GLYCOLIC ACID POLYMERS AND METHOD OF PRODUCING THE SAME

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ABSTRACT
Described herein is a feasible, significantly simplified production method that avoids challenging lactonization steps and converts a low molecular weight aliphatic polyester, consisting of hydroxy acids and a comonomer, whose molecular weight has been increased by step-growth polymerization reactions. The molecular weight of the aliphatic polyester, based on comparison of initial and final weight average molecular weights ($M_w, M_n$), increased significantly at a rate which permits the use of reactive extrusion to produce high molecular weight aliphatic polyesters in a simple, economically feasible manner.
Fig. 1
Fig. 2

Fig. 3
Case 1.

Case 2.

Case 3.

Fig. 4
GLYCOLIC ACID POLYMERS AND METHOD OF PRODUCING THE SAME

CROSS-REFERENCE TO RELATED APPLICATIONS


TECHNICAL FIELD

[0002] The present invention relates to polyesters. In particular, the present invention concerns the synthesis difunctional aliphatic polyesters, which may exhibit a low molecular weight limiting their use in practise, conversion of such polymers to a higher molecular weight polymeric material and production methods thereof leading to products derived mainly from hydroxy acids, particularly but not exclusively, from glycolic acid.

BACKGROUND ART


[0004] A common disadvantage of conventional condensation polymerization is that a polymer exhibiting a low molar mass is typically formed in the process. The low molecular weight prevents the polymer to achieve sufficiently good properties to be useful in a myriad of applications. Similar methods to produce polyglycolic acid are polycondensation of an alkyl glycolate and desalting polycondensation of a glycolic acid salt. Equivalent reactions can be used for lactic acid, albeit with a frequent disadvantage where the desired optical purity of the precursor is lost due to racemization during the polycondensation reactions.

[0005] As known by those familiar with the art, higher molecular weight materials with subsequently improved properties can be obtained by ring-opening polymerization of the equivalent lactone or cyclic ester. Improvements to these processes have similarly a long history (GB 825,335, U.S. Pat. No. 3,442,871). However, the preparation of pure glycolide which yields high molecular weight polyglycolic acid is difficult to achieve. For example, preparation of pure glycolide has been described in 1987 (cf. U.S. Pat. No. 4,650,851). Numerous process improvements have been reported (cf. U.S. Pat. No. 5,223,630; U.S. Pat. No. 7,235,673). Processes to prepare lactide, a closely related molecule, result in poor collected yields of glycolide and high amounts of undesired side products which make the glycolide forming process a bottleneck for wider use of the material. Thus, preparation of glycolide in high yields requires the use of solvents and additives which must be separated from the product and present a technical obstacle for wider use of the materials.

[0006] Aliphatic polyesters, including polyglycolic acid and polyactic acid, are biodegradable materials as these are degraded through hydrolysis and/or through microbial or enzymatic pathways. The biodegradability and biocompatibility make these useful in many applications. Poly(α-hydroxy acid)s have been under research for medical devices as surgical sutures and artificial skins since the 1960s, where one area of interest has been on polyglycolic acid (Vert, M. et al., Makromol Chem Suppl 1981, 5, 30-41). Beyond medical applications polyglycolic acid has been proposed as a barrier material for containers (U.S. Pat. No. 4,424,242, U.S. Pat. No. 4,565,851). Later in 1988 polyglycolic acid copolymers have been applied also for packaging applications (cf. U.S. Pat. No. 4,729,927).

[0007] Utilization of step growth copolymerization with for example diisocyanates is a well-known technology. Such methods have been used for lactic acid based materials (U.S. Pat. No. 5,380,813) with a clear disadvantage as the optically pure precursor undergoes racemization during the condensation process losing its crystallinity, thus severely limiting its applicability due to a low glass transition temperature as the sole thermal transition for the material.

SUMMARY

[0008] It is an object of this invention to provide a method to produce a high molecular weight copolymer consisting of polyglycolic acid segments that exhibits properties making it useful in applications including but not limiting to packaging of various articles or fluids, as a barrier material or in medical applications as implantable material or drug delivery vehicle.

[0009] A second object of this invention is to provide material compositions consisting of polyglycolic acid segments that are able to form crystalline structures after a step-growth molecular weight extension step.

[0010] It is a third objective of this invention to provide methods to prepare a suitable α,ω-difunctional polyglycolic acid polymer prepared by but not limited to a condensation process which can be efficiently utilized in step-growth polymerizations to increase its molecular weight.

[0011] A further objective is to provide suitable comonomers for the step-growth polymerization or chain extension of the described polyglycolic acid polymer.

[0012] One more object of this invention is to carry out the step-growth copolymerization for molecular weight increase in such a manner that the weight percentage of the polyglycolic acid segments is as high as possible in order to retain its useful properties.

[0013] Still a further object is to avoid glycolide as an intermediate for the polymer through preparation of telechelic hydroxy-terminated prepolymers of glycolic acid by a condensation process, and which can be applied by useful methods and process to polymerize hydroxyacid monomers to high molar weight polymers which are processable to applications.

[0014] Final object is to use said materials as films, sheets, fibers, powders or molded articles in applications.

[0015] These and other objects, which jointly with existing materials and methods are achieved in the present description are described and claimed herein.

[0016] The present invention is based on the formation of a polyglycolic acid polymer, or optionally copolymer, which can undergo step-growth reactions to increase its molecular weight. By a subsequent step-growth polymerization step of a polyglycolic acid polymer, or optionally copolymer, high molecular weights can be achieved which are difficult to achieve for polyglycolic acid, particularly by condensation processes as known for those familiar to the art. In particular, significant benefits are achieved when said polyglycolic acid polymer has been prepared by a condensation process thus avoiding the cumbersome synthesis of glycolide and its subsequent ring-opening polymerization.
[0017] Particularly preferred compositions of the polyglycolic acid polymer and copolymer include, in addition to glycolic acid:

[0018] one or more organic molecule, linear, branched, cyclic, aromatic or polymeric containing two or more hydroxy, carboxylic acid, thiol or amino groups

[0019] a hydroxy acid comonomer, or its equivalent lactone, of general formula

\[ \text{HO} (\text{CH}_2)_m \text{CHR} (\text{CH}_2)_n \text{COOH} \]

[0020] wherein

[0021] \( R \) is independently selected from a group consisting of hydrogen, linear alkyl, alkynyl alkynyl, branched alkyl, cyclic alkyl and aryl moieties, and

[0022] \( n \) and \( m \) are integers which can independently vary between 0 and 20

[0023] Preferred classes of the chemicals that can be used to increase the molecular weight of the polyglycolic acid polymer and copolymer include: disiocyanate, bisepoxy compounds, dialdehydes, diamines, diketenes, phosphoric acid esters and bisoxazolines. Particularly preferred classes are disiocyanates and bisepoxy compounds.

[0024] More specifically, the present (pre)polymers comprise the steps of providing glycolic acid; providing a bifunctional monomer; subjecting said glycolic acid to condensation polymerization in the presence of said monomer; and continuing polymerization to provide a polymeric chain formed by residues derived from glycolic acid and said bifunctional monomer.

[0025] The method according to the present invention for producing (pre)polymers comprises the steps of providing glycolic acid; providing a bifunctional monomer; subjecting said glycolic acid to polymerization in the presence of a catalyst and said bifunctional monomer; and continuing polymerization to provide a polymeric chain formed by residues derived from glycolic acid and said monomer, and the glycolic acid polymers according to the present invention comprising at least two prepolymers, preferably 5 to 100 prepolymers, linked together.

[0026] The method of producing the polymers comprises subjecting a prepolymer to step-growth polymerization in the presence of a chain extender to provide a linear polymer having a molecular weight (Mn) of more than 10,000 g/mol, in particular about 20,000 to 1,000,000 g/mol and the novel uses for forming polymeric films, sheets, fibers, powders or moulded articles, in particular for forming compression-moulded, injection-moulded, blow-moulded products, and other formed or moulded products.

[0027] Considerable advantages are obtained by the present invention. The present invention provides for controlled preparation of polyglycolic acid polymers by condensation polymerization in a first step, optionally in combination with an increase in molecular weight in a second step, which preferably is being carried out after the first step. The increase in molecular weight can be achieved using chemicals that extend the polymer chains to longer units yielding high molecular weight linear or optionally branched polymers.

[0028] These polymers retain melting transitions and hence also find use in higher temperature applications. Thus, various embodiments of the invention are useful for producing polyglycolic acid based materials with high molecular weights.

[0029] Lactic acid is optically active, and condensation polymerization typically leads to racemization of the optically pure monomers. As a result, corresponding polymers (PLAs) are not crystalline and their use is limited. The present glycolic acid polymers are not impaired by such features.

[0030] The polymers obtained can be used as such or in blends with other polymers in a range of applications.

[0031] Next, the invention will be examined more closely with the aid of detailed description with reference to the appended drawings.

**BRIEF DESCRIPTION OF DRAWINGS**

[0032] FIG. 1 shows the molecular weight increase of polymers with varying degree of hydroxyl-termination based on Examples 8, 9, 10, 11 and 12;

[0033] FIG. 2 shows the effect of the amount of chain extender used as in Example 12;

[0034] FIG. 3 shows the molecular weight increase of polymers when a bisepoxy functional compound is used as in Example 13;

[0035] FIG. 4 shows three plausible polymer structures for a polymer prepared by condensation polymerization of glycolic acid and a bifunctional comonomer (e.g. hexanediol): Case 1—OH/COON-terminated polymer chain, Case 2—α, ω-OH-terminated polymer chain, and Case 3—macroyclic polymer chain with no terminating group(s);

[0036] FIG. 5 is a quantitative 13C NMR spectrum of a polyhydroxy acid polymer from range of 174-165 ppm;

[0037] FIG. 6 is a quantitative 13C NMR spectrum of a polyhydroxy acid polymer from range of 67-59 ppm and 25-24 ppm; and

[0038] FIG. 7 is an NMR showing completion of the step growth polymerization reaction in 2 minutes.

**DETAILED DESCRIPTION**

[0039] Described herein is a feasible, significantly simplified production method that avoids challenging lactonization steps and converts a low molecular weight aliphatic polyester, consisting of hydroxy acids and a comonomer, whose molecular weight has been increased by step-growth polymerization reactions, providing glycolic acid;

[0040] The method comprises providing a bifunctional monomer, subjecting said glycolic acid to condensation polymerization in the presence of the monomer and preferably a catalyst; and continuing polymerization to provide a polymeric chain formed by residues derived from glycolic acid and said bifunctional monomer. Preferably, in particular when carrying our condensation polymerization in the presence of an esterification catalyst, water formed during condensation polymerization is continuously removed.

[0041] The molecular weight of the aliphatic polyester, based on comparison of initial and final weight average molecular weights (\( M_{n,1}, M_{n,2} \)), increased significantly at a rate which permits the use of reactive extrusion to produce high molecular weight aliphatic polyesters in a simple, economically feasible manner.

[0042] In one embodiment, the present technology provides for telechelic polymers of glycolic acid. The term “telechelic” is used for indicating that the present polymers or prepolymers are capable of being subjected to polymerization through their reactive end-groups. The end groups typically exhibit the same (chemical) functionality.

[0043] In a preferred embodiment, the polymers of the present kind are capable of being used as prepolymers.
Typically the present polymers comprise at least 5, for example 5 to 250, preferably 6 to 100, in particular 10 to 50 residues of glycolic acid.

The polymers comprise in one preferred embodiments essentially linear polymer chains. In another preferred embodiment, the polymers comprise a branched polymeric chain.

As will be discussed in more detail below, in a preferred embodiment, the present polymers have a degree of crystallinity of at least 50%, preferably at least 40%, in particular at least 50% of the crystallinity or melting enthalpy of the prepolymer.

Further, the present polymers comprise typically at least 80 mol-%, preferably at least 90 mol-%, of residues derived from glycolic acid.

To reach the aim of providing a telechelic polymer, the polymer according to the present technology suitably comprises 0.1 to 20 mol-%, preferably 0.5 to 10 mol-%, more preferably 1 to 5 mol-%, of residues derived from a suitably terminated, e.g. a difunctional, comonomer, typically a hydroxy-terminated comonomer.

In a particularly interesting embodiment, the molar ratio between residues derived from glycolic acid and residues derived from a comonomer is 1000:1 or less, for example 500:1 or less, suitably 50:1 or less, advantageously 45:1 to 55:1, in particular 48:1 to 52:1.

In one embodiment, the comonomers are selected exclusively from comonomers which are hydroxy-terminated comonomers (diols, triols, tetraols, pentaols), such as propanediol, butanediol, hexanediol, pentaerythritol and oligomeric polyethylene glycol and combinations thereof, to produce a hydroxy-terminated telechelic polymer.

In another embodiment, at least a part of the comonomers (optionally in combination with any of the above, in particular diols) is selected from the group of dicarboxylic acid, dithiol, and diamine and mixtures thereof.

Formula 1 shows an example of a telechelic, hydroxyl-terminated glycolic acid polymer of the instant kind. In the formula m and n are integers which stand for a value of 1 to 100.

Various features of the synthesis of the polyhydroxy acid polymers according to the present technology will be examined in the following.

It has been found that polyhydroxy acid polymers, in particular polyesters of a kind consisting to a high degree of polyglycolic acid, are suitable for a subsequent step growth polymerization process which extends the molecular weight of the resultant material to levels which are required for its acceptable use in applications.

The polyhydroxy acid polymer can be obtained as a product of condensation polymerization of glycolic acid, optionally one or more hydroxy acid, or the corresponding lactone, as a comonomer, and one or more suitable acid difunctional compound.

In one embodiment, the step of preparing a hydroxyl-terminated telechelic polymer of glycolic acid, comprises the steps of providing glycolic acid; providing a hydroxy-terminated monomer; subjecting the glycolic acid to condensation polymerization in the presence of an esterification catalyst and the hydroxy-terminated monomer; continuously removing water formed during condensation polymerization; and continuing polymerization to provide a polymeric chain formed by residues derived from glycolic acid and the hydroxy-terminated monomer.

The esterification catalyst can be an organic or inorganic compound. The catalyst can be an organic or inorganic acid. The catalyst can also be a metal compound of tin, zinc, lead, titanium, antimony, cerium, germanium, cobalt, magnesium, iron, aluminium, magnesium, calcium and strontium. For example, metal alkoxides, organic acid salts of metal, chelates and metal oxides can be used. Particularly useful catalysts are organic zinc, tin and titanium compounds, such as zinc, tin or titanium octoate, and alkyl ester titanate, titanium oxy acetyl acetonate, and titanium oxalate.

The amount of the catalyst is preferably from 0.001 to 0.5% by weight of the glycolic acid together with the comonomers.

According to one preferred embodiment, the content of the glycolic acid in the polyhydroxy acid polymer is so high that the condensed segments of the glycolic acid repeating units are able to form crystals in the polyhydroxy acid polymer and in the subsequent polymer formed after a step growth polymerization process. Hence, it is preferred that the glycolic acid weight ratio to the total weight of monomers is, at the start of the polymerization, at least more than 50%, preferably more than 70% and most preferably more than 90%.

Another hydroxy acid, or the corresponding lactone thereof, may be used in a preferred composition as a comonomer to adjust the properties of the polyhydroxy acid polymer. Such hydroxy acids have the general formula

\[ \text{HOC(CH}_2\text{CHR(CH}_2\text{)}_n\text{COOH} \]

wherein R is independently selected from a group consisting of hydrogen, linear alkyl, linear alkenyl linear alkyl, branched alkyl, cyclic alkyl, cyclic alkenyl, aromatic (consisting of 1-6 rings) and alkylaromatic (consisting of 1-6 rings) moieties; and

n and m are integers which can vary between 0 and 20.

Examples of hydroxy acid comonomers include lactic acid, 3-hydroxypropionic acid, 2-hydroxybutanoic acid, 3-hydroxybutanoic acid, 4-hydroxybutanoic acid, 5-hydroxypentanoic acid, 6-hydroxyhexanoic acid, and benzoic acid.

According to another preferred embodiment of the invention the weight ratio of added hydroxy acid comonomers is less than 30%, more preferably less than 20% and most preferably less than 10%.

Suitable acid, \( \alpha, \omega \)-difunctional compounds are used to generate polyhydroxy acid compositions which in turn yield and a suitable acid, \( \alpha, \omega \)-difunctional polyhydroxy acid materials. The ability to form \( \alpha, \omega \)-difunctional polyhydroxy acid materials is critical to achieve successful increase in molecular weight in the subsequent step growth polymerization step. In the examples below it is demonstrated that having a high
degree of α,ω-difunctionality in the polyhydroxy acid materials has a direct impact on the success of the step growth polymerization step.

[0071] The properties and molecular weight of α,ω-difunctional polyhydroxy acid material can be adjusted based on the ratio of the hydroxy acid and the α,ω-difunctional compound. The amount of the α,ω-difunctional compound will directly impact on the resultant molecular weight of the material. An increased use of α,ω-difunctional compounds results in a decreased molecular weight of the resultant α,ω-difunctional polyhydroxy acid material. Also, the properties of α,ω-difunctional compound will have an impact on the resultant α,ω-difunctional polyhydroxy acid material. For instance, the use of α,ω-terminal polymeric material in the condensation process will yield a copolymer with unique properties.

According to a further preferred embodiment of the invention, the content of the α,ω-difunctional compounds is less than 20%, preferably less than 15% and most preferably less than 10%.

[0072] As noted above, such an organic added α,ω-difunctional compound may be linear, branched, cyclic, aromatic or polymeric containing hydroxy, carboxylic acid, thio or aminogroups. Examples of such compounds are diols, dicarboxylic acids and their anhydrides, diamines and polymeric materials having α,ω-difunctionality, wherein the functionality is a dihydroxy, diacid, diithio or diamines.

[0073] Examples of such classes of compounds are ethylene glycol, 1,3-propanediol, 1,4-butanediol, 1,5-pentanediol, 1,6-hexanediol, 1,7-heptanediol, 1,8-octanediol, 1,9-nonanediol, 1,10-decanediol, 1,2-propanediol, 1,2-butanediol, 2,3-butanediol, 1,3-butanol, 1,2-pentanediol, neopentyl glycol, equivalent thiols, oxalic acid, malonic acid, maleic acid, maleic anhydride itaconic acid, succinic acid, succinic anhydride, glutaric acid, adipic acid, pimelic acid, suberic acid, azelaic acid, sebacic acid, ethylene diamine, 1,3-propylene diamine, 1,4-butanedi amine, 1,5-pentanedi amine, 1,6-hexanedi amine, telechelic polyethylene glycol, telechelic polypropylene glycol, polytetramethyl ethylene. These examples are not exclusive and can be complemented with related compounds, linear or branched as well as cyclic or aromatic and derivatives thereof.

[0074] In a further preferred embodiment, the present invention also permits the preparation of α,ω-multiterminated polyhydroxy acids. Such materials are obtained when the α,ω-difunctional compounds is replaced by an equivalent having three or more, or dissimilar functionalities, thus permitting the preparation of α,ω-multiterminated polyhydroxy acids which are capable of step growth conditions yield thermo-setting materials as known for those familiar to the art. Particularly preferred are compounds having three or more similar functionalities. Unambiguous examples of such compounds are glycerol, pentaerythritol, trimethylol propane and the like.

[0075] One further embodiment of this invention is the production process how to obtain α,ω-difunctional polyhydroxy acid materials. Polyglycolic acid as a homopolymer has a melting transition above 210-220°C, which necessitates high reaction temperatures if one wants to conduct the polymerization process in a molten state.

[0076] Basically, this may require that the reaction media be heated to 230-250°C, which under prolonged times lead to darkening of the material, in particular if oxygen is present. A discoloured product is of disadvantage in consumer applications when optically attractive materials are preferred and hence yellowish or even brown-like materials are unfavourable.

[0077] It has surprisingly been found that the melt condensation polymerization of glycolic acid can be performed at temperatures below the melting point of the product polyglycolic acid (which is typically 210-220°C, as mentioned above). The polycondensation temperature is gradually increased and the absolute pressure decreased during the typical melt polycondensation stage.

[0078] It is beneficial to avoid too high polymerisation temperatures that easily generate unwanted brown colour for the polymer.

[0079] Thus, according to a preferred embodiment of the present technology, α,ω-difunctional polyhydroxy acid materials can be obtained by a process where the condensation polymerization is carried out sequentially first in the molten state and then followed by a condensation process that takes place in a solid state at 200°C or less for a prolonged period of time. The use of lower polymerization temperatures in the solid-state polymerization does not substantially affect the colour of the material. Prolonged polymerization times at high temperatures will cause undesired yellowing or darkening of the product. The solid state polymerization may optionally be followed by a final step at 240°C or higher for a shortest possible time to obtain a freely flowing product that can be easily handled and transferred. Reduced pressure is beneficial for the progress of the condensation polymerisation during all the steps, and in the later stages of polymerisation high vacuum can be applied (below 50 mbar absolute pressure). A preferred time for the first melt polymerization is 36 h or less and 24 h or less for the solid polymerization prior to the final dehydration, and condensation process. A further method to increase the content of the α,ω-difunctional polyhydroxy acid materials is to increase the content of the α,ω-difunctional comonomer in the feed, which increases the probability of obtaining an α,ω-difunctional polyhydroxy acid material.

[0080] In a particularly preferred embodiment, the polymerisation temperatures at the end of the melt stage polycondensation are 180-220°C, preferably 190-210°C. Typically, operation at these temperature ranges is possible after 6 to 10 h from the beginning of the polycondensation stage.

[0081] The instant embodiment also makes it possible to produce the telechelic prepolymer with high enough molecular weight and uniform enough and group functionalization without solid state polymerisation stage.

[0082] Another method to produce the α,ω-difunctional polyhydroxy acid materials, as known for those familiar to the art, is the process in a solvent where the said solvent is capable of dissolving the polymer that is formed and simultaneously assist in removal of the condensation product byzeotropic distillation or the like. Examples of such solvents are dimethylformamide, dimethylacetamide, diphenyl ether and dimethylsulphoxide.

[0083] In the above discussed embodiments, the monomers and the difunctional compound can be added simultaneously or sequentially. In one embodiment, glycolic acid monomers are first subjected to condensation polymerization, optionally together with comonomers, to provide oligomers (comprising 2 to 8 units) and only then difunctional compound is added. In another embodiment, glycolic acid is first condensation polymerized, the comonomer is added, and polymerization is continued. Finally difunctional compound is added
and polymerization is continued. In a third embodiment, glycolic acid monomers are first condensation polymerized, then difunctional compound is added and polymerization is continued. Finally, comonomer is added and polymerization is continued.

[0084] In one more embodiment the temperature for the melt polymerization can be adjusted by the amount of \(\alpha,\omega\)-difunctional comonomer. Typically, increased concentrations of the \(\alpha,\omega\)-difunctional comonomer result in a decreased melting temperature for the \(\alpha,\omega\)-difunctional polyhydroxy acid material, which permit the use of decreased polymerization times and hence may impart in the desired appearance of the product.

Molecular Weight Increase of the Polyhydroxy Acid by Step Growth Polymerization

[0085] In studies it has been found that the degree of \(\alpha,\omega\)-difunctionality should be considered when the outcome of the step growth polymerization is evaluated. For successful use in application, it has been determined that the weight average molecular weight \(M_w\) of the material should exceed 50,000 g/mol as determined by SEC. If the \(M_w\) is lower than this, the material is unable to form continuous articles, such as thin films, which are useful in packaging applications. This can be easily tested as solvent castings from HFIP by monitoring whether continuous films can be achieved or if the film forms cracks during drying. Based on this it has been concluded that the \(M_w\) should at minimum double its value during the step growth polymerization.

[0086] In the production of the present \(\alpha,\omega\)-difunctional polyhydroxy acid materials, three types of materials can theoretically be obtained. This is depicted in FIG. 4 as a result of a condensation polymerization of glycolic acid and 1,6-hexanediol. If the \(\alpha,\omega\)-difunctional compound is present in the polymer molecule, an \(\alpha,\omega\)-difunctional polyhydroxy acid material is obtained. However, if an \(\alpha,\omega\)-difunctional is not included in all polymer molecules, a material exhibiting both carboxylic acid and hydroxy functions are present in addition to the \(\alpha,\omega\)-difunctional polyhydroxy acid material. As the third possible structure is a macrocycle which is formed when the carboxylic acid and hydroxy functions of a polymer molecule intramolecurarly react.

[0087] As known for those familiar to the art, either carboxyl or hydroxy functions can react with related comonomers used for the step-growth molecular weight increase step. In some cases a comonomer may be able to react with both carboxyl and hydroxy functions. For instance, the reaction with a hydroxy group and an isocyanate yields a carbamate, or urethane, structure while the reaction with a carboxylic acid and an isocyanate yields an amide structure with carbon dioxide as a by-product. However, as can be seen from FIG. 1, the OH/COONH ratio of terminal groups, or the degree of \(\alpha,\omega\)-difunctionality, has a profound impact on the result of the step growth polymerization. Hence to achieve a sufficient molecular weight increase, in one further embodiment of this invention the degree of the \(\alpha,\omega\)-difunctionality is more than 60%, preferably more than 70% and most preferably more than 80%, when the degree of \(\alpha,\omega\)-difunctionality is determined from a quantitative \(^{1}H\) NMR spectrum.

[0088] The difunctionality is calculated from the ratios of OH and COOH terminal polymer chain as follows based on three plausible polymer structures that can be formed in the condensation polymerization process of a copolymer of an exemplary diol, 1,6-hexanediol, and glycolic acid (cases 1 to 3). Assignations of the signals from spectra are presented in FIGS. 5 and 6. Signals from glycolide are assigned as G.

[0089] In the first case there is no hexanediol in the polymer chain and therefore it is OH/COONH-terminated. A COOH-terminated end group gives characteristic peaks at \(-170.2\) ppm and \(-60.7\) ppm (carbons 10 and 9). Other assigned signals are the repeating unit at \(-168.0\) and \(-60.9\) ppm (carbons 5 and 6). OH-end group at \(-172.8\) ppm and \(-59.9\) ppm (carbons 1 and 2), and the glycolic acid unit next to the end-group at 168.7 and other signal possibly overlapping with repeating unit (carbons 3 and 4).

[0090] In the second case there is one hexanediol unit most likely in the middle of the chain. Hexanediol unit gives characteristic signals at \(-66.8\) ppm, \(-27.7\) and \(-24.8\) ppm (carbons 13, 14 and 15), and the glycolic acid monomers next to hexanediol unit give signals at \(-169.3\) and \(-61.3\) ppm (carbons 11 and 12).

[0091] In the third case it is assumed that the repeating unit is the same in the whole cyclic structure and therefore it gives one signal in carbonyl region and one signal in aliphatic region. The signals are most likely fused into the signals of the repeating units at \(-168.0\) and \(-59.8\) ppm (carbons 5 and 6), and therefore it might increase the value of calculated number average molecular weight \(M_n\).

[0092] Number average molecular masses were calculated from integrals obtained from spectra as follows:

\[
M_n = \frac{\sum M_n}{\sum} = \frac{n_{OH,COON} + n_{OH,COOH}}{n_{OH,COON} + n_{OH,COOH}}
\]  

\[n_{OH,COON} = \frac{A_{168\text{ppm}}}{2}
\]

\[n_{OH,COOH} = \frac{A_{170.2\text{ppm}}}{2}
\]

\[M_{OH,COON} = \frac{A_{168\text{ppm}}}{2} \left( M_RU + M_gal \right)
\]

\[M_{OH,COOH} = \frac{A_{170.2\text{ppm}}}{2} \left( M_RU + 3M_gal \right)
\]

[0093] As noted above, preferred classes of the chemicals that can be used to increase the molecular weight of the polyglycolic acid polymer and copolymer in a step growth polymerization process include: diisocyanates, bisepoxy compounds, dialdehydes, diketenes, phosphoric acid derivatives and bisoxazolines.

[0094] Examples of these are alkyl diisocyanates, such as butane diisocyanate, hexamethylene diisocyanate, aromatic isocyanates such as tolylene-2,4-diisocyanate, tolylene-2,5-diisocyanate, toluylene-2,6-diisocyanate, 1,4-phenylene diisocyanate, 1,3-phenylene diisocyanate, m-xylene diisocyanate, poly(hexamethylene diisocyanate).

[0095] Examples of bis-epoxy function compounds are aliphatic, aromatic and polymeric diglycidyl ethers, such as ethylene glycol diglycidyl ether, propylene glycol diglycidyl ether, neopentyl glycol diglycidyl ether, bisphenol A diglycidyl ether, and polyethylene glycol diglycidyl ether.

[0096] Examples of phosphoric acid esters are trimethylphosphate, triphenylphosphate fatty acid phosphates (e.g. stearyl phosphates), and 2,6-di-tert-butyl-4-methylphenyl phosphate.

[0097] According to present invention the compounds used in the step growth polymerization step are chosen in such a way that the molecular weight increase step can be carried out
in a twin screw extruder. Analysis has shown that appropriate manufacturing of α,ω-diunctional polyhydroxy acid mate-
rial combined with step growth polymerization comonomer results in rapid completion of the reaction under conditions of
twin screw extrusion (FIG. 7). Hence in an embodiment of the present technology the step growth polymerization step can be 
completed in less than 30 minutes, preferably less than 10 minutes and most preferably in less than 5 minutes.

As will be discussed below (cf. Example 19), gly-
colic acid polymers, for example produced by a method 
according to the present invention, exhibit properties of elon-
gation in combination with high modulus (i.e. high strength 
properties). Such materials are particularly useful in biaxial 
orientation or thermoforming applications because they will 
yield layers that are not prone to having discontinuities. Such 
polymers typically exhibit a molecular ratio between the pre-
polymer and the coupling agent, e.g. a disiocyanate, amount-
ing to 1:0.9 to 1:1.3, in particular 1:1 to 1:1.1.

The obtained telechelic material from the conden-
sation process can be fed into the extruder by appropriate 
hoppers as known for those familiar to the art. It is preferable 
that the hopper has a moisture free atmosphere in order to 
prevent undesired hydrolysis to take place. The telechelic prepolymer can be rendered or prepared with suitable equip-
ment to a grain size that can be easily handled by the hopper 
feed mechanism. The extruder configuration will preferably 
be such that it consists of four or more segments. The first 
segment is set up by screw configuration and temperatures so 
that the telechelic prepolymer melts. The second segment 
consists of an inlet for addition of the comonomer used for 
the step-growth polymerization, temperatures that permit to keep 
the material flowing in the segment and a screw configuration 
suitable for efficient mixing of the telechelic polymer and 
the step-growth prepolymer. The third segment consists of an 
inlet for addition of suitable additives for stabilizing addi-
tives, temperatures that permit to keep the material flowing 
in the segment and a screw configuration suitable for efficient 
mixing of the formed copolymer. An exhaust to remove 
potentially gaseous products also can be included in the third 
segment, or in a separate segment. Other segments may be 
added according to need if more additions or exhausts are 
required. The final segment consists of a screw configuration that 
will permit the material to efficiently exit the extruder die. 
The preferred temperatures for the segments used in the step 
growth polymerization in an extruder are 215-280°C, more 
preferably 220-250°C or most preferably 220-240°C.

For convenient addition of the comonomer used in 
the step-growth polymerization of the telechelic prepoly-
mers, it is preferred that such comonomers are in liquid form 
so they can be added to the extruder by pumps. Hence, 
comonomers that are solids at room temperature can be dis-
solved in suitable solvents which do not interfere with the 
step-growth polymerization reaction and can be easily evapo-
rated in the third segment of the step-growth process. Suitable 
solvents are polar or apolar solvents such as THF, DMSO, 
alkanes, toluene, dichloromethane. Polar solvents such as 
alcohols are not preferred.

For those familiar to the art, the amount and type of 
the comonomer used for the step growth polymerization will 
have a profound impact on the product properties. For instan-
tce, when using diisocyanates as comonomers, the obtained 
material may lack sufficient properties if too little of the 
diisocyanate is used. Similarly, the obtained material may 
be crosslinked if an extensive amount of the diisocyanate 
comonomer is used. While theories of step-growth polymeriz-
ation teach that optimal amount of the reactive comonomers 
is in an equimolar amount, it may be beneficial to deviate 
from this according to need. For instance a slight excess of 
diisocyanate may yield a polymer which possesses long-
chain branching in a suitable amount that can have a benefi-
cial effect on the melt viscosity and processability of the 

Further to retain the useful properties of polygly-
colic acid, it is important to choose the comonomers in such 
a way that crystallinity is preserved in the material. Under 
conditions described herein, such materials are formed.

For the sake of completeness it should be noted out 
that it is also possible carry out the step growth polymeriza-
tion step with prepolymer obtained by ring opening poly-
merization of corresponding lactones (glycolides), although 
the present condensation polymerization of monomers is pre-
ferred.

Various other materials or additives can be mixed 
into the material during or after the step-growth polymeriza-
tion process. Such materials may be polymers to yield blends, 
fillers and reinforcing fibres such as silica or CaCO₃, plastici-
cizers, stabilizers against light, thermally or hydrolytically 
induced degradation, glass fibres or lignocellulosic fibres.

The polymer produced can be formed using known 
processing methods for thermoplastic or solution formable 
polymers. Examples are extrusion to films, sheets, profiles, 
pipes or fibres; solvent casting or fibre spinning from solu-
tion. Moulded articles can be produced e.g. by injection 
moulding, blow moulding or thermoforming. The polymer 
can be applied as single layer material or as laminates or 
multi-layer structures.

It should finally be pointed out that although bulk 
 polymerization is described below in more detail, it is also 
possible to carry out the polymerization steps as solution 
polymerization and emulsion polymerization.

EXAMPLES

Characterization of Materials

Molecular weights and molecular weight distribu-
tions were determined with size exclusion chromatography 
(SEC). A Waters system equipped with two 7.8 mm x 300 mm 
Styragel HR 4E and HR 5E columns and Waters 2414 Refract-
ive Index Detector connected in series was used. Hexafluoro-
propanoic acid (HFIP, 5 mM CF₂COONa) was used as an elu-
ent and was delivered at a rate of 1.0 ml/min. The results 
were calculated against monodisperse polymethylmethacrylate 
standards.

Differential scanning calorimetry (DSC) was used to 
determine thermal transitions of the prepared polymers 
using Mettler Toledo DSC820 STAR® SW 9.20 instrument 
under nitrogen atmosphere. Samples were heated twice from 
0 to 250°C at a rate of 10°C/min. Thermal transitions were 
recorded from the second heating scan.

The microstructure of polymers was analysed by 1H 
NMR and 13C NMR using a Bruker 500 MHz spectrometer. 
Samples were dissolved in a 2:1 mixture of hexafluoroiso-
propanol and deuterated chloroform (CDCl₃). 13C NMR 
were acquired using broad band proton decoupling and relax-
ation delay of 3 s. Chromium (III) acetylacetonate was added 
as a relaxation reagent. Chemical shift scale was calibrated to 
TMS.
[0110] The oxygen barrier properties of prepared materials were evaluated by solution coating on Performa White board (Stora Enso). Coatings of polymer solutions (5 wt-%) in hexafluoroisopropanol were performed using an Erichsen bar coater (30 μm bar) theoretically yielding a film thickness of 1.5 μm. To ensure solvent free coatings, the coated boards were dried in ambient conditions for 30 min and subsequently in a circulating air oven at 100°C for 15 min. The oxygen transmission rate (OTR) was measured from two or more parallel samples using humid gases at room temperature (23°C, 50% relative humidity) with Systech M800i and expressed as cm³/m² day.

Example 1

[0111] 500 g solid glycidic acid, 15.6 g hexanediol (2 mol-%) and 0.26 g SnOct₃ (0.05 m-%) were added to a 1000 mL flask and reaction was performed similarly as in example 1. Temperature was increased gradually from 130°C to 190°C and pressure was decreased gradually from 500 mbar to 30 mbar during four hours. When target temperature and pressure were achieved, reaction was continued for 24 hours. Temperature was increased to 230°C and reaction was continued for two hours. Yield 392 g, Mₙ (NMR) 2000 g/mol, Mₚ (GPC) 15000 g/mol, T₆ 24°C, Tₚ 90°C, ΔH_m 15 J/g, T_m 203°C, ΔH_m -99 J/g, 65% OH-terminated.

Example 2

[0112] 500 g solid glycidic acid, 23.3 g hexanediol (3 mol-%) and 0.26 g SnOct₃ (0.05 m-%) were added to a 1000 mL flask and reaction was performed similarly as in example 1. Yield 405 g, Mₙ (NMR) 1500 g/mol, Mₚ (GPC) 10000 g/mol, Mₚ (GPC) 14 400 g/mol, T₆ 19°C, Tₚ 95°C, ΔH_m 51 J/g, T_m 203°C, ΔH_m -95 J/g, 81% OH-terminated.

Example 3

[0113] 500 g solid glycidic acid, 31.2 g hexanediol (4 mol-%) and 0.27 g SnOct₃ (0.05 m-%) were added to a 1000 mL flask and reaction was performed similarly as in example 1. Yield 410 g, Mₙ (NMR) 1300 g/mol, Mₚ (GPC) 10 300 g/mol, Mₚ (GPC) 14 400 g/mol, T₆ 14°C, Tₚ 88°C, ΔH_m 47 J/g, T_m 196°C, ΔH_m -97 J/g, 87% OH-terminated.

Example 4

[0114] 70% glycidic acid solution with high purity was distilled at 100°C and 200 mbar to remove the water prior usage. 1732 g (22.3 mol) distilled glycidic acid solution, 41 g (2 mol-%) H₂O₂ and 0.89 g (0.05 m-%) SnOct₃ were added to a 2 L flask and reaction was carried out similarly as presented in example 3. 1397.9 g white polymer was obtained. Polymer was grinded using standard Wiley mill with 2 mm sieve. Grinded PPG was added to a 250 mL flask and the flask was connected to a rotovapor supplied with argon flow. PPA was kept 24 h at 150°C. Temperature was increased to 230°C during 1 h and PPA was kept at 230°C for 5 hours. Yield 524 g, Mₙ (NMR) 3 800 g/mol, Mₚ (GPC) 14 700 g/mol, Mₚ (GPC) 26 900 g/mol, T₆ 36°C, Tₚ 2°C, ΔH_m, T_m 212°C, ΔH_m -87 J/g, 100% OH-terminated.

Example 5

[0115] 500 g solid glycidic acid, 13.2 g succinimide anhydride (2 mol-%) and 0.26 g SnOct₃ (0.05 m-%) were added to a 1000 mL flask and reaction was performed similarly as in example 1. Yield 387 g, Mₙ (NMR) 1 300 g/mol, Mₚ (GPC) 10 400 g/mol, Mₚ (GPC) 14 200 g/mol, T₆ 33°C, Tₚ 103°C, ΔH_m 42 J/g, T_m 210°C, ΔH_m -98 J/g.

Example 6

[0116] 500 g solid glycidic acid, 27.6 g PEG (Mw ~210 g/mol, 2 mol-%) and 0.26 g SnOct₃ (0.05 m-%) were added to a 1000 mL flask and reaction was performed similarly as in example 1. Yield 347 g, Mₙ (NMR) 1 800 g/mol, Mₚ (GPC) 13 100 g/mol, Mₚ (GPC) 1 3400 g/mol, T₆ 24°C, Tₚ 91°C, ΔH_m 36 J/g, T_m 208°C, ΔH_m -98 J/g.

Example 7

[0117] 500 g solid glycidic acid, 17.9 g pentaerytrol (2 mol-%) and 0.26 g SnOct₃ (0.05 m-%) were added to a 1000 mL flask and reaction was performed similarly as in example 1. Yield 390 g, Mₙ (NMR) 1 500 g/mol, Mₚ (GPC) 13 900 g/mol, Mₚ (GPC) 16 000 g/mol, T₆ 32°C, Tₚ 30°C, ΔH_m, T_m 190°C, ΔH_m -21 J/g.

Table 1

<table>
<thead>
<tr>
<th>Prepolymer properties</th>
<th>Mn (NMR) g/mol</th>
<th>Mn (GPC) g/mol</th>
<th>Mw (GPC) g/mol</th>
<th>T₆ °C</th>
<th>Tₚ °C</th>
<th>ΔH_m J/g</th>
<th>T_m °C</th>
<th>ΔH_m J/g</th>
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<tbody>
<tr>
<td>Example 1</td>
<td>2000</td>
<td>10800</td>
<td>15000</td>
<td>24</td>
<td>90</td>
<td>15</td>
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<td>10000</td>
<td>14000</td>
<td>19</td>
<td>95</td>
<td>51</td>
<td>203</td>
<td>-95</td>
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<tr>
<td>Example 3</td>
<td>1300</td>
<td>10300</td>
<td>14000</td>
<td>14</td>
<td>88</td>
<td>47</td>
<td>196</td>
<td>-97</td>
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<td>Example 4</td>
<td>3800</td>
<td>14700</td>
<td>26900</td>
<td>36</td>
<td>—</td>
<td>212</td>
<td>-87</td>
<td>—</td>
</tr>
<tr>
<td>Example 5</td>
<td>1300</td>
<td>10400</td>
<td>14200</td>
<td>33</td>
<td>103</td>
<td>42</td>
<td>210</td>
<td>-98</td>
</tr>
<tr>
<td>Example 6</td>
<td>1800</td>
<td>13100</td>
<td>13400</td>
<td>24</td>
<td>91</td>
<td>36</td>
<td>208</td>
<td>-98</td>
</tr>
<tr>
<td>Example 7</td>
<td>1500</td>
<td>13900</td>
<td>16000</td>
<td>32</td>
<td>—</td>
<td>190</td>
<td>-21</td>
<td>—</td>
</tr>
</tbody>
</table>

Example 8

[0118] 55 g of prepolymer prepared in Example 1 and equivalent amount of hexamethylene di-isocyanate (HDI) were added to a Brabender melt mixing equipment. Samples were taken at 1, 2, 3 and 5 minutes. Results are presented in FIG. 1. A solution coated film on Performa White board displayed an OTR of 4600 cm³/m² day.

Example 9

[0119] 55 g of prepolymer prepared in Example 2 and an equivalent amount of hexamethylene di-isocyanate (HDI) were added to a Brabender melt mixing equipment. Samples were taken at 1, 2, 3 and 5 minutes. Results are presented in FIG. 1. A solution coated film on Performa White board displayed an OTR of 34 cm³/m² day.

Example 10

[0120] 55 g of prepolymer prepared in Example 3 and an equivalent amount of hexamethylene di-isocyanate (HDI) were added to a Brabender melt mixing equipment. Samples were taken at 1, 2, 3 and 5 minutes. Sample taken at 5 minutes was crosslinked an insoluble. Results are presented in FIG. 1. A solution coated film on Performa White board displayed an OTR of 365 cm³/m² day.
Example 11

[0121] 55 g of prepolymer prepared in Example 4 and an equivalent amount of hexamethylene di-isocyanate (HMDI) were added to a Brabender melt mixing equipment. Samples were taken at 1, 2, and 5 minutes. Results are presented in Table 1. Thermal properties of starting material: \( T_g \) 36°C, \( T_{c-v} \) 126°C, \( \Delta T_{c-v} \) 87 J/g. Thermal properties after 1 minute: \( T_g \) 36°C, \( T_{c-v} \) 126°C, \( \Delta T_{c-v} \) 24 J/g, \( T_m \) 204°C, \( \Delta H_m \) –52 J/g. Thermal properties after 2 minutes: \( T_g \) 36°C, \( T_{c-v} \) 123°C, \( \Delta T_{c-v} \) 29 J/g, \( T_m \) 205°C, \( \Delta H_m \) –52 J/g. Thermal properties after 5 minutes: \( T_g \) 36°C, \( T_{c-v} \) 118°C, \( \Delta T_{c-v} \) 26 J/g, \( T_m \) 206°C, \( \Delta H_m \) –55 J/g.

Example 12

[0122] 55 g of prepolymer prepared in Example 1 and hexamethylene di-isocyanate (HMDI) with ratios of 0.75, 1.0 and 1.25 were added to a Brabender melt mixing equipment. Samples were taken at 1, 2, 5, 10, and 15 minutes. Results are presented in Fig. 2. Solution coated films on Performa White board displayed OTRs of 83, 92 and 83 cm³/m² day for HMDI:prepolymers ratios of 0.75, 1.0 and 1.25, respectively.

Example 13

[0123] 55 g of prepolymer prepared in Example 1 and bisphenol A diglycidyl ether (BPADGE) with ratios of 0.75, 1.0 and 1.25 were added to a Brabender melt mixing equipment. Samples were taken at 1, 2, 5, 10, and 15 minutes. Results are presented in Fig. 3. Solution coated films on Performa White board displayed OTRs of 226 cm³/m² day.

Example 14

[0124] Glycolic acid (100 kg), and hexanediol (2 mol-%) were added to a multipurpose reactor at 25-70°C over a period of 2 h. Then, SnOct₂ (0.5% m-%) was added to the molten mixture of glycolic acid and hexane. The melt condensation reaction was performed. Temperature was increased gradually during the reaction and the pressure was reduced slowly to <50 mbar. To prevent formation of glycolic such as an incremental process was chosen. After completing the reaction, the polymer (76 kg, ~100% yield) was placed in metal containers, allowed to cool down to room temperature. During cooling the material crystalized, and to obtained solid was ground to a coarse powder through a sieve (Ø 4 mm).

[0125] The coarse polymer powder was then placed into a 100L Lodige reactor. The reaction was continued in solid state by sequentially increasing the temperature from ~25°C to ~120°C and then ~160°C. During this reaction step, the polymer was held under constant, reduced pressure. Yield 63 kg, \( M_n \) (NMR) 2 800 g/mol, \( M_w \) (GPC) 8 700 g/mol, \( M_p \) (GPC) 17 500 g/mol, \( T_g \) 27°C, \( T_{c-v} \) 125°C, \( \Delta T_{c-v} \) 210°C, \( \Delta H_m \) –84 J/g, 100% O2-terminated.

Example 15

[0126] The step growth polymerization of the PGA polymer prepared in example 14 and hexamethylene diisocyanate was carried out in a twin screw extruder. Prior to the reaction the polymer was dried overnight in 120°C and allowed to cool to room temperature under reduced pressure to minimize excess moisture. The PGA prepolymer was fed to an extruder at 3 kg/h. The hopper used was continuously purged with dry \( N_2 \). A pump was used to dose hexamethylene di-isocyanate to the extruder at molar rate of 0.9x compared to PGA. The product was collected on a cooled conveyer belt and collected. In this was a polymer with Mn (GPC) 51 300 g/mol, \( M_w \) (GPC) 211 400 g/mol, \( T_g \) 30°C, \( T_{c-v} \) 265°C, \( \Delta H_m \) –48 J/g, modulus 8.5 GPa, tensile stress at yield 72.1 MPa, and tensile strain at break 3.1% was obtained.

Example 16

[0127] Example 15 was repeated by dosing hexamethylene di-isocyanate to the extruder at molar rate of 1.0x compared to PGA. Following polymer was obtained: Mn (GPC) 47 500 g/mol, \( M_w \) (GPC) 245 900 g/mol, \( T_g \) 31°C, \( T_{c-v} \) 265°C, \( \Delta H_m \) –58 J/g, Modulus 6.4 GPa, Tensile stress at yield 80.4 MPa, Tensile strain at break >100%.

Example 17

[0128] Example 15 was repeated by dosing hexamethylene di-isocyanate to the extruder at molar rate of 1.05x compared to PGA. Following polymer was obtained: Mn (GPC) 52 300 g/mol, \( M_w \) (GPC) 202 700 g/mol, \( T_g \) 32°C, \( T_{c-v} \) 265°C, \( \Delta H_m \) –63 J/g, Modulus 7.1 GPa, Tensile stress at yield 78.6 MPa, Tensile strain at break >100%.

Example 18

[0129] Example 15 was repeated by dosing hexamethylene di-isocyanate to the extruder at molar rate of 1.15x compared to PGA. Following polymer was obtained: Mn (GPC) 45 500 g/mol, \( M_w \) (GPC) 203 600 g/mol, \( T_g \) 31°C, \( T_{c-v} \) 264°C, \( \Delta H_m \) –60 J/g, Modulus 6.5 GPa, Tensile stress at yield 74.0 MPa, Tensile strain at break >100%.

Example 19

<table>
<thead>
<tr>
<th>Mn (GPC) (g/mol)</th>
<th>Mw (GPC) (g/mol)</th>
<th>( T_g ) (°C)</th>
<th>Modulus (GPa)</th>
<th>Tensile stress at Yield (MPa)</th>
<th>Tensile strain at Break (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Example 15</td>
<td>51300</td>
<td>211400</td>
<td>30</td>
<td>8.5</td>
<td>72.1</td>
</tr>
<tr>
<td>Example 16</td>
<td>47500</td>
<td>245900</td>
<td>31</td>
<td>6.4</td>
<td>80.4</td>
</tr>
<tr>
<td>Example 17</td>
<td>52500</td>
<td>202700</td>
<td>32</td>
<td>7.1</td>
<td>78.6</td>
</tr>
<tr>
<td>Example 18</td>
<td>45500</td>
<td>203600</td>
<td>31</td>
<td>6.5</td>
<td>74.0</td>
</tr>
</tbody>
</table>

Example 10

[0130] Copolymerization of glycolic acid and hexanediol was carried out in a similar fashion as described in Example 7. The obtained polymer was then copolymerized with 1,6-hexamethylene diisocyanate in a twin screw extruder. The molar ratio between the prepolymer and the diisocyanate was 1:1.

[0131] Standard mechanical test specimens were prepared by injection molding and the resultant polymer exhibited the following mechanical properties:

[0132] Tensile strength: 78 MPa

[0133] Tensile modulus: 7.5 GPa

[0134] Elongation: >100%

[0135] The present high-molecular weight aliphatic polyesters exhibit decreased discoloration, good heat and chemical resistance as well as good gas barrier properties, e.g. excellent oxygen and carbon dioxide barrier properties, and can be used alone or as composite materials or multi-layered structures in a wide variety of fields as extruded, compres-
sion-moulded, injection-moulded, blow-moulded products, and other formed or moulded products.

[0136] Further, the polymers prepared herein can be extruded to threads, spun to fibers or with suitable additives extruded to foams. A particular, but not an exclusive list of examples, is the following: multilayer films obtained after extrusion optionally combined with a subsequent blowing or biaxial orientation process for packaging applications, laminated optionally multi-layered films for packaging applications, injection molded articles for use in oil or gas drilling, injection molded articles for various parts or consumer articles, injection molded articles for use in degradable implants in bone repair, foamed products for insulation of sound or against temperature, spun fibers for cloths, textiles and sutures.

[0137] The instant polymers can be used for coating of specimens, preferably specimens selected from the group of solid objects and webs, in particular by thermal powder coating or extrusion coating.

[0138] The process may also be optimized for reactive extrusion in such way that the above products be directly produced after the described step-growth polymerization process.

[0139] Based on the above, it is possible to prepare materials which have high modulus imparting in improved rigidity of the material and may permit reduction of materials in multilayer structures, which contributes to cost saving and environmental benefits, which structures have high elongation useful in biaxial orientation or thermoforming applications yielding layers that are not prone to exhibit discontinuities; and which materials have high optical clarity in transparent films.

[0140] While the present disclosure has been illustrated and described with respect to a particular embodiment thereof, it should be appreciated by those of ordinary skill in the art that various modifications to this disclosure may be made without departing from the spirit and scope of the present disclosure.

What is claimed is:

1. A telechelic polymers of glycolic acid obtained by a process comprising the steps of providing glycolic acid; providing a difunctional monomer; subjecting said glycolic acid to condensation polymerization in the presence of said monomer; and continuing polymerization to provide a polymeric chain formed by residues derived from glycolic acid and said difunctional monomer.

2. The polymers according to claim 1, comprising at least 5, for example 5 to 250, preferably 6 to 100, in particular 10 to 50 residues of glycolic acid.

3. The polymers according to claim 1, comprising an essentially linear polymeric chain or a branched polymeric chain.

4. The polymers according to claim 1, having a degree of crystallinity of at least 50%, preferably at least 40%, in particular at least 50%.

5. The polymer according to claim 1, comprising at least 80 mol-%, preferably at least 90 mol-%, of residues derived from glycolic acid.

6. The polymer according to claim 1, comprising 0.5 to 10 mol-%, preferably 1 to 5 mol-%, of residues derived from a difunctional comonomer, e.g. a hydroxy-terminated comonomer.

7. The polymer according to claim 5, wherein the molar ratio between residues derived from glycolic acid and residues derived from the comonomer is 1000:1 or less, for example 500:1 or less, suitably 300:1 or less, advantageously 45:1 to 55:1, in particular 48:1 to 52:1.

8. The polymer according to claim 6, wherein the hydroxy-terminated comonomer is selected from the group of diols, such as propanediol, butanediol, hexanediol, pentaerythritol and oligomeric polyethylene glycol and combinations thereof.

9. The polymer according to claim 1, which is a telechelic hydroxy-terminated polymer, preferably exhibiting exclusively comonomer residues of hydroxy-terminated comonomers.

10. The polymer according to claim 7, wherein the comonomer is selected from a dicarboxylic acid, dithiol, or diamine.

11. The polymer according to claim 1, wherein the glycolic acid is subjected to condensation polymerization in the presence of the monomer and a catalyst, preferably an esterification catalyst.

12. The polymer according to claim 1, comprising 0 to 20, preferably 0 to 10 mol-% of residues of at least a second hydroxy acid, e.g. lactic acid.

13. The polymer according to claim 1, having a molecular weight (Mn) of 500 to 25,000, in particular about 750 to 15,000 g/mol.

14. The polymer according to claim 1, capable of being used as a prepolymer for producing polymers.

15. A method of producing a telechelic polymer of glycolic acid, comprising the steps of providing glycolic acid; providing a difunctional monomer; subjecting said glycolic acid to polymerization in the presence of a catalyst and said difunctional monomer; and continuing polymerization to provide a polymeric chain formed by residues derived from glycolic acid and said monomer.

16. The method according to claim 15, wherein a hydroxyl-terminated polymer chain is formed by the steps of providing glycolic acid; providing a hydroxy-terminated monomer; subjecting said glycolic acid to condensation polymerization in the presence of an esterification catalyst and said hydroxy-terminated monomer; continuously removing water formed during condensation polymerization; and continuing polymerization to provide a polymeric chain formed by residues derived from glycolic acid and said hydroxy-terminated monomer.

17. A method of producing a telechelic polyester polymer of glycolic acid, which preferably is hydroxy-terminated, optionally in combination with a method according to claim 15, wherein polymerization of the corresponding monomers is carried out in an initial solid state at 200° C. or less for a pre-longed period of time under reduced pressure, followed by a final dehydration, the initial polymerization being carried out for a time of less than 36 h and dehydration step being carried out at a time of less than 24 h.

18. The method according to claim 15, wherein condensation polymerization, optionally in combination with a dehydration step, is continued until a polymer comprising least 5, for example 5 to 250, advantageously 6 to 100, in particular 10 to 50 residues of glycolic acid is obtained.

19. A method according to claim 1 for preparing glycolic acid homo- or copolymers.
20. The method according to claims 15, wherein condensation polymerization is carried out at a temperature of 120 to 250°C, optionally in an inert atmosphere or under reduced pressure.

21. The method according to claim 15, wherein condensation polymerisation of glycolic acid is performed at temperatures below the melting point of the product polyglycolic acid, in particular condensation polymerisation of glycolic acid is performed at temperatures below 210°C.

22. The method according to claim 15, wherein the temperature of the polymerisation of glycolic acid is gradually increased and the absolute pressure decreased.

23. The method according to claim 15, wherein condensation polymerization is carried out in the presence of a catalyst selected from the group of zinc, tin or titanium octoate.

24. The method according to claim 15, wherein condensation polymerization is carried out in the presence of a comonomer, e.g. a second hydroxy acid, such as laetic acid.

25. The method according to claim 15, wherein polymerization is continued to provide an essentially linear hydroxy-terminated polymeric chain.

26. The method according to claim 15, wherein polymerization is continued to provide an essentially branched hydroxy-terminated polymeric chain.

27. The method according to claim 15, wherein polymerization is carried out in a solvent.

28. A glycolic acid polymer comprising at least two prepolymers, preferably 5 to 100 prepolymers, according to claim 1 linked together.

29. A polymer according to claim 28, comprising prepolymers linked together with chain extenders, e.g. chain extenders selected from the group of diepoxides or diisocyanates.

30. A polymer according to claim 28, retaining at least 30%, preferably at least 40% of the crystallinity or melting enthalpy of the prepolymer.

31. A polymer according to claim 28, having a molecular weight (Mn) of more than 10,000 g/mol, in particular about 20,000 to 1,000,000 g/mol.

32. A polymer according to claim 28, having an essentially linear polymer structure or an essentially branched polymer structure.

33. A method of producing a polymer according to claim 28, comprising subjecting a prepolymer to step-growth polymerization in the presence of a chain extender to provide a linear polymer having a molecular weight (Mn) of more than 10,000 g/mol, in particular about 20,000 to 1,000,000 g/mol.

34. The method according to claim 33, wherein the chain extenders are selected from the group of diepoxides and diisocyanates.

35. The method according to claim 33, carried out in an extruder, e.g. in a twin screw extruder for achieving step growth polymerization.

36. Use of polymers according to claim 28 for forming polymeric films, sheets, fibers, powders or moulded articles, in particular for forming compression-moulded, injection-moulded, blow-moulded products, and other formed or moulded products.

37. The use according to claim 36, wherein the polymers are used for forming products exhibiting properties of good heat and chemical resistance as well as good gas barrier properties, e.g. excellent oxygen and carbon dioxide barrier properties.

38. The use according to claim 36 for coating of specimens, preferably selected from the group of solid objects and webs, in particular by thermal powder coating or extrusion coating.

39. The use according to claim 36, wherein the polymer is used directly from reactive processing for extrusion or moulding.