to the products' ability to take up fluids as well as the capacity to retain them.

Configuration of the Fibers

The fibers disclosed herein can be in many different configurations. The fibers can be multiconstituent. Constituent, as used herein, is defined as meaning the chemical species of matter or the material. Multiconstituent fiber, as used herein, is defined to mean a fiber containing more than one chemical species or material. Generally, fibers may be of monocomponent or multicomponent in configuration. Component, as used herein, is defined as a separate part of the fiber that has a spatial relationship to another part of the fiber. The term multicomponent, as used herein, is defined as a fiber having more than one separate part in spatial relationship to one another. The term multicomponent includes bicomponent, which is defined as a fiber having two separate parts in a spatial relationship to one another. The different components of multicomponent fibers are arranged in substantially distinct regions across the cross-section of the fiber and extend continuously along the length of the fiber.

Spunbond structures, staple fibers, hollow fibers, shaped fibers, such as multi-lobal fibers and multicomponent fibers can all be produced by using the compositions and methods disclosed herein. Multicomponent fibers, commonly a bicomponent fiber, can be in a side-by-side, sheath-
core, segmented pie, ribbon, or islands-in-the-sea configuration. The sheath may be continuous or non-continuous around the core. The ratio of the weight of the sheath to the core is about 5:95 to about 95:5. The fibers disclosed herein can have different geometries that include round, elliptical, star shaped, rectangular, and other various eccentricities.

The fibers disclosed herein can also be splittable fibers. Rheological, thermal, and solidification differential behavior can potentially cause splitting. Splitting may also occur by a mechanical means such as ringrolling, stress or strain, use of an abrasive, or differential stretching, and/or by fluid induced distortion, such as hydrodynamic or aerodynamic.

For a bicomponent fiber, a composition as disclosed herein can be both the sheath and the core with one of the components containing more oil and/or additives than the other component. Alternatively, the composition disclosed herein can be the sheath with the core being some other materials, e.g., pure polymer. The composition can alternatively be the core with the sheath being some other polymer, e.g., pure polymer. The exact configuration of the fiber desired is dependent upon the use of the fiber.

**Processes of Making the Compositions as Disclosed herein**

Melt mixing of the polymer, starch, and oil: The polymer, TPS, and oil and/or wax can be suitably mixed by melting the polymer and TPS in the presence of the oil and/or wax. It should be understood that when the thermoplastic polymer and TPS are melted, the wax will also be in the molten state. In the melt state, the polymer, TPS, and oil and/or wax are subjected to shear which enables a dispersion of the oil into the polymer and/or TPS. In the melt state, the oil and/or wax and polymer and/or TPS are significantly more compatible with each other.

The melt mixing of the thermoplastic polymer, TPS, and oil and/or wax can be accomplished in a number of different processes, but processes with high shear are preferred to generate the preferred morphology of the composition. The processes can involve traditional thermoplastic polymer processing equipment. The general process order involves adding the thermoplastic polymer and TPS to the system, melting the thermoplastic polymer and TPS, and then adding the oil and/or wax. However, the materials can be added in any order, depending on the nature of the specific mixing system.

For the disclosed processes, the thermoplastic starch (TPS) is prepared prior to mixing with a thermoplastic polymer and/or an oil and/or wax. U.S. Patent Nos. 7,851,391, 6,783,854 and 6,818,295 describe processes for producing TPS. However, TPS can be made in-line and the thermoplastic polymer and oil/wax combined in the same production process to make the
compositions as disclosed herein in a single step process. For example, the starch, starch plasticizer and thermoplastic polymer are combined first in a twin-screw extruder where TPS is formed in the presence of the thermoplastic polymer. Later, the oil/wax is introduced into the TPS/thermoplastic polymer mixture via a second feeding location.

**Single Screw Extruder:** A single screw extruder is a typical process unit used in most molten polymer extrusion. The single screw extruder typically includes a single shaft within a barrel, the shaft and barrel engineered with certain screw elements (e.g., shapes and clearances) to adjust the shearing profile. A typical RPM range for single screw extruder is about 10 to about 120. The single screw extruder design is composed of a feed section, compression section and metering section. In the feed section, using fairly high void volume flights, the polymer is heated and supplied into the compression section, where the melting is completed and the fully molten polymer is sheared. In the compression section, the void volume between the flights is reduced. In the metering section, the polymer is subjected to its highest shearing amount using low void volume between flights. For this work, general purpose single screw designs were used. In this unit, a continuous or steady state type of process is achieved where the composition components are introduced at desired locations, and then subjected to temperatures and shear within target zones. The process can be considered to be a steady state process as the physical nature of the interaction at each location in the single screw process is constant as a function of time. This allows for optimization of the mixing process by enabling a zone-by-zone adjustment of the temperature and shears, where the shear can be changed through the screw elements and/or barrel design or screw speed.

The mixed composition exiting the single screw extruder can then be pelletized via extrusion of the melt into a liquid cooling medium, often water, and then the polymer strand can be cut into small pieces. There are two basic types of molten polymer pelletization process used in polymer processing: strand cutting and underwater pelletization. In strand cutting the composition is rapidly quenched (generally much less than 10 seconds) in the liquid medium then cut into small pieces. In the underwater pelletization process, the molten polymer is cut into small pieces then simultaneously or immediately thereafter placed in the presence of a low temperature liquid which rapidly quenches and crystallizes the polymer. These methods are commonly known and used within the polymer processing industry.

The polymer strands that come from the extruder are rapidly placed into a water bath, most often having a temperature range of 1°C to 50°C (e.g., normally is about room temperature, which is 25°C). An alternate end use for the mixed composition is further processing into the
desired structure, for example fiber spinning or injection molding. The single screw extrusion process can provide for a high level of mixing and high quench rate. A single screw extruder also can be used to further process a pelletized composition into fibers and injection molded articles. For example, the fiber single screw extruder can be a 37 mm system with a standard general purpose screw profile and a 30:1 length to diameter ratio.

For example, the fiber single screw extruder is a 37mm system with a standard general purpose screw profile and a 30:1 length to diameter ratio. In the single screw extruder case, already produced TPS and thermoplastic polymer can be combined with the oil/wax, or already produced TPS can be combined with oil/wax that is already dispersed within a thermoplastic polymer. In the first case, an already produced TPS formulation can be melted and the oil/wax additive directly injected into the single screw extruder, followed directly by fiber spinning or final end-use product. The mixing is achieved directly within the single screw extruder. In a second case, the oil/wax is added into the TPS in a second step after the base TPS formulation is produced, similar to the procedure for adding it to a thermoplastic polymer, such as, for example, polypropylene.

**Twin Screw Extruder:** A twin screw extruder is the typical unit used in most molten polymer extrusion, where high intensity mixing is required. The twin screw extruder includes two shafts and an outer barrel. A typical RPM range for twin screw extruder is about 10 to about 1200. The two shafts can be co-rotating or counter rotating and allow for close tolerance, high intensity mixing. In this type of unit, a continuous or steady state type of process is achieved where the composition components are introduced at desired locations along the screws, and subjected to high temperatures and shear within target zones. The process can be considered to be a steady state process as the physical nature of the interaction at each location in the single screw process is constant as a function of time. This allows for optimization of the mixing process by enabling a zone-by-zone adjustment of the temperature and shear, where the shear can be changed through the screw elements and/or barrel design.

The mixed composition at the end of the twin screw extruder can then be pelletized via extrusion of the melt into a liquid cooling medium, often water, and then the polymer strand is cut into small pieces. There are two basic types of molten polymer pelletization process, strand cutting and underwater pelletization, used in polymer processing. In strand cutting the composition is rapidly quenched (generally much less than 10s) in the liquid medium then cut into small pieces. In the underwater pelletization process, the molten polymer is cut into small pieces then simultaneously or immediately thereafter placed in the presence of a low temperature
liquid which rapidly quenches and crystallizes the polymer. An alternate end use for the mixed composition is further processing into the desired structure, for example fiber spinning or injection molding.

Three different screw profiles can be employed using a Baker Perkins CT-25 25mm corotating 40:1 length to diameter ratio system. This specific CT-25 is composed of nine zones where the temperature can be controlled, as well as the die temperature. Four liquid injection sites as also possible, located between zone 1 and 2 (location A), zone 2 and 3 (location B), zone 4 and 5 (location C), and zone 6 and 7 (location D).

The liquid injection location is not directly heated, but indirectly through the adjacent zone temperatures. Locations A, B, C and D can be used to inject the additive. Zone 6 can contain a side feeder for adding additional solids or used for venting. Zone 8 contains a vacuum for removing any residual vapor, as needed. Unless noted otherwise, the melted wax is injected at location A. The wax is melted via a glue tank and supplied to the twin-screw via a heated hose. Both the glue tank and the supply hose are heated to a temperature greater than the melting point of the wax (e.g., about 80°C).

Two types of regions, conveyance and mixing, are used in the CT-25. In the conveyance region, the materials are heated (including through melting which is done in Zone 1 into Zone 2 if needed) and conveyed along the length of the barrel, under low to moderate shear. The mixing section contains special elements that dramatically increase shear and mixing. The length and location of the mixing sections can be changed as needed to increase or decrease shear as needed.

Two primary types of mixing elements are used for shearing and mixing. The first are kneading blocks and the second are thermal mechanical energy elements. The simple mixing screw has 10.6% of the total screw length using mixing elements composed of kneading blocks in a single set followed by a reversing element. The kneading elements are RKB 45/5/12 (right handed forward kneading block with 45° offset and five lobes at 12mm total element length), followed by two RKB 45/5/36 (right handed forward kneading block with 45° offset and five lobes at 36mm total element length), that is followed by two RKB 45/5/12 and reversing element 24/12 LH (left handed reversing element 24mm pitch at 12mm total element length).

The Simple mixing screw mixing elements are located in zone 7. The Intensive screw is composed of additional mixing sections, four in total. The first section is single set of kneading blocks is a single element of RKB45/5/36 (located in zone 2) followed by conveyance elements into zone 3 where the second mixing zone is located. In the second mixing zone, two RKB 45/5/36 elements are directly followed by four TME 22.5/12 (thermomechanical element with
22.5 teeth per revolution and total element length of 12mm) then two conveyance elements into
the third mixing area. The third mixing area, located at the end of zone 4 into zone 5, is
composed of three RKB 45/5/36 and a KB45/5/12 LH (left handed forward reversing block with
45° offset and five lobes at 12mm total element length. The material is conveyed through zone 6
into the final mixing area comprising two TME 22.5/12, seven RKB 45/5/12, followed by SE
24/12 LH. The SE 24/12 LH is a reversing element that enables the last mixing zone to be
completely filled with polymer and additive, where the intensive mixing takes place. The
reversing elements can control the residence time in a given mixing area and are a key
contributor to the level of mixing.

The High Intensity mixing screw is composed of three mixing sections. The first mixing
section is located in zone 3 and is two RKB45/5/36 followed by three TME 22.5/12 and then
conveyance into the second mixing section. Prior to the second mixing section three RSE 16/16
(right handed conveyance element with 16mm pitch and 16mm total element length) elements are
used to increase pumping into the second mixing region. The second mixing region, located in
zone 5, is composed of three RKB 45/5/36 followed by a KB 45/5/12 LH and then a full
reversing element SE 24/12 LH. The combination of the SE 16/16 elements in front of the
mixing zone and two reversing elements greatly increases the shear and mixing. The third
mixing zone is located in zone 7 and is composed of three RKB 45/5/12, followed by two TME
22.5/12 and then three more RKB45/5/12. The third mixing zone is completed with a reversing
element SE 24/12 LH.

An additional screw element type is a reversing element, which can increase the filling
level in that part of the screw and provide better mixing. Twin screw compounding is a mature
field. One skilled in the art can consult books for proper mixing and dispersion. These types of
screw extruders are well understood in the art and a general description can be found in: Twin
Although specific examples are given for mixing, many different combinations are possible using
various element configurations to achieve the needed level of mixing.

For in-line production of TPS, 70wt% solids sorbitol solution can be used to destructure
and plasticize the starch to produce TPS. A side feeder can be installed in Zone 6 to vent off the
majority of the moisture from the starch and liquid sorbitol. The thermoplastic polymer (e.g.,
polypropylene or other thermoplastic polymers as described herein) can then added to the
destructured starch. The oil/wax can be heated and added into the compounding system at
location C or D. In the case where the TPS formulation and the oil/wax are added in the same
process, use of a longer L:D ratio extruder is preferred to increase mixing and enable the various
process steps to be separated. Extruder ratio above 40:1 are contemplated, preferably up to 60:1
and even longer are considered.

Properties of Compositions

The compositions as disclosed herein can have one or more of the following properties
that provide an advantage over known thermoplastic compositions. These benefits can be present
alone or in a combination.

Shear Viscosity Reduction: Viscosity reduction is a process improvement as it can allow
for effectively higher polymer flow rates by having a reduced process pressure (lower shear
viscosity), or can allow for an increase in polymer and/or TPS molecular weight, which improves
the material strength. Without the presence of the oil/wax, it may not be possible to process the
polymer and/or TPS with a high polymer flow rate at existing process conditions in a suitable
way. Alternatively, the presence of the oil/wax can enable lower process temperatures, which
can reduce degradation of the various components (for instance, the TPS component).

Sustainable Content: Inclusion of sustainable materials into the existing polymeric
system is a strongly desired property. Materials that can be replaced every year through natural
growth cycles contribute to overall lower environmental impact and are desired.

Pigmentation: Adding pigments to polymers often involves using expensive inorganic
compounds that are particles within the polymer matrix. These particles are often large and can
interfere in the processing of the composition. Using an oil and/or wax as disclosed herein,
because of the fine dispersion (as measured by droplet size) and uniform distribution throughout
the thermoplastic polymer and/or TPS allows for coloration, such as via traditional ink
compounds. Soy ink is widely used in paper publication) that does not impact processability.

Fragrance: Because the oils and/or waxes, for example SBO or HSBO, can contain
perfumes much more preferentially than the base thermoplastic polymer and/or TPS, the present
composition can be used to contain scents that are beneficial for end-use. Many scented candles
are made using SBO based or paraffin based materials, so incorporation of these into the polymer
for the final composition is useful.

Morphology: The benefits are delivered via the morphology produced in production of
the compositions. The morphology is produced by a combination of intensive mixing and rapid
crystallization. The intensive mixing comes from the compounding process used and rapid
crystallization comes from the cooling process used. High intensity mixing is desired and rapid crystallization is used to preserves the fine pore size and relatively uniform pore size distribution.

Water Resistance: Adding a hydrophobic material to a TPS material improves water resistance of the starch.

Surface Feel: The presence of the oil/wax can change the surface properties of the composition, often making it feel softer.

Improved Spinning Performance: Adding the oil has shown to improve spinning of fibers, enabling a finer diameter filament to be achieved vs the neat polymer the additive has been admixed into during composition preparation.

Processes for Makin2 Fibers

Fibers can be spun from a melt of the compositions as disclosed herein. In melt spinning, there is no mass loss in the extrudate. Melt spinning is differentiated from other spinning, such as wet or dry spinning from solution, where a solvent is being eliminated by volatilizing or diffusing out of the extrudate resulting in a mass loss.

Spinning can occur at 120°C to about 320°C, preferably 185°C to about 250°C and most preferably from 200°C to 230°C. Fiber spinning speeds of greater than 100 meters/minute are preferred. Preferably, the fiber spinning speed is about 1,000 to about 10,000 meters/minute, more preferably about 2,000 to about 7,000 meters/minute, and most preferably about 2,500 to about 5,000 meters/minute. The polymer composition is spun fast to avoid brittleness in the fiber.

Continuous fibers can be produced through spunbond methods or meltblowing processes or non-continuous (staple fibers) fibers can be produced. The various methods of fiber manufacturing can also be combined to produce a combination technique.

The homogeneous blend can be melt spun into multicomponent fibers on conventional melt spinning equipment. The equipment will be chosen based on the desired configuration of the multicomponent. Commercially available melt spinning equipment is available from Hills, Inc. located in Melbourne, Florida. The temperature for spinning is about 100°C to about 320°C. The processing temperature is determined by the chemical nature, molecular weights and concentration of each component. The fibers spun can be collected using conventional godet winding systems or through air drag attenuation devices. If the godet system is used, the fibers can be further oriented through post extrusion drawing at temperatures of about 25°C to about
200°C. The drawn fibers may then be crimped and/or cut to form non-continuous fibers (staple fibers) used in a carding, airlaid, or fluidlaid process.

For example, a suitable process for spinning bicomponent sheath core fibers using the composition in the sheath and a different composition in the core is as follows. A composition is first prepared through compounding containing 10wt% SBO and a second composition is first prepared through compounding containing 30wt% SBO. The 10wt% SBO component extruder profile may be 180°C, 200°C and 220°C in the first three zones of a three heater zone extruder. The transfer lines and melt pump heater temperatures may be 220°C for the first composition. The second composition extruder temperature profile can be 180°C, 230°C and 230°C in the first three zones of a three heater zone extruder. The transfer lines and melt pump can be heated to 230°C. In this case, the spinneret temperature can be 220°C to 230°C.

**Fine Fiber Production**

In one embodiment, the homogenous blend is spun into one or more filaments or fibers by melt film fibrillation. Suitable systems and melt film fibrillation methods are described in U.S. Pat. Nos. 6,315,806, 5,183,670, and 4,536,361, to Torobin et al., and U.S. Pat. Nos. 6,382,526, 6,520,425, and 6,695,992, to Reneker et al. and assigned to the University of Akron. Other melt film fibrillation methods and systems are described in the U.S. Pat. Nos. 7,666,343 and 7,931,457, to Johnson, et al., U.S. Pat. No. 7,628,941, to Krause et al., and U.S. Pat. No. 7,722,347, to Krause, et al. Methods and apparatus described in above patents provide nonwoven webs with uniform and narrow fiber distribution, reduced or minimal fiber defects. Melt film fibrillation process comprises providing one or more melt films of the homogenous blend, one or more pressurized fluid streams (or fiberizing fluid streams) to fibrillate the melt film into ligaments, which are attenuated by the pressurized fluid stream. Optionally, one or more pressurized fluid streams may be provided to aid the attenuation and quenching of the ligaments to form fibers. Fibers produced from the melt film fibrillation process using one of embodiment homogenous blend would have diameters typically ranging from about 100 nanometer (0.1 micrometer) to about 5000 nanometer (5 micrometer). In one embodiment, the fibers produced from the melt film fibrillation process of the homogenous blend would be less than 2 micrometer, more preferably less than 1 micrometer (1000 nanometer), and most preferably in the range of 100 nanometer (0.1 micrometer) to about 900 nanometer (0.9 micrometer). The average diameter (an arithmetic average diameter of at least 100 fiber samples) of fibers of the homogenous blend produced using the melt film fibrillation would be less than 2.5 micrometer, more preferably less than 1 micrometer, and most preferably less than
0.7 micrometer (700 nanometer). The median fiber diameter can be 1 micrometer or less. In an
embodiment, at least 50% of the fibers of the homogenous blend produced by the melt film
fibrillation process may have diameter less than 1 micrometer, more preferably, at least 70% of
the fibers may have diameter less than 1 micrometer, and most preferably, at least 90% of the
fibers may have diameter less than 1 micrometer. In certain embodiments, even 99% or more
fibers may have diameter less than 1 micrometer when produced using the melt film fibrillation
process.

In the melt film fibrillation process, the homogenous blend is typically heated until it
forms a liquid and flows easily. The homogenous blend may be at a temperature of from about
120°C to about 350°C at the time of melt film fibrillation, in one embodiment from about 160°C
to about 350°C, and in another embodiment from about 200°C to about 300°C. The temperature
of the homogenous blend depends on the composition. The heated homogenous blend is at a
pressure from about 15 pounds per square inch absolute (psia) to about 400 psia, in another
embodiment from about 20 psia to about 200 psia, and in yet another embodiment from about 25
psia to about 100 psia.

Non-limiting examples of the pressurized fiberizing fluid stream are gases such as air or
nitrogen or any other fluid compatible (defined as reactive or inert) with homogenous blend
composition. The fiberizing fluid stream can be at a temperature close to the temperature of the
heated homogenous blend. The fiberizing fluid stream temperature may be at a higher
temperature than the heated homogenous blend to help in the flow of the homogenous blend and
the formation of the melt film. In one embodiment, the fiberizing fluid stream temperature is
about 100°C above the heated homogenous blend, in another embodiment about 50°C above the
heated homogenous blend, or just at temperature of the heated homogenous blend. Alternatively,
the fiberizing fluid stream temperature can be below the heated homogenous blend temperature.

In one embodiment, the fiberizing fluid stream temperature is about 50°C below the heated
homogenous blend, in another embodiment about 100°C below the heated homogenous blend, or
200°C below heated homogenous blend. In certain embodiments, the temperature of the
fiberizing fluid stream may be ranging from about -100°C to about 450°C, more preferably,
ranging from about -50°C to 350°C, and most preferably, ranging from about 0°C to about
300°C. The pressure of the fiberizing fluid stream is sufficient to fibrillate the homogenous
blend into fibers, and is above the pressure of the heated homogenous blend. The pressure of the
fiberizing fluid stream may range from about 15 psia to about 500 psia, more preferably from
about 30 psia to about 200 psia, and most preferably from about 40 psia to about 100 psia.
fiberizing fluid stream may have a velocity of more than about 200 meter per second at the location of melt film fibrillation. In one embodiment, at the location of melt film fibrillation, the fiberizing fluid stream velocity will be more than about 300 meter per second, i.e., transonic velocity; in another embodiment more than about 330 meter per second, i.e., sonic velocity; and in yet another embodiment from about 350 to about 900 meters per second (m/s), i.e., supersonic velocity from about Mach 1 to Mach 3. The fiberizing fluid stream may pulsate or may be a steady flow. The homogenous blend throughput will primarily depend upon the specific homogenous blend used, the apparatus design, and the temperature and pressure of the homogenous blend. The homogenous blend throughput will be more than about 1 gram per minute per orifice, for example in a circular nozzle. In one embodiment, the homogenous blend throughput will be more than about 10 gram per minute per orifice and in another embodiment greater than about 20 gram per minute per orifice, and in yet another embodiment greater than about 30 gram per minute per orifice. In an embodiment with the slot nozzle, the homogenous blend throughput will be more than about 0.5 kilogram per hour per meter width of the slot nozzle. In another slot nozzle embodiment, the homogenous blend throughput will be more than about 5 kilogram per hour per meter width of the slot nozzle, and in another slot nozzle embodiment, the homogenous blend throughput will be more than about 20 kilogram per hour per meter width of the slot nozzle, and in yet another slot nozzle embodiment, the homogenous blend throughput will be more than about 40 kilogram per hour per meter width of the slot nozzle. In certain embodiments of the slot nozzle, the homogenous blend throughput may exceed about 60 kilogram per hour per meter width of the slot nozzle. There will likely be several orifices or nozzles operating at one time which further increases the total production throughput. The throughput, along with pressure, temperature, and velocity, are measured at the orifice or nozzle for both circular and slot nozzles.

Optionally, an entraining fluid can be used to induce a pulsating or fluctuating pressure field to help in forming fibers. Non-limiting examples of the entraining fluid are pressurized gas stream such as compressed air, nitrogen, oxygen, or any other fluid compatible (defined as reactive or inert) with the homogenous blend composition. The entertaining fluid with a high velocity can have a velocity near sonic speed (i.e. about 330 m/s) or supersonic speeds (i.e. greater than about 330 m/s). An entraining fluid with a low velocity will typically have a velocity of from about 1 to about 100 m/s and in another embodiment from about 3 to about 50 m/s. It is desirable to have low turbulence in the entraining fluid stream 14 to minimize fiber-to-fiber entanglements, which usually occur due to high turbulence present in the fluid stream. The temperature of the entraining fluid 14 can be the same as the above fiberizing fluid stream, or a
higher temperature to aid quenching of filaments, and ranges from about -40°C to 40°C and in another embodiment from about 0°C to about 25°C. The additional fluid stream may form a "curtain" or "shroud" around the filaments exiting from the nozzle. Any fluid stream may contribute to the fiberization of the homogenous blend and can thus generally be called fiberizing fluid stream.

The spunlaid processes in the present invention are made using a high speed spinning process as disclosed in US Patents Nos 3,802,817; 5,545,371; 6,548,431 and 5,885,909. In these melt spinning processes, extruders supply molten polymer to melt pumps, which deliver specific volumes of molten polymer that transfer through a spinpack, composed of a multiplicity of capillaries formed into fibers, where the fibers are cooled through an air quenching zone and are pneumatically drawn down to reduce their size into highly attenuated fibers to increase fiber strength through molecular level fiber orientation. The drawn fibers are then deposited onto a porous belt, often referred to as a forming belt or forming table.

**Spunlaid Process**

The fibers forming the base substrate in the present invention are preferably continuous filaments forming spunlaid fabrics. Spunlaid fabrics are defined as unbonded fabrics having basically no cohesive tensile properties formed from essentially continuous filaments. Continuous filaments are defined as fibers with high length to diameter ratios, with a ratio of more than 10,000:1. Continuous filaments in the present invention that compose the spunlaid fabric are not staple fibers, short cut fibers or other intentionally made short length fibers. The continuous filaments, defined as essentially continuous, in the present invention are on average, more than 100 mm long, preferably more than 200 mm long. The continuous filaments in the present invention are also not crimped, intentionally or unintentionally. Essentially discontinuous fibers and filaments are defined as having a length less than 100 mm long, preferably less than 50 mm long.

The spunlaid processes in the present invention are made using a high speed spinning process as disclosed in US Patents Nos 3,802,817; 5,545,371; 6,548,431 and 5,885,909. In these melt spinning processes, extruders supply molten polymer to melt pumps, which deliver specific volumes of molten polymer that transfer through a spinpack, composed of a multiplicity of capillaries formed into fibers, where the fibers are cooled through an air quenching zone and are pneumatically drawn down to reduce their size into highly attenuated fibers to increase fiber strength through molecular level fiber orientation. The drawn fibers are then deposited onto a porous belt, often referred to as a forming belt or forming table.
The spunlaid process in the present invention used to make the continuous filaments will
contain 100 to 10,000 capillaries per meter, preferably 200 to 7,000 capillaries per meter, more
preferably 500 to 5,000 capillaries per meter. The polymer mass flow rate per capillary in the
present invention will be greater than 0.3GHM (grams per hole per minute). The preferred range
is from 0.35GHM to 2GHM, preferably between 0.4GHM and 1GHM, still more preferred
between 0.45GHM and 8GHM and the most preferred range from 0.5GHM to 0.6GHM.

The spunlaid process in the present invention contains a single process step for making
the highly attenuated, uncrimped continuous filaments. Extruded filaments are drawn through a
zone of quench air where they are cooled and solidified as they are attenuated. Such spunlaid
processes are disclosed in US 3338992, US 3802817, US 4233014 US 5688468, US 6548431B1,
1340843B1 and EP 1323852B1 can also be used to produce the spunlaid nonwovens. The highly
attenuated continuous filaments are directly drawn down from the exit of the polymer from the
spinneret to the attenuation device, wherein the continuous filament diameter or denier does not
change substantially as the spunlaid fabric is formed on the forming table.

Preferred polymeric materials include, but are not limited to, polypropylene and
polypropylene copolymers, polyethylene and polyethylene copolymers, polyester and polyester
copolymers, polyamide, polyimide, poly(lactic acid), polyhydroxyalkanoate, polyvinyl alcohol,
ethylene vinyl alcohol, polyacrylates, and copolymers thereof and mixtures thereof, as well as the
other mixture presented in the present invention. Other suitable polymeric materials include
thermoplastic starch compositions as described in detail in U.S. publications 2003/0109605A1
and 2003/0091803. Still other suitable polymeric materials include ethylene acrylic acid,
polyolefin carboxylic acid copolymers, and combinations thereof. The polymers described in US
Patents 6746766, US 6818295, US 6946506 and US Published Application 03/0092343.

Common thermoplastic polymer fiber grade materials are preferred, most notably polyester based
resins, polypropylene based resins, poly(lactic acid) based resin, polyhydroxyalkanoate based
resin, and polyethylene based resin and combination thereof. Most preferred are polyester and
polypropylene based resins.

One additional element in the present invention is the ability to utilize mixture
compositions above 40 weigh percent (wt%) wax in the extrusion process, where the masterbatch
level of wax is combined with a lower concentration (down to 0wt%) thermoplastic composition
during extrusion to produce a wax content within the target range.
In the process of spinning fibers, particularly as the temperature is increased above 105°C, typically it is desirable for residual water levels to be 1%, by weight of the fiber, or less, alternately 0.5% or less, or 0.15% or less.

5 Articles

The fibers can be converted to nonwovens by different bonding methods. Continuous fibers can be formed into a web using industry standard spunbond type technologies while staple fibers can be formed into a web using industry standard carding, airlaid, or wetlaid technologies. Typical bonding methods include: calendar (pressure and heat), thru-air heat, mechanical entanglement, hydrodynamic entanglement, needle punching, and chemical bonding and/or resin bonding. The calendar, thru-air heat, and chemical bonding are the preferred bonding methods for the starch polymer fibers. Thermally bondable fibers are required for the pressurized heat and thru-air heat bonding methods.

10 The fibers of the present invention may also be bonded or combined with other synthetic or natural fibers to make nonwoven articles. The synthetic or natural fibers may be blended together in the forming process or used in discrete layers. Suitable synthetic fibers include fibers made from polypropylene, polyethylene, polyester, polyacrylates, and copolymers thereof and mixtures thereof. Natural fibers include cellulotic fibers and derivatives thereof. Suitable cellulotic fibers include those derived from any tree or vegetation, including hardwood fibers, softwood fibers, hemp, and cotton. Also included are fibers made from processed natural cellulotic resources such as rayon.

15 The fibers of the present invention may be used to make nonwovens, among other suitable articles. Nonwoven articles are defined as articles that contain greater than 15% of a plurality of fibers that are continuous or non-continuous and physically and/or chemically attached to one another. The nonwoven may be combined with additional nonwovens or films to produce a layered product used either by itself or as a component in a complex combination of other materials, such as a baby diaper or feminine care pad. Preferred articles are disposable, nonwoven articles. The resultant products may find use in filters for air, oil and water; vacuum cleaner filters; furnace filters; face masks; coffee filters, tea or coffee bags; thermal insulation materials and sound insulation materials; nonwovens for one-time use sanitary products such as diapers, feminine pads, and incontinence articles; biodegradable textile fabrics for improved moisture absorption and softness of wear such as micro fiber or breathable fabrics; an electrostatically charged, structured web for collecting and removing dust; reinforcements and
webs for hard grades of paper, such as wrapping paper, writing paper, newsprint, corrugated paper board, and webs for tissue grades of paper such as toilet paper, paper towel, napkins and facial tissue; medical uses such as surgical drapes, wound dressing, bandages, dermal patches and self-dissolving sutures; and dental uses such as dental floss and toothbrush bristles. The fibrous web may also include odor absorbents, termite repellants, insecticides, rodenticides, and the like, for specific uses. The resultant product absorbs water and oil and may find use in oil or water spill clean-up, or controlled water retention and release for agricultural or horticultural applications. The resultant starch fibers or fiber webs may also be incorporated into other materials such as saw dust, wood pulp, plastics, and concrete, to form composite materials, which can be used as building materials such as walls, support beams, pressed boards, dry walls and backings, and ceiling tiles; other medical uses such as casts, splints, and tongue depressors; and in fireplace logs for decorative and/or burning purpose. Preferred articles of the present invention include disposable nonwovens for hygiene and medical applications. Hygiene applications include such items as wipes; diapers, particularly the top sheet or back sheet; and feminine pads or products, particularly the top sheet.

EXAMPLES

Polymers: U.S. Patent No. 6,783,854 provides a comprehensive list of polymers that are compatible with TPS, although not all have been tested. Current polymeric mixtures have the basic following composition, although it is not limited to the one type described below.

30wt% TPS: Is a mixture of 70wt% polypropylene and 30wt% TPS. The TPS is 70% starch and 30% sorbitol. 10wt% of the polypropylene is maleated PP, Polybond 3200. The remaining PP can be any number of materials, but those used in the present work is 50wt% Basell Profax PH-835 and 50wt% Basell Metocene MF650W.

45wt% TPS: Is a mixture of 70wt% polypropylene and 30wt% TPS. The TPS is 70% starch and 30% sorbitol. 10wt% of the polypropylene is maleated PP, Polybond 3200. The remaining PP can be any number of materials, but those used in the present work is Basell Moplen HP-562T.

Oils/Waxes: Specific examples used were: Soy Bean Oil (SBO); Hydrogenated Soy Bean Oil (HSBO); Partially Hydrogenated Soy Bean Oil (PHSBO); Partially Hydrogenated Palm Kernel Oil (PKPKO); candle with pigmentation and fragrance added; and Standard green Soy Bean Green Ink Pigment.

Compositions were made using a Baker Perkins CT-25 Screw twin screw extruder, with the zones set as noted in the below table:
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<th>Table 1</th>
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<td>30</td>
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<tr>
<td>31</td>
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</tbody>
</table>
For examples 3, 6, and 26, it was noted that the oil was surging at the end of the CT-25 extruder. Examples 3 and 6 failed to properly pelletize. For examples 17-20, 25, and 27, vacuum eliminated blooming at strand outlet of the extruder.

Examples 1-29 demonstrate that one can add oils and waxes to TPS. In Examples 1-29, the TPS resin has been pre-compounded to destructure the starch. Although not required, the oil and wax in Examples 1-29 were added in a second compounding step. What was observed was that with a stable composition (e.g., able to be extruded and/or pelletized), strands from the B&P 25mm system could be extruded, quenched in a water bath at 5°C and cut via a pelletizer without interruption. The twin-screw extrudate was immediately dropped into the water bath.

During stable extrusion, no significant amount of oil/wax separated from the formulation strand (>99wt% made it through the pelletizer). Saturation of the composition can be noted by separation of the polymer and oil/wax from each other at the end of the twin-screw. The saturation point of the oil/wax in the composition can change based on the oil/wax and polymer combination, along with the process conditions. The practical utility is that the oil/wax and polymer remain admixed and do not separate, which is a function of the mixing level and quench rate for proper dispersion of the additive. Specific Examples where the extrusion became unstable from high oil/wax inclusion are Example 3 and 6.

Fibers can be produced by melt spinning a composition of any one of Examples 1-45. Fiber were melt spun with several composition examples.

The specific melt spinning equipment was a specially designed bicomponent extrusion system that consists of two single extruders, followed by a melt pump after each extruder. The two melt streams are combined into a sheath/core spinpack purchased from Hills Inc. The spinpack had 144 holes with capillary orifice diameter of 0.35mm. The fibers extruded through the spinpack were quenched on two sides using a 1 m long quench system that blows air. The fibers are attenuated using a high pressure aspirator that draws the filaments down. The as-spun fibers were deposited onto a belt and collected to measure the final as-spun filament diameter. The as-spun filament diameter is an average of 10 measurements made under a light microscope. The reported fiber diameter is the minimum fiber diameter that could be achieved without any filament breaks over five minutes for the entire 144 filaments being extruded. The mass throughput used was 0.5 grams per capillary per minute (ghm). The specific fibers made and the processes for making them are shown in Table 2.
### Table 2

<table>
<thead>
<tr>
<th>Examples</th>
<th>Sheath Material</th>
<th>Core Material</th>
<th>Sheath Ratio</th>
<th>Core Ratio</th>
<th>Sheath Extruder</th>
<th>Core Extruder</th>
<th>Final Beam</th>
<th>Diameter (micron)</th>
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<td>32</td>
<td>50/50 Blend PH-835 and MF650W</td>
<td>Example 14</td>
<td>30</td>
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<tr>
<td>33</td>
<td>50/50 Blend PH-835 and MF650W</td>
<td>Example 15</td>
<td>30</td>
<td>70</td>
<td>180</td>
<td>190</td>
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<td>190</td>
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<tr>
<td>37</td>
<td>Example 31</td>
<td>Example 15</td>
<td>30</td>
<td>70</td>
<td>180</td>
<td>190</td>
<td>200</td>
<td>200</td>
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</tbody>
</table>
Examples 46-63 show the results from producing useful fibers and the benefit of improved spinnability by adding oil. The examples show that utilizing polypropylene with oil in the core or into the sheath and core improves the spinnability and enable finer filaments to be produced. Finer fibers can improve softness, barrier properties and wicking behavior.

Spunbond nonwovens were made by using the porous collection belt and adjusting the belt speed to target 20 grams per square meter (gsm). The collected fibers were first passed through a heated press roll at 100°C at 50 PLI (pounds per linear inch) and then a heated calendering system for the final thermal point bonding, followed by winding the continuous spunbond nonwoven onto a roll for later property measurements. The heated calendering system consisted of a heated engraved roll and heated smooth roll. The heated engraved roll had 18% raised bonding area. The calender roll pressure was held constant at 350PLI and the line speed of the forming belt was held constant at 38 meters per minute.

The tensile properties of base substrates and structured substrates were all measured the same way. The gauge width is 50 mm, gauge length is 100 mm in the MD and 50mm in the CD and the extension rate is 100 mm/min. The values reported are for strength and elongation at peak, unless stated otherwise. Separate measurements are made for the MD and CD properties. The typical units are Newton (N), and they are Newtons per centimeter (N/cm). The values presented are the average of at least ten measurements. The perforce load is 0.2 N. The samples should be stored at 23 ± 2°C and at 50 ± 2% relative humidity for 24 hours with no compression, then tested at 23 ± 2°C and at 50 ± 2%. The tensile strength as reported here is the peak tensile strength in the stress-strain curve. The elongation at tensile peak is the percent elongation at which the tensile peak is recorded.

Examples 64-103 show that useful spunbond nonwovens can be produced. The specifics of Examples 64-103 are shown in Table 3. The examples show that an optimum bonding temperature is achieved at a particular fiber composition.

### Table 3

<table>
<thead>
<tr>
<th>Example</th>
<th>Sheath Material</th>
<th>Core Material</th>
<th>Fiber Example</th>
<th>Engraved Roll</th>
<th>Smooth Roll</th>
<th>Temperature (°C)</th>
<th>Average Basis Wt (gsm)</th>
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All documents cited in the Detailed Description of the Invention are, in relevant part, incorporated herein by reference; the citation of any document is not to be construed as an
admission that it is prior art with respect to the present invention. To the extent that any meaning or definition of a term in this document conflicts with any meaning or definition of the same term in a document incorporated by reference, the meaning or definition assigned to that term in this document shall govern.

The dimensions and values disclosed herein are not to be understood as being strictly limited to the exact numerical values recited. Instead, unless otherwise specified, each such dimension is intended to mean both the recited value and a functionally equivalent range surrounding that value. For example, a dimension disclosed as "40 mm" is intended to mean "about 40 mm".

While particular embodiments of the present invention have been illustrated and described, it would be obvious to those skilled in the art that various other changes and modifications can be made without departing from the spirit and scope of the invention. It is therefore intended to cover in the appended claims all such changes and modifications that are within the scope of this invention.
CLAIMS

What is claimed is:

1. A fiber produced by melt spinning a composition comprising an intimate admixture of
   (a) a thermoplastic starch;
   (b) a thermoplastic polymer; and
   (c) an oil, wax, or combination thereof present in an amount of 5 wt% to 40 wt%, based upon the total weight of the composition.

2. The fiber of claim 1, wherein the thermoplastic polymer comprises a polyolefin, a polyester, a polyamide, copolymers thereof, or combinations thereof.

3. The fiber of claim 2, wherein the thermoplastic polymer is selected from the group consisting of polypropylene, polyethylene, polypropylene co-polymer, polyethylene co-polymer, polyethylene terephthalate, polybutylene terephthalate, polylactic acid, polyhydroxyalkanoates, polyamide-6, polyamide-6,6, and combinations thereof.

4. The fiber of any one of claims 1 to 3, wherein the thermoplastic polymer comprises polypropylene.

5. The fiber of any one of claims 1 to 4, wherein the thermoplastic polymer comprises 20 wt% to 90 wt% of the composition, based upon the total weight of the composition.

6. The fiber of any one of claims 1 to 5, wherein the oil, wax or combination thereof comprises a lipid.

7. The fiber of claim 6, wherein the lipid comprises a monoglyceride, diglyceride, triglyceride, fatty acid, fatty alcohol, esterified fatty acid, epoxidized lipid, maleated lipid, hydrogenated lipid, alkyd resin derived from a lipid, sucrose polyester, or combinations thereof.

8. The fiber of any one of claims 1 to 7, wherein the oil, wax, or combination thereof is selected from the group consisting of soy bean oil, epoxidized soy bean oil, maleated soy bean oil, corn oil, cottonseed oil, canola oil, beef tallow, castor oil, coconut oil, coconut seed oil, corn germ oil, fish oil, linseed oil, olive oil, oiticica oil, palm kernel oil, palm oil, palm seed oil, peanut oil, rapeseed oil, safflower oil, sperm oil, sunflower seed oil, tall oil, tung oil, whale oil, tristearin, triolein, tripalmitin, 1,2-dipalmitoolein, 1,3-
dipalmitoolein, 1-palmito-3-stearo-2-olein, 1-palmito-2- stearo-3-olein, 2-palmito-1-stearo-3-olein, trilinolein, 1,2-dipalmitolinolein, 1-palmito-dilinolein, 1-stearo- dilinolein, 1,2-diacetopalmitin, 1,2-distearo-olein, 1,3-distearo-olein, trimyristin, trilaurin, capric acid, caproic acid, caprylic acid, lauric acid, lauroleic acid, linoleic acid, linolenic acid, myristic acid, myristoleic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, and combinations thereof.

9. The fiber of any one of claims 1 to 8, wherein the oil, wax, or combination thereof is dispersed within the thermoplastic polymer such that the oil, wax, or combination thereof has a droplet size of less than 10 µm within the thermoplastic polymer.

10. The fiber of claim 9, wherein the droplet size is less than 1 µm.
What is claimed is:

1. A fiber produced by melt spinning a composition comprising an intimate admixture of
   (a) a thermoplastic starch;
   (b) a thermoplastic polymer; and
   (c) an oil, wax, or combination thereof present in an amount of 5 wt% to 40 wt%, based upon the total weight of the composition, said oil, wax, or combination thereof being dispersed within said thermoplastic polymer such that the oil, wax, or combination thereof has a droplet size of less than 10 μm within the thermoplastic polymer.

2. The fiber of claim 1, wherein the thermoplastic polymer comprises a polyolefin, a polyester, a polyamide, copolymers thereof, or combinations thereof.

3. The fiber of claim 2, wherein the thermoplastic polymer is selected from the group consisting of polypropylene, polyethylene, polypropylene co-polymer, polyethylene co-polymer, polyethylene terephthalate, polybutylene terephthalate, polylactic acid, polyhydroxyalkanoates, polyamide-6, polyamide-6,6, and combinations thereof.

4. The fiber of any one of claims 1 to 3, wherein the thermoplastic polymer comprises polypropylene.

5. The fiber of any one of claims 1 to 4, wherein the thermoplastic polymer comprises 20 wt% to 90 wt% of the composition, based upon the total weight of the composition.

6. The fiber of any one of claims 1 to 5, wherein the oil, wax or combination thereof comprises a lipid.

7. The fiber of claim 6, wherein the lipid comprises a monoglyceride, diglyceride, triglyceride, fatty acid, fatty alcohol, esterified fatty acid, epoxidized lipid, maleated lipid, hydrogenated lipid, alkyd resin derived from a lipid, sucrose polyester, or combinations thereof.

8. The fiber of any one of claims 1 to 7, wherein the oil, wax, or combination thereof is selected from the group consisting of soy bean oil, epoxidized soy bean oil, maleated soy bean oil, corn oil, cottonseed oil, canola oil, beef tallow, castor oil, coconut oil,
coconut seed oil, corn germ oil, fish oil, linseed oil, olive oil, oiticica oil, palm kernel oil, palm oil, palm seed oil, peanut oil, rapeseed oil, safflower oil, sperm oil, sunflower seed oil, tall oil, tung oil, whale oil, tristearin, tripalmitin, 1,2-dipalmitoolein, 1,3-dipalmitoolein, 1-palmito-3-stearo-2-olein, 1-palmito-2-stearo-3-olein, 2-palmito-1-stearo-3-olein, trilinolein, 1,2-dipalmitolinolein, 1-palmito-dilinolein, 1-stearo-dilinolein, 1,2-diaceptopalmitin, 1,2-distearo-olein, 1,3-distearo-olein, trimyristin, trilaurin, capric acid, caproic acid, caprylic acid, lauric acid, lauroleic acid, linoleic acid, linolenic acid, myristic acid, myristoleic acid, oleic acid, palmitic acid, palmitoleic acid, stearic acid, and combinations thereof.

9. The fiber of any one of claims 1 to 8, wherein the droplet size of said oil, wax, or combination thereof is less than 1 µm.
**INTERNATIONAL SEARCH REPORT**

**International application No**

PCT/US2012/038303

**A. CLASSIFICATION OF SUBJECT MATTER**

INV. D01F6/46 D01F6/90 D01F6/92 D01F9/00

According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

D01F

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**

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<td>FR 2 92Q 432 AI (CONTI CELINE [FR] ; PONTANIER AURELIA [FR]) 6 March 2009 (2Q09-Q3-O6) page 2, line 11 - page 3, line 34 claims 1-14</td>
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Further documents are listed in the continuation of Box C. X See patent family annex.

* Special categories of cited documents:
- "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
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- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed
- "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
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- "Y" document of particular relevance; the claimed invention cannot be 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
- "Z" document member of the same patent family

Date of the actual completion of the international search 27 July 2012

Date of mailing of the international search report 07/08/2012

Name and mailing address of the ISA/

European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk

Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016

Authorized officer Verschuren , Jo
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