REVIEW PAPER



Modern biopolyamide-based materials: synthesis and modification

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Received: 4 May 2018 / Revised: 30 November 2018 / Accepted: 12 February 2019 / Published online: 9 March 2019 © The Author(s) 2019

Abstract

The wider use of renewable feedstock in structural applications, where high mechanical performance is required, can be achieved by the application of recently developed engineering biopolymers and their further modification by micro- and nanoparticles. In this review, we present the current state of the art of biopolyamide materials for structural and functional applications. The overview includes all stages of the manufacturing—from the synthesis of building blocks, through the synthesis of polymers and its physical modification, with special emphasis on the properties of the obtained engineering biocomposites as a final product of modern polymer technology. In the first part, the synthetic routes of bio-derived counterparts of common polyamides as well as specialty polymers with functional properties arising from the complex structure of biochemicals were exemplified. The development of environmentally friendly composites and nanocomposites based on biopolyamides and natural fillers, such as plant fibers or cellulosic nanofibers, was of particular interest due to preserved sustainable character of such materials.

Keywords Biopolyamides · Biocomposites · Natural fibers · Cellulose fibers

Introduction

Nowadays, there is a strong need to reduce the consumption of fossil resources in the production of polymeric materials. In order to provide a new generation of renewable, eco-friendly polymeric materials the classical polymer technology needs to be alternated in numerous fields, including production of raw materials, polymerization techniques, and physical modification of biopolymers. The term 'biopolymers' refers to materials partially or fully based on renewable substrates. Among various biopolymers, polyamides synthesized from biomass-derived monomers constitute an

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interesting class with outstanding mechanical performance, such as high hardness, impact strength, and abrasion resistance, barrier and thermal properties, as well as good processability by, e.g., extrusion and injection molding. Recent research in the field of biopolyamides showed efforts to synthesise and apply monomers from renewable resources [1, 2], adapt enzyme technology for polymerization processes [3] and physically modify biopolyamides by compounding with natural reinforcing fillers and functional additives.

The areas of research and development reflect the specific market demands for biopolyamides as an engineering materials, where mechanical performance combined with functional properties is of major interest. With this respect, biopolyamide composites emerge as an attractive final materials that could be introduced in the machine, automotive, building, electrical and electronics, and consumer goods industries.

Currently, the main attention is paid to biopolyamide composites reinforced by plant fibers and cellulosic nanoparticles, since they offer improvement of mechanical performance without sacrificing the original renewable character of the material.

However, some other modifications imparting novel properties to biopolyamides are also investigated. Recently, Gorrasi et al. obtained composites containing thymol as a natural photostabilizer encapsulated in halloysite nanotubes. Both, the modulus and the photo-degradation time were increased for these composites [4]. The efficiency of natural compound introduced as a stabilizer was increased by using nanotechnology approach. Improvements of the biopolyamides mechanical and electrical behavior have recently been achieved as a result of incorporation of carbon nanoparticles. For instance, polyamide 6-based masterbatch containing carbon nanotubes was melt-mixed with a neat bioPA 4.10 and graphene nanoplatelets were added into biopolyamide 11 [5, 6]. Still, numerous alternative synthesis paths and composite materials need to be validated for biopolyamides in terms of properties and cost-effectiveness.

Main building blocks for biopolyamides

The development of bio-based feedstock is fundamental for the further progress in bioplastic production and is considered as a distinct and dynamically evolving field of chemical technology. New sustainable technologies of extraction and synthesis of a variety of biomass-derived monomers are reported in the literature [7, 8].

There are three main groups of monomers for the production of biopolyamides: (i) diamines and diacids undergoing polycondensation, where both monomers or only diacid come from renewable feedstock, (ii) amino-carboxylic acids capable for polycondensation (iii) lactams transformed into polyamides via ring-opening polymerization [9].

The main source of bio-based monomers for polyamides is castor oil, which makes up 40–60% of the castor bean [10]. Castor oil, as most of natural feedstocks, has a complex composition. It consists of a mixture of saturated and unsaturated fatty acids, and the main component is C18 ricinoleic acid. Methods of castor oil extraction from beans include mechanical pressing, solvent extraction and a combination

of these techniques [11, 12]. Further transformation of ricinoleic acid (1,2-hydroxyoctadec-9-enoic acid) leads to monomers useful in further synthesis of biopolyamides. Three functional groups in ricinoleic acid structure, namely hydroxyl, carboxyl and unsaturated bond are crucial for its further chemical derivatization [11, 13, 14]. They also make castor oil polar and therefore, soluble in alcohols and compatible with selected plasticizers [11, 15].

In Fig. 1 different routes leading to various types of biopolyamides are schematically showed [16].

Monomers used in biopolyamides synthesis can be obtained partially or fully from biomass. Currently, biomass-derived monomers with the highest industrial meaning for the synthesis of biopolyamides are 11-aminoundecanoic acid, 1,8-octanedicarboxylic acid, 1,10-decanediamine, adipic acid, caprolactam and 1,4-butanediamine. The concepts of their synthesis from natural derivatives were briefly described below.

11-Aminoundecanoic acid

11-Aminoundecanoic acid is obtained through the pyrolysis of castor oil at 450–600 °C/1 atm, in the absence of air and in the presence of water vapor. Instead of castor oil, the transesterification product of ricinoleic acid, metyl ricinoleate, can be used as a starting material [10, 17–19]. After that step, methyl undecenoate and

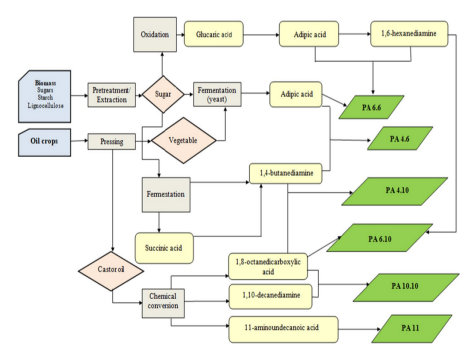


Fig. 1 Biopolyamides formation processes. Adapted with permission from Ref. [16]

heptaldehyde are obtained. The first of mentioned products is then hydrolyzed to obtain undecenoic acid, which in the next step reacts with hydrogen bromide. This reaction is carried out in a nonpolar solvent to facilitate reverse addition reaction resulting in bromoundecanoic acid formation. The last step, in which solid 11-aminoundecanoic acid is obtained, is an ammonia treatment of bromoundecanoic acid [10]. To form bioPA11 from 11-aminoundecanoic acid, water solution polycondensation followed by reactant water extraction is carried out. The manufacturing of 11-aminoundecanoic acid is well-established technology and biopolyamide 11, as a commercial material produced for decades under the trade name Rilsan, is an exception among biopolyamides.

Sebacic acid

1,8-Octanedicarboxylic acid, commonly known as a sebacic acid, is a monomer used for synthesis of biopolyamides such as PA 4.10, PA 5.10, PA 6.10 and PA 10.10. Alkali treatment of castor oil is a process which leads to separation of intermediates containing 8 and 10 carbon atoms in a molecular chain. The alkali splitting is carried out using NaOH or KOH and a catalyst at elevated temperatures [17, 19, 20]. Diamond et al. reported that the treatment of ricinoleic acid methyl ester with alkali, using 2:1 molar ratio of alkali to the ester, in the temperature range between 250 and 270 °C, leads to the formation of sebacic acid and 2-octanol [21]. Using lead tetroxide as a catalyst leads to higher yields of sebacic acid [19]. The synthesis of different types of polyamides using this diacid is performed via polymerization with diamines and the type of obtained polyamide depends on the number of carbon atom in diamine chain [22].

Adipic acid

On an industrial scale, adipic acid is produced currently from benzene. This process involves intermediate products—cyclohexane, cyclohexanol, and cyclohexanone. Nevertheless, there have been developed some bio-based pilot processes for this diacid production. According to Niu et al., there is a possibility of adipic acid production from glucose. The fermentation is conducted using modified *Escherichia coli* [23]. Additionally, there are some yeast (Candida yeast) which can be used for adipic acid production from vegetable oils and sugars [16, 24]. Another approach includes catalytic oxidation of glucose to glucaric acid, which then can be reduced to adipic acid [25]. Additionally, there are attempts to produce bio-based hexanediamine via conversion of adipic acid. Polycondensation of adipic acid and hexanediamine obtained from renewable resources leads to the production of fully bio-based PA 6.6 [16, 25].

1,10-Decanediamine

1,10-Decanediamine is a monomer which can be obtained from sebacic acid in a number of steps. At first, the sebacic acid is exposed to gaseous ammonia, so the

diammonium sebacate is formed. The next step is a dehydration process, which leads to dinitrile octamethylene dicyanide formation. After hydrogenation in the presence of potassium hydroxide and a nickel catalyst the final product—decamethylene diamine (DMDA), is obtained [2, 10, 26]. Polycondensation of this diamine with diacids is a useful path toward biopolyamide synthesis.

1,4-Butanediamine and 1,5-pentamethylenediamine

Microbial synthesis is a promising technology for low molecular mass building blocks for polymers, including diamines. For example, bio-based butane diamine can be obtained via biochemical route by fermentation of glucose with metabolically engineered strain of *Escherichia coli* [16, 27]. Diaminopentane can be produced by *Carynebacterium glutamicum* from xylose, a major constituent of hemicellulose [28] and from L-lysine [29]. Short chain diaminopentane when condensed with long-chain bio-based sebacic acid produced polyamides with comparable mechanical strength as petrochemical PA 6 and PA 6.6 at lower specific density [30].

Caprolactam

Bio-based caprolactam can be obtained from lysine derived from glucose. The process comprises heating of L-lysine salt in a solution containing an alcohol and then deaminating of the obtained intermediate [31]. Buntara et al. described a method of caprolactam synthesis from 5-hydroxymethylfurfural obtained from glucose or fructose. The first step involves 1,6-hexanediol formation, which then is converted into caprolactone. The last step is the reaction of caprolactone with ammonia to form the desired product [32].

Synthesis of biopolyamides from other bio-based monomers

There were numerous other bio-based monomers reported that underwent polycondensation and ring-opening reactions and were useful in engineering of biopolyamide properties. The type and content of the special monomers influence the basic physical characteristics of biopolyamides, such as glass transition temperature, crystallinity, melting point, mechanical performance, low-temperature behavior, and often impart specific functional properties, e.g., adhesive or optical properties. The groups of other bio-resourced building blocks and their influence on biopolyamides properties were discussed below.

Polycondensation reactions in biopolyamides synthesis can be performed using bio-based dicarboxylic acids and diamines or bifunctional amino acids. In general, the synthesis of fully bio-based dyadic polyamides is more difficult than monadic ones, because of the difficulties in biodiamines synthesis [2].

There are some bio-based monomers from fatty acids which are useful in polycondensation reactions. They offer alternative synthesis paths for long-chain polyamides which are of industrial interest due to reduced hydrogen bond density and, as a result, reduced water uptake, high dimensional stability of molded parts in variable humidity conditions, lower melting point and thus easier processing. The example is brassylic acid (1,11-undecanedicarboxylic acid) derived from the crambe seed oil [2, 33] by ozonolysis of erucic acid, a C_{22} straight-chain monosaturated acid constituting 55% to 60% of the acids in the crambe oil. A biotechnological processing of palm oil leads to ω -amino lauric acid, which is a monomer for polyamide 12 [34].

Another approach was presented by Kolb et al. who described a way of PA 16 preparation using long-chain monomer–methyl 16-aminohexadecanoate. This monomer was prepared by transformation of the mesylated fatty acids methyl esters (FAME), to the azide which was then reduced to obtain the building block for PA 16 [35]. Stempfle et al. synthesized long-chain polyamides via polycondensation reaction using previously prepared C_{19} and $C_{23} \alpha, \omega$ -diamines and α, ω -diacids, the monomers obtained from unsaturated fatty acids esters. The melting point of PA 19.23 and PA 23.23 was 156 °C and 152 °C, respectively [2, 36].

Additionally, there is a possibility of unsaturated polyamides synthesis through polycondensation reaction of substrates originating from natural fatty acids. Examples of such compounds are Z-octadec-9-enedioic acid [37], dimethyl(E)-icos-10-enedioate obtained by self-metathesis of methyl-10-undecenoate [38] and dimethyl 1,19-nonadecanedioate [39]. Unsaturated linear aliphatic polyamides could be potentially used as thermoreactive sealants, barrier and adhesive films or high-temperature rigid materials. They have also reactive species susceptible for further chemical derivatization.

Unsaturated monomers were also used for the synthesis of branched aliphatic polyamides [40]. For this purpose, unsaturated fatty acid methyl esters were subjected to hydrobromination, then the obtained bromides were further converted into azides, which were subsequently reduced to amines. The obtained aminoacids had long alkyl chains in the proximity of amine functionality that strongly influenced the intermolecular interactions of resulting biopolyamides. The melting temperatures of branched bioPA were found too low (40 and 89 °C) for practical application of homopolymers, but branched monomers could be useful for the synthesis of copolymers with tailored thermal properties. Another strategy for derivatization of unsaturated fatty acid methyl esters was direct synthesis of keto-fatty acid esters by Wacker oxidations employing oxygen as a sole re-oxidant and amine functionalization of the thus obtained keto-fatty acid esters by reductive amination [41]. In another work, unsaturated bond of fatty acid-derived methyl 10-undecenoate, methyl oleate, and methyl erucate was utilized in the thiol-ene addition reaction of cysteamine hydrochloride to obtain amine functional branched monomers [42]. Monomers were further used to obtain homo and copolymers with adjustable thermal and solubility properties.

Introduction of cyclic aliphatic structures into the PA backbone also interferes the crystalline phase formation and molecular cohesion of polymer leading to a more flexible material. For example, cycloaddition of two unsaturated fatty acids were applied to obtain monomers with cyclic moieties [43, 44].

Another group of raw materials for bio-based polymers are carbohydrates, with the cellulose obtained from non-food resources as a main potential feedstock for the industrial scale production. A big advantage of carbohydrates is that they offer a wide range of functionalities that can be implemented into the biopolymers. The examples of renewable monomers from carbohydrates are furan-2,5-dicarboxylic acid [45], carbohydrates as sources of tartaric acid [46], as well as rosin acids [2].

Furan-2,5-dicarboxylic acid (FDCA)-based furanic-aliphatic polyamides were recognized as sustainable alternatives to polyphthalamides used as high-performance heat resistant materials [47, 48]. Bio-resourced counterparts showed comparable thermal properties in terms of glass transition and melting behavior but lower melting temperatures (c.a. 160 °C) that allows energy savings during processing. Moreover, Jiang et al. demonstrated effectiveness of enzymatic polymerization by Novozym 435 with achievable high molecular mass at relatively low temperature (90 $^{\circ}$ C), as compared to conventional polycondensation techniques. They found that further lowering of polycondensation temperature is limited by the effectiveness of water and methanol (by-product) removal, the decreased solubility of FDCAbased polymers, and the restricted chain mobility of the biopolyamide below its T_o. Structural analysis showed further necessity for the process optimization in terms of polymer dispersity. Thus, application of lipase can be a solution to a challenging problem of FDCA decarboxylation and extensive N-methylation of polyamides observed in the course of conventional polycondensation of this bio-based monomer at elevated temperatures.

There is also a protocol for biopolyamides synthesis using bio-monomer–itaconic acid, produced by the fermentation of *Aspergillus terreus*. Bio-derived itaconic acid was also used in polycondensation reactions with aromatic diamines by Ali et al. to produce degradable biopolyamies [49–51]. Another recently raised building block for sustainable polymers is succinic acid derived from glucose. The present market for succinic acid is rather small but, interestingly, the production of bio-based succinic acid is much cheaper than its petrochemical counterpart and thus experiences fast growth [8]. Polyamides obtained from succinic acid, such as PA 24, PA 44 and PA 46, show high thermal resistance and mechanical strength [52–54].

Monadic polyamides can be produced via ring-opening polymerization of lactams. Apart from the bio-based ε -caprolactam, there are some lactam derivatives which can be useful in ring-opening polymerization reactions. Winkler et al. [55] prepared a novel modified ε -caprolactone starting from 1,4-cyclohexadiene (CHD) derived from polyunsaturated fatty acid methyl esters. CHD was oxidized to form cyclohex-2-en-1-one, which was subjected to the Michael addition reaction with thiols. The product was then oxidized via Baeyer–Villiger reaction. Then, after oxime formation and Beckmann rearrangement, lactam monomer designed for the synthesis of polyamides with side groups, is formed [2, 55].

The ring-opening polymerization was used by Winnacker et al. [56] to synthesize optically active polyamides using ε -lactams from terpene β -pinene—Fig. 2.

Most of the desired properties of biopolyamides such as biodegradability, water solubility, specific thermal behavior, mechanical strength and ductility are related to crystallinity. Thus, crystallinity control is one of the major tasks in biopolyamide engineering. For instance, recently van Velthoven et al. [57] proposed a new biopolyamide that contained moieties of isoidide-based diamine and dimerized fatty acid to obtain rigid but amorphous biopolymer. The chemical components were rationally selected to provide the planar structure of the monomer, enhancing the rigidity

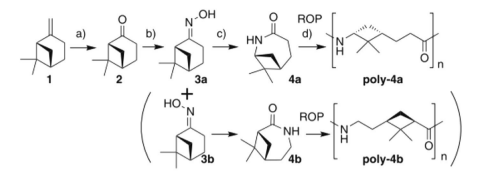


Fig.2 Synthesis of polyamide poly-4 from β -pinene; **a** KMnO₄, Al₂O₃, H₂O/CH₂Cl₂. **b** NH₄OH·HCl, NaHCO₃, MeOH/H₂O. **c** Polyphosphoric acid. **d** ROP=ring-opening polymerization. Reproduced with permission from Ref. [56]

of the polymer backbone for increased glass transition temperature, as well as steric hinderance combined with the introduction of the odd–even effect that hampered the regular packing of polymer chains into the crystal structure. Numerous other research works were focussed on the developed crystal structures, crystallization kinetic and temperatures of phase transitions of biopolyamides [58–60]. Though, further investigation on the structure-properties relationship of new synthetized biopolymers and its composites is still needed for better understanding of fundamental aspects and validation of future applications.

Using bio-based polyamides allows synthesis of high-performance, environmentally friendly materials. However, because of a decrease in intermolecular interactions with an increasing length of the alkyl spacer, long-chain biopolyamides exhibit worse properties, e.g., lower density, strength or stiffness, than polyamides with shorter hydrocarbon chains (e.g, PA 6) [9]. On the other hand, lesser amide groups density imparts lower water uptake and higher dimensional stability in the environment with varying humidity. Examples of specific biopolyamides' properties in relation to the monomer type are collected in Table 1. The physical properties of polyamides, such as glass transition and melting temperature, mechanical strength and water absorption, can be adjusted by copolymerization of monomers with different chain length and structure [33, 61].

Biopolyamide composites

The development of new biopolyamide materials is observed both in the area of chemical synthesis, as well as further physical modification. The chemical and physical modifications reflect the specific market demands for biopolyamides as an engineering material, where mechanical performance combined with functional properties is of major interest. With this respect, biopolyamide composites emerge as an attractive final materials dedicated to the machine, automotive, building, electrical and electronics, and consumer goods industries, etc.

Table 1 Examples of specific bi	Table 1 Examples of specific biopolyamide properties in relation to the bio-monomer type	type
Compound name	Chemical structure of monomer	Desired properties of biopolyamide
1,5-Diaminopentane	H ₂ NNH ₂	6% lower density at comparable mechanical strength (bioPA 5.10 vs petrochemi- cal PA 6 and PA 6.6) [30] Increased moisture regain for fibers giving more comfort in wearability, higher exhaustion of acid dyse [62]
11-Aminoundecanoic acid	HO	Modified melt properties for easier processability by synthesis of star structures [63]
1,13-Tridecane diamine	² HN	Reduced water absorption [64]
Methyl 9(10)-aminostearate and methyl 13(14)-aminodo- cosanoate	Mars	Branched, higher elasticity, lowering of melting point of copolymers [40]
1,18-Octadec-9-enedioic acid	10 ⁻¹⁰ -0-10-10-10-10-10-10-10-10-10-10-10-10-1	Long-chain unsaturated thermoreactive, susceptible for further chemical derivati- zation [37, 38]
Dimethyl(E)-icos-10-enedioate	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
Rapeseed oil-based dimer acid	ноос(H ₂ C),HC=HC H ₃ C(H ₂ O) ₅ HC=HC (GH ₂) ₅ CH ₃	Lowering of softening point, transparent and flexible at ambient temperature [44]
Piperazine	IZZI	Lowering of softening point [43]
Furan-2,5-dicarboxylic acid	HO JOH OH	Bio-resourced alternatives for the phtalamides [48]

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Table 1 (continued)		
Compound name	Chemical structure of monomer	Desired properties of biopolyamide
2,3-Di- <i>O</i> -methyl-L-tartaric acid	HO HO OMEO OH	Optical properties in solution [65]
2,3,4-Tri-O-methyl-arabinaric acid (p, L and their equimolar mixture)	HO OME MEO OH	Adding hydrophilicity and biodegradability [66]
Isoidide diamine	H N N N N N N N N N N N N N N N N N N N	Amorphous polyamides with $T_{\rm g}$ values up to 100 °C [57]
Rosin	HOOC HOOC	Photoactive polyamideimides with photoinduced birefringence, good solubility in organic solvents and processability by spin coating [67]

Bio-based polyamides reinforced by natural fibers

Using natural fibers as fillers in bio-based polymers gives an opportunity to obtain fully green, sustainable materials. Besides their renewable nature, natural fibers offer a wide range of valuable properties such as low density, low cost, higher strength per unit mass in comparison with inorganic fillers [68]. Moreover, these fibers are nontoxic and non-abrasive to the equipment used for composites preparation. However, the main difficulty arises from their polar and hydrophilic nature, which makes it difficult to obtain a good dispersion of fibers in hydrophobic polymer matrices. Another challenge of making such composites is that the processing temperature of some polyamides is higher than a degradation temperature of natural fillers [69].

Properties of natural fibers

Generally, vegetable natural fibers can be divided into three groups: seed fibers (e.g., cotton), bast fibers (e.g., jute) and leaf fibers (e.g., sisal) [70]. They all have an elongated shape with diameters in the range of 15–30 μ m and lengths in the range of 1–50 mm, but there are high fluctuations in both dimensions. As short fibers are considered the ones that have lengths up to 5 mm, and as long fibers—those with lengths up to 50 mm. Application of short fibers results in composite materials with rather isotropic properties, while the long ones lead to materials with anisotropic properties, imposed by specific orientation of fibers [71].

The chemical composition of natural fibers is influenced not only by the plant type but also by the climatic conditions, plant's age and processing conditions. The main components of natural fibers are cellulose, hemicellulose, and lignin. Other ingredients are extractives, starch, inorganics, and protein [72]. Generally, in plant fibers cellulose content increases after processing as the result of the non-cellulose components removal. Due to the high crystallinity degree of cellulose, there is a correlation between cellulose content and stiffness and strength of fibers [71, 73]. Mechanical properties depend also on the aspect ratio of fibers and openness of the fibrous material [72].

Investigation on the thermal decomposition of ten types of natural fibers (woodmaple, wood-pine, bagasse, bamboo, cotton stalk, hemp, jute, kenaf, rice husk and rice straw), provided by Yao et al. [68], showed that for most of them thermal decomposition occurs in the temperature range between 215 and 310 °C. The degradation process runs in two stages—the first one is associated with hemicellulose degradation, and the second one with degradation of lignin [70]. Results of the thermogravimetric analysis indicate that some problems may occur when natural fibers are processed at elevated temperatures to obtain composites with thermoplastic polymers. In order to improve the thermal stability of natural fibers and, moreover, their compatibility with hydrophobic polymer matrices, surface modification has to be performed by a physical or chemical method. Physical methods include corona, gamma and cold plasma treatment, whereby chemical methods include the use of coupling agents, such as silanes, impregnation with compounds compatible with the polymer matrix, mercerization (alkali treatment) and chemical coupling through "linkers" which are able to form chemical bonds with both the fiber and the polymer matrix in composite material [69, 72, 74]. Preparation of polymeric composites reinforced by natural fibers requires also careful drying of the filler, as fibers are characterized by high water absorption. The presence of water on the surface of processed components results in voids caused by water evaporation and separation of the surfaces of fibers and the matrix. Consequently, mechanical properties of polymeric materials decrease [70, 72]. Water content must be kept especially low when natural fibers are processed with polymers susceptible for hydrolysis at elevated temperatures, such as polyamides. Despite the difficulties arising from relatively low thermal stability and high water absorption, polymer composites with natural fibers are a vast group of composite materials that have been studied for the last three decades.

Properties of biopolyamide composites with natural fibers

Researches considered different approaches to develop biopolyamide/natural fibers—composites with improved properties in comparison with unmodified materials. One of the paths includes manufacturing of the composite materials in which matrix is a blend of bio-based polyamide and some polyolefins, e.g., polypropylene. Another solution is addition of the mixture of natural and synthetic fibers into the bioPAs. Moreover, some researchers used unmodified natural fibers as the reinforcement for the polyamides, while others conducted modification of fibers surfaces before dispersing them into the polymers. In this section, the influence of natural fibers on the properties of different biopolyamides will be reviewed.

The morphology and structure of biopolyamides reinforced by natural fibers

Biopolyamide 11 and its blends with polypropylene reinforced by wood fibers, carbon fibers, and the mixture of both fillers have been investigated by Armioun et al. [75]. The morphology studies of the fractured surfaces of composite materials showed that those which contained only wood fibers had better adhesion between the filler and the matrix than composites containing the mixture of natural and synthetic fibers, though the latter had lower values of the impact strength. The main fracture mechanism in case of composites containing only wood fibers was fiber breakage, while in hybrid composites there were also fiber pullouts—Fig. 3. The same fracture behavior was observed for blends of biopolyamide and polypropylene [75, 76].

Investigations done by Kuciel et al. on composites made of Hiprolon 211 and flax fibers proved a good fiber-matrix adhesion, which was confirmed by the improvement of mechanical properties. SEM images confirmed that biopolyamide composites can be processed with no thermal degradation of the fibers, as there were no visible signs of thermal damage and deformations of the fibers [9].

Another approach was reported by Nishitani et al. Investigations carried out on biopolyamide 10.10 reinforced by hemp fibers modified with sodium hydroxide solution, sodium chlorite solution (NaClO₂) and surface treated using ureidosilane coupling agent, showed that fibers after treatment with NaClO₂ and silane coupling agent had better adhesion to the matrix than those which were treated only with NaOH [77].

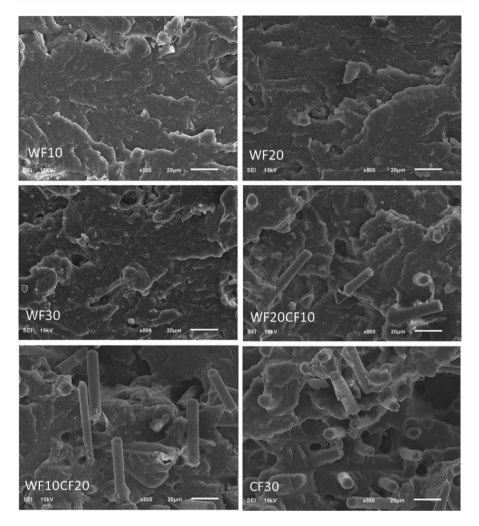


Fig.3 SEM photographs of bioPA 11 composites containing wood fibers (WF) and carbon fibers (CF). Reproduced with permission from Ref. [75]

Besides using the natural fibers, some researchers reported the use of processed cellulose fibers as the reinforcement for biopolyamides. The influence of such fibers on the mechanical and thermal properties of composites was investigated but also the length of the fibers after processing was measured. The processing methods used for composites preparation included extrusion and injection molding.

Feldmann and Bledzki have investigated the composites of polyamide 6.10 and 10.10 with 15 and 30 wt% of cellulosic fibers. The SEM pictures of those materials showed that there was low compatibility of cellulosic fibers and hydrophobic 6.10 polyamide matrix which resulted in fibers pullouts after material breakage and the presence of the gaps between the filler and the matrix. Authors prepared also the

composite containing 15 wt% of glass fibers which exhibited better adhesion with polyamide [78].

On the other hand, partial compatibility of cellulose fibers and dimer fatty acidbased polyamide was showed by Hablot et al. for the composite containing 20 wt% of cellulosic fibers. Though some holes after fibers pullouts were observed by the authors, SEM investigations indicated a good wetting of the fibers by the polyamide as the majority of fibers were cut at the level of the fracture surface [79].

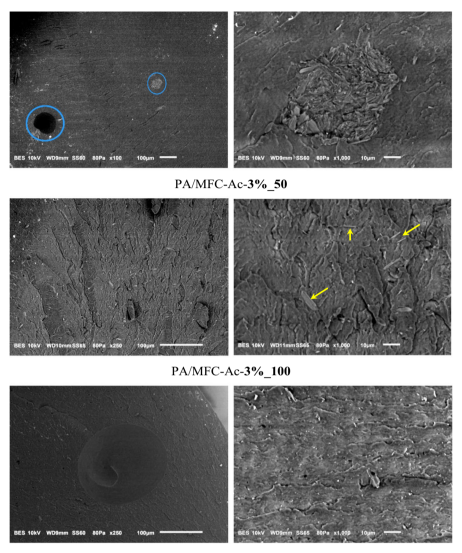
Moreover, the influence of processing parameters on the biocomposite properties was studied by Leszczynska et al. Bio-based polyamide 4.10 was modified with acetylated microfibrillated cellulose (MFC-Ac). Authors investigated the influence of screw speed during extrusion on the dispersion degree of the fibers and resulting dynamic mechanical properties. SEM analysis showed that using screw speed of 50 rpm was not enough to disperse the filler evenly. Higher values of screw speed (100 and 150 rpm) resulted in better dispersion, although at the highest speed the bubbles and defects were formed due to local overheating during processing [80]. Figure 4 shows SEM microphotographs of composites obtained at different screw speeds containing 3 wt% of the acetylated microfibrillated cellulose.

Morphology studies of composite materials prepared with different natural fillers showed that applying proper surface treatment leads to materials with good adhesion of the filler to the matrix, and thus materials in which efficient stress transfer from matrix to the fiber will be achieved.

Mechanical properties of biopolyamides reinforced by natural fibers

The main role of natural fibers in polymer composites is to improve their mechanical performance. Investigations on mechanical properties of biopolyamide 11 reinforced by wood fibers showed that with the increase in fibers content, the tensile strength, tensile modulus, flexural strength and flexural modulus increased. In the case of biopolyamide 11 containing 30 wt% of wood fibers, these parameters were 43.3 MPa, 1.78 GPa, 69.9 MPa and 1.80 GPa, while for neat PA 11 35.4 MPa, 1.04 GPa, 53.5 MPa and 1.10 GPa, respectively [75]. The same dependence of the tensile strength and fiber content was observed for polyamide 11 with stone groundwood fibers. However, after addition of 60 wt% of fibers the tensile strength decreased [81, 82]. Improvement of tensile properties in composites is related to the ability of transferring the stress between the polymer and the fiber and indicates a good interfacial adhesion between the filler and the matrix.

Studies of shear properties of biopolyamide 11 and flax fiber composites provided by Le Duigou et al. [83] proved that there is a linear (increasing) dependence between the shear stiffness and the fiber content, whereas there was no strong influence of the natural fibers content on the shear strength of the biopolyamide. Generally, incorporation of natural fibers into bio-based polyamides improved mechanical properties such as tensile and flexural strength and modulus. These observations were confirmed in other studies on biocomposites containing flax and beech fibers [9, 74]. Unfortunately, the impact strength decreases after addition of the filler, which is the weak point of composites containing natural fibers. Both notched and unnotched impact strengths have significantly lower values for composite materials



PA/MFC-Ac-3%_150

Fig. 4 SEM photographs of PA 4.10/MFC-Ac composites prepared at different screw speeds. **a** 50 rpm, **b** 100 rpm, **c** 150 rpm. Reproduced with permission from Ref. [80]

than for the neat polyamide 11. Investigation of mechanical performance of dimer fatty acid-based polyamide composites showed that the Yung's modulus and the yield stress of the materials increased with increasing content of cellulose fibers which indicated good dispersion of the fibers and sufficient interactions between the filler and the matrix [79, 84].

Cellulose fibers efficiently improve mechanical performance of polyamides, and thus, such materials are another alternative for obtaining environmentally friendly

materials for high-performance applications. However, some reports showed superior mechanical behavior of biopolyamide reinforced by synthetic fibers or mixed natural/synthetic reinforcement as compared with natural fillers. Although the composites comprising a mixture of natural and synthetic fibers showed inferior properties as compared to composites containing solely carbon fibers, they were still interesting from the viewpoint of the cost/weight ratio [74-76]. In hybrid composites, with the increasing content of the carbon fiber, the tensile and flexural strength of the composites was enhanced. The same trend was observed for tensile and flexural modulus. Such results arise from the higher strength and modulus of carbon fibers in comparison with the natural ones. Tensile modulus for the hybrid composite containing 10% of wood fibers and 20% of carbon fiber was improved by 168%, in comparison with composite with only wood fibers, and the flexural modulus was enhanced by 142%. It is worth to mention that incorporation of synthetic fibers also improves notched and unnotched impact strength of the composites containing natural fibers. The same results were obtained for composites where the matrix was made of biopolyamide and polypropylene blend [75, 76].

The influence of compounding process conditions on the mechanical properties of composites made of PA 6.10 and PA 10.10 was investigated by Feldmann and Bledzki. Single-step and two-step compounding techniques were applied, and it was found that tensile and impact strength have higher values in the one-step process due to the lower thermal and mechanical stresses imparted on the material than in the two-step approach. Further investigations of mechanical properties on these biopolyamides containing 15 and 30 wt% of cellulosic fibers were compared with composites containing 30 wt% of glass fibers. Composites with 30 wt% of cellulosic fibers had two times higher values of tensile strength (120 MPa) than neat polyamide, however, still lower than that of composites containing glass fibers. On the other hand, biopolyamide reinforced with natural fibers was superior to composite with glass fibers in terms of elongation at break and notched impact strength [78].

The mechanical properties of biopolyamide 10.10 containing chopped cellulose fibers were also investigated by Nikiforov et al, who compared the properties of such materials with the composites of PA 10.10 containing carbon and glass fibers. Results showed that the highest values of tensile elastic modulus were obtained for composites containing carbon fibers (6.34 GPa for 6 vol% of fibers, and 10.15 GPa for 12 vol%). At the same volumetric filling, the composites containing cellulose fibers had values of elastic modulus 2.27 and 2.74 GPa, respectively [85].

Thermal and thermomechanical properties of biopolyamide composites

Understanding of thermal properties of composites reinforced by natural fibers is an important issue because of the relatively low thermal stability of these fibers in comparison with polymer matrix. Nishitani et al. investigated thermal properties of biopolyamide 10.10 reinforced by hemp fibers treated with various chemicals, e.g., NaOH, NaClO₂, 3-(2-aminoethylamino) propyltrimethoxy silane. The results of the thermogravimetric analysis (TGA) showed that the thermal stability of neat polyamide is higher than composites in the whole temperature range. Additionally, degradation of unmodified polyamide was a single-stage process, while for the composite materials there were two stages of thermal decomposition. Authors explain that the first stage, occurring between 80 and 200 °C, is related to the degradation of hemicellulose and lignin, as well as to the absorbed water evaporation. The second stage, at 300 °C, is probably the degradation process of cellulose and polymeric matrix. Moreover, TG data revealed that in the case of composites with modified hemp fibers, thermal stability of composites containing fibers treated with the combination of sodium hydroxide and amino silane is higher than those in which fibers were treated with only one of the chemicals, and also higher than those containing raw fibers [77].

The investigations carried out on biopolyamide 11 reinforced by wood fibers showed some similar effects as those discussed above, namely the thermal decomposition of composites proceeded in two stages. The first stage, which started at 200 °C, was associated with the degradation of fibers, while the second one at 350 °C with the polyamide degradation. This indicated that thermal stability of composites containing wood fibers is lower than neat biopolyamide 11. Nonetheless, authors noticed that with an increase in the fiber content, the onset temperature of the second decomposition stage slightly got higher as a result of possibly good interaction of fibers and polyamide. It was also observed that the rate of the thermal decomposition was decreasing after the addition of synthetic fibers, and thus, enhancing the thermal stability of composites. [75].

Some studies of the thermal behavior of biopolyamides composites with natural fibers indicated that using raw fibers would result in deterioration of materials thermal stability. The perspective for resolving this problem is a prior modification of the fibers before composites processing or manufacturing of biocomposites containing natural and synthetic additives. Chemical modifications of natural fibers presented in the literature include alkali, acetyl, silane, benzyl, acryl, permanganate, peroxide, isocyanate, titanate, zirconate and acrylonitrile treatments and use of maleated anhydride grafted coupling agent. Additionally, environmentally friendly enzyme treatment has been gaining more and more interest recently [86, 87]. Except improvement of fibers thermal stability, chemical modification enhanced compatibility between fibers and matrix, and thus positively affected their mechanical performance.

Thermal properties of different biopolyamides and composites of bioPA 4.10 containing 1, 3, 5 wt% of cellulosic fibers were further investigated by Leszczynska et al. [80] and Pagacz et al. [89]. The approach employed by the authors included modification of cellulose fibers before composites preparation. Surface acetylation of microfibrillated cellulose (MFC) by acetic anhydride resulted in substantial thermal stability enhancement due to the non-cellulose components removal and partial esterification of hydroxyl groups; the onset temperature of degradation was reported to be 277 °C for the raw fibers, while for the fibers after modification this temperature was above 316 °C [80]. Melting temperatures of composites determined by DSC method were slightly decreased as the cellulose content increased, while the crystallization temperatures were shifted to the higher values after addition of the filler due to heterogeneous nucleation of polyamide crystallization. Dynamic mechanical analysis showed that the highest value of storage modulus was obtained for the composite containing 5 wt% of cellulosic filler extruded at screw speed of

150 rpm, although at this screw speed material's deterioration may occur due to overheating effects [80, 88, 89]. Moreover, Leszczynska et al. investigated the effect of surface modification of microfibrillated cellulose by acid chlorides on the structural and thermomechanical properties of biopolyamide 4.10 nanocomposites produced by melt blending. Dynamic mechanical analysis (DMA) showed that there is a dependence on composite performance on the length of alkyl chain attached to the surface of cellulose fibers. Modification of the fibers with acid chlorides with longer alkyl chains resulted in decrease in storage modulus in comparison with the shorter ones (Fig. 5) [90].

Also some studies of biopolyamide composites containing unmodified cellulosic fibers are presented in the literature. Investigations provided on dimer fatty acidbased polyamides showed that the introduction of cellulosic fibers resulted in a significant increase in glass transition temperature, from -10 to -3 °C for pure polyamide and composite containing 20 wt% of the fibers, respectively. Increasing of the glass transition temperature with increasing cellulose fiber content may be explained by the formation of interactions between polyamide carbonyl and cellulose hydroxyl groups which reduced the mobility of polymer chains. It is worth to mention that the reduction of polymer chains mobility also induced the decrease in the crystallization temperature [91]. Results obtained from dynamical mechanical thermal analysis showed that the addition of cellulosic fibers into dimer fatty acid biopolyamides results in increase in storage modulus with increasing cellulose content [79, 84, 91].

Other properties of biopolyamides reinforced by natural fibers

Other properties which are important for the efficient processing and further applications of composite materials based on biopolyamides and natural fibers are melt flow index (MFI), heat deflection temperature, and water uptake. As incorporation of fibers causes a decrease in the mobility of macromolecules, the values of MFI are lower for composite materials in comparison with unmodified polyamide and decrease with the increasing fiber content. The temperature at which polymer deforms under a particular load is the heat deflection temperature (HDT). According to Armioun et al. HDT for biopolyamide 11 was 44 °C and no significant improvement of this parameter was observed after incorporation of natural fibers. High water absorption is one of the disadvantages of natural fibers because the presence of water during composites processing can result in dimensional instability, surface defects, and decrease in mechanical properties. In composite materials based on biopolyamide 11 and wood fibers, the water uptake increased with an increasing fiber content. The highest value of water absorption was observed for composite containing the highest addition of wood fibers (30 wt%), for which this parameter increased by 70% in comparison with neat PA11 [75, 76].

Studies of different biopolyamides reinforced with natural fibers indicated that such green materials can be manufactured and used as an alternative for some petrochemical plastics. The main approach included manufacturing of hybrid composites containing natural and synthetic fibers to achieve a compromise between the mechanical and thermal performance and a cost, as well as density and environmental aspects. Beside the most widely described composites of bioPAs with

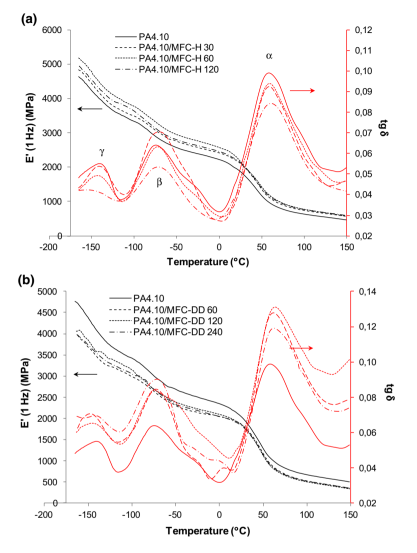


Fig. 5 Bending storage modulus and tg δ for PA 4.10 composites containing 3 wt% of MFC modified with hexanoyl chloride (**a**) and dodecanoyl chloride (**b**) obtained after different times of reaction. Reproduced with permission from Ref. [90]

natural fibers, also biopolyamides composites containing processed cellulose fibers are described in the literature as a promising material.

Biopolyamides reinforced by other additives

Although particular attention is paid to the bio-based polyamide composites with natural or cellulose fibers or mixtures of those natural fillers with glass or carbon fibers, there are also reports on biopolyamides composites reinforced by carbon fibers, boron nitride microparticles, composites containing nutshell powder modified with nanosilver particles or rice husk and nanoclay [92–95]. For instance, Hajibeygi and Omidi-Ghallemohamadi have synthesized a semi-aromatic bio-based polyamide via triphenyl phosphite-activated polycondensation reaction and prepared Mg–Al layered double hydroxide containing composites with enhanced thermal stability [96]. There are also some investigations of thermal stability and flammability of polyamide 11 composites with lignins or nanoboehmite, which causes the increase in thermal stability and reduction of heat release and mass loss rate [97, 98]. Such materials offer novel applications for bio-based polyamides.

Carbon fibers

Mechanical properties of biopolyamide 10.10 reinforced by carbon fibers were described by Kuciel et al. Authors investigated two different types of polyamide 10.10. The filler contents were 10 and 30 wt% and 20 and 40 wt%, respectively. Mechanical performance of all composite materials was better than those of unmodified polyamides. Values of such parameters as tensile strength and modulus or flexural strength and modulus increased with an increasing fiber content, e.g., for neat biopolyamide those parameters were 51.4 MPa, 2030 MPa, 62.5 MPa and 1676 MPa, respectively, while for its composite containing 40 wt% of carbon fibers they took values of 184.9 MPa, 23,645 MPa, 283.0 MPa, 19,830 MPa, respectively [92]. The substantial improvement of mechanical performance was also observed for biopolyamide 11 reinforced by carbon fibers [75, 92]. Another advantage from carbon fibers incorporation into bio-based polyamide matrix is the reduction of water absorption [76, 92].

Boron Nitride

In order to produce environmentally friendly materials useful in electronic packaging applications, Mosanenzadeh et al. investigated the properties of bio-based polyamide 30 and 90 that were melt blended with boron nitride microparticles and then compression molded—Fig. 6.

The choice of this filler comes from its high thermal conductivity and electrical resistivity. Authors analyzed such properties as thermal conductivity, thermal expansion and electrical insulation, as well as mechanical and rheological properties of composites. Results showed that the effective thermal conductivity of polyamides and boron nitride composites increases with an increasing filler amount. Higher values of this parameter were obtained for the samples in which PA 30 was used as a matrix. Moreover, because of the insulative properties of both components, the

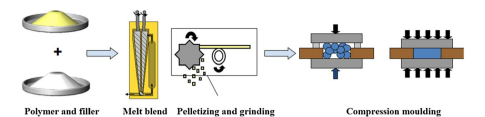


Fig. 6 Preparation of biobased polyamides composites containing boron nitride particles. Reproduced with permission from Ref. [94]

composites were shown to be electrical insulators. Mechanical properties, such as hardness and impact strength, were reduced after addition of the boron nitride filler, but at the same time the thermal properties were improved with increasing filler content [94].

Rice husk ash and nanoclay

Mechanical properties of composite materials consisting of bioPA 10.10 or bioPA 6.10 and rice husk ash and nanoclay were investigated by Battegazzore et al. Results showed that the addition of rice husk ash results in significant improvement of Young's modulus, reduction of tensile strength and large reduction of deformation at break which may be related to the low efficiency of stresses transferring from matrixes to the filler. The best results of mechanical performance and interactions between polyamides and fillers were obtained after additional incorporation of nanoclay to the composite. Such results were related to the aspect ratio and sub-micrometric exfoliation of montmorillonite [95].

Lignins

The influence of the lignin type and content on the thermal properties and fire retardancy of biopolyamide 11 composites were investigated by Mandlekar et al. The authors used four types of lignins that differed in resource, purity and chemical nature. The investigations showed that the addition of the sulfonated lignins provided better thermal stability and higher char formation than the addition of kraft lignins (Fig. 7). Additionally, flame retardancy tests showed that the addition of 15 wt% of both filler types is optimum to improve the flame retardancy [97].

Summary

A wide range of new biopolyamides with promising mechanical performance and functional properties, but with varied and not fully recognized industrial potential has been developed in the past years from renewable building blocks. The progress in the synthesis of biopolyamides relies on the development in production of biobased platform chemicals, advanced polymerization techniques and understanding

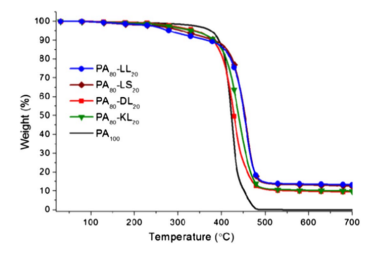


Fig. 7 TG curves, in nitrogen for PA/lignin microcomposite containing 20 wt% of lignin. Reproduced with permission from Ref. [97]

of structure–property relationship of newly obtained macromolecules. The monomers used for the synthesis of biopolyamides included diacids, diamines, aminoacids and lactams obtained from renewable resources, with a chemical structure analogue to commonly used petrochemical monomers, but also numbers of more sophisticated bio-resourced compounds. The important advantage of bio-based over the petrochemical sourced monomers is that the former often bear functional groups that impart new functional properties to obtained specialty polymers. Making use of biochemicals, a complex and diverse group, is both promising, due to new possible properties of polymers unavailable through petrochemical routes, and challenging due to efficiency and cost issues.

In order to fulfil specific market demands for biopolyamides as an engineering material, where mechanical performance combined with functional properties is of major interest, the chemical approach is supplemented with physical modification of biopolyamides. In the context of composite manufacturing, the main attention was paid to biopolyamides reinforced with natural fibers. That environmentally friendly composite materials offer enhanced mechanical properties including tensile and flexural strength and modulus. However, there are still some challenges in fabrication of bioPAs composites with natural fibers. A relatively low thermal stability of natural fibers in comparison with biopolyamide matrices was the main difficulty in the preparation scheme, which induced surface modification and careful selection of components and processing parameters. Another problem was high water absorption of natural fibers which caused technological difficulties during processing. Thus, chemical modification conducted on the fibers prior to composites processing or using blends of biopolyamides with hydrophobic polyolefines as matrices for biocomposites were applied. The use of hybrid reinforcement composed of natural fibers and synthetic ones was shown as an alternative approach for manufacturing biopolyamide materials with improved thermal and mechanical behavior. Hybrid

composites are currently a feasible compromise between mechanical properties and the cost, density and environmental aspects. Biopolyamides filled with natural fibers, including cellulosic ones, can be a good choice for structural and high-performance applications, where other sustainable materials do not fulfil demands, such as natural fiber plastic composites (NFPCs) based on polyolefins or polylactide. Biopolyamides modified with functional additives as materials for special applications, e.g., in electronic, packaging or textiles industries, are in the initial phase of development.

Acknowledgements This research work was supported by Polish National Science Centre under the Grant No. UMO/2015/19/B/ST8/01060.

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