Synthesis and Study of Thermo Stability Properties of Novel Polyamides

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Abstract

Aromatic and aliphatic diamines are one of the main components in polyamides, polyimides, polyurea and polyurethanes. To prepare high performance and well featured materials, attempts is directed to the novel diamine synthesis routes in which soluble thermo-resistant, soluble and easy process polymers are obtained. To solubility improvement and decreasing the glass transition temperature, existing flexible bonds in polymer repeating unit are necessary. These bonds decrease the internal rotational energies. Using heterocyclic rings or introducing of polar functional groups such as amide in the main synthetic polymer chain results in convenient polarity and solubility of polymer. Until now, massive investigations have been conducted to prepare aromatic monomers with high solubility and processing capabilities. In the current study, our main aim is to prepare diamines which are applicable in thermo-resistant polymers including polyamide, polyimide, polyurea and polyurethanes.

Keywords: Diamine; Thermo-resistant polymer; Polyamide; Polyimide; Polyurethane

Introduction

Amines being a main component of amino acids, peptides and containing nitrogen compounds are the most abundant organic molecules. Primary and secondary amines are less acidic and form weaker hydrogen bonds with alcohols and ethers compare to alkaloids. On the contrary, they are stronger base and nucleophiles compared to the alcohols and ethers. Diamines are amines with two functional amine groups. Hexamethylene diamine is an important amine which is used in nylon manufacture and it reacts with adipic acid to produce nylon 6, 6. This nylon is used in socks, stretch cloths and textile fibers. Many investigations have been performed on diamines that some of them are referred in continue. Effect of EDTA and Municipal solid waste compost on different amount of lead and cadmium in soil and plant decontamination of these elements by sunflower was investigated [1]. A cationic catalyst of iron complex has catalyzed an amination of aryl-aziridine to high yield production a diamine. This method is compatible with both existing functional groups in amine and aziridine [1]. Amination of meso-en-phenyl aziridineis progressed by 1% of mentioned catalyst to produce a diamine with high yield [2]. Activating of aziridine thorough a SN_2 reaction is the main reason of urea formation [11]. In this study, we deal with synthesis and characterization of novel monomer which are used in various polymers. Diol, diacid, diisocyanate and diamines are typical monomers which are applicable in preparation of different polymers such as polyamide, polyurethane and polyimides. Diamines occupied here, are among of the most important monomers which have been used during recent years. There are two kinds of diamines: aliphatic and aromatic diamines. Polymers prepared from aromatic monomers are commonly more thermo-resistant and less soluble. Therefore, synthesis of semi-aromatic diamines seems to be more soluble than aromatic ones. So, semi-aromatic diamines seem to be more thermo-resistant and less soluble. Therefore, synthesis of semi-aromatic diamines is the aim of many investigations, as they have suitable solubility and thermal resistance.

In addition, novel polyamides having high thermal resistivity were synthesized successfully. In addition to high thermal resistance, these polyamides had improved solubility and showed high processing properties. To synthesis these polyamides, first a diamine with well-defined precursor was prepared and then reacted with different acyl chloride and dicarboxylic acids to obtain these novel polyamides. It was clear that such thermal resistivity is due to amide structure and rigid phenyl rings. Also, solubility and processability will improve by introducing aliphatic groups into the polyamide structure.

Experimental

Materials and instruments

Nitric acid Fuming, Benzyl, 4-Nitrobenzaldehyde, Ammonium acetate, Hydrazine hydrate, dichloride acid and solvents were purchased from Merck Co, and Fluorene prepared from Sigma Aldrich Co. 1H-NMR and 13C-NMR spectra were recorded on a 500 MHz
Bruker Advance Dpx400 apparatus using DMSO-d6 as solvent and tetra methyl silane as standard. FTIR spectra were recorded by a FT-IR Spectrometer Spectrum RXI from Perkin Co. The CHNS-Rapid Heraeus was used for elemental analysis. Thermal gravimetric analysis was performed by TGA device model PL-TGA from Polymer Laboratories. Inherent viscosity was measured with an Ubbelohde suspended-level viscometer at room temperature and melting point was measured using Electrothermal model Engineering LTD.

Monomer synthesis

**Synthesis of 2, 7-dinitrofluorene:** Initially, mixture of 10 g (0.06 mol) fluorene and 50 ml glacial acetic acid at 0-5°C were entered a three-neck balloon equipped with mechanical stirrer while stirring for 0.5 h (Scheme 1). While the solution was stirring, 50 ml fuming nitric acid was added during 45 min. After nitric acid addition, reaction temperature ascended to 65°C and after that mixture allowed to cool down and stand for a night. An orange precipitate is formed through this procedure. Next, the mixture was transferred to the 500 ml water/ice and stirred for 1 h. The mixture was filtered and washed several times with deionized water. The product was dissolved in 150 ml chloroform and after washing several times with water and salt, the orange phase was separated and dried with magnesium sulfate. After solvent removal, orange viscose phase was separated and dissolved in 100 ml hexane. After placing the container into ice-water, the yellow precipitate which is related to 2, 7-dinitrofluorene was appeared. Yield of reaction was 79% and melting point was about 295-300°C.

**Scheme 1: Structure of 2, 7-dinitrofluorene.**

**Fluorene based diamine synthesis:** Mixture of 7 g (0.028 mol) 2, 7-dinitrofluorene, 0.5 g palladium/C (5%) and 150 ml ethanol was transferred to a 250 ml balloon equipped with stirrer and condenser while refluxing (Scheme 4). Mixture was stirred under nitrogen atmosphere and then 4.1 mmol triphenyl phosphate which have already been dissolved in a solvent mixture (30 ml NMP and 6 ml pyridine) containing dissolved LiCl (0.6 g) and CaCl₂ (1.8 g) was added to the reaction vessel. The mixture was stirred under nitrogen for 7 h at 110°C then the mixture was transferred into the methanol and filtered, after refluxing by methanol dried at 120°C. Reaction yield was 90% and melting point was over 400°C (PA1).

**Scheme 4: Structure of Schiff base diamine monomer (LH₄).**

20 ml hydrazine hydrate (85%) dissolved in 25 ml ethanol was drop wise added to the reaction container during 1.5 h. The mixture was refluxed for 2 h and another 0.1 g off rest catalysis added at different time distances during the refluxing time. After filtering, obtained precipitate was dried at 60°C for 5 h. Then the precipitate was dissolved in 100 ml THF and refluxed at 70°C. After filtering, the resultant solution was transferred to the cold water and opaque white crystallized product appeared. Finally, precipitate was dried at 60°C for 5 h. Yield of reaction was 78% and the melting point was about 245-248°C.

**Polyamide synthesis:** For the synthesis of polyamides, dicarboxylic acid and di-acyl chloride were used. Schematic of synthesis procedure is shown in Scheme 5.

**Scheme 2: Structure of 2, 7-diaminofluorene.**

**Polyamide synthesis via dicarboxylic acid:** Firstly, 2 mmol fluorene based diamine and 2 mmol biphenyl-4, 4-dicarboxylic acid were transferred into a two-necked balloon equipped with condenser, under nitrogen atmosphere and then 4.1 mmol triphenyl phosphate which have already been dissolved in a solvent mixture (30 ml NMP and 6 ml pyridine) containing dissolved LiCl (0.6 g) and CaCl₂ (1.8 g) was added to the reaction vessel. The mixture was stirred under nitrogen for 7 h at 110°C then the mixture was transferred into the methanol and filtered, after refluxing by methanol dried at 120°C. Reaction yield was 90% and melting point was over 400°C (PA1).
Polyamide synthesis by di-acyl chloride: Triethyl amine was added to a 100 ml balloon containing 2 mmol dissolved diamine in 20 ml THF at 0°C under nitrogen atmosphere while stirring. 0.5 g and then 2 mmol dissolved isophthaloyl dichloride in 10 ml THF, was added to the reaction mixture dropwise. After 3 h stirring at room temperature, the mixture was transferred into the cold water where the product filtered and washed with NaHCO₃ and water. Finally, the obtained polymer was dried at 70°C. The final product melting point was 380°C (PA3).

In another synthesis, we prepared the polyamide in a similar way of PA3, but instead of isophthaloyl dichloride, adipoyl dichloride (PA4) was used.

Results and Discussion

Monomer synthesis and characterization

The monomers, 1-2, were prepared according to the method given in the literature. The characteristics data obtained from FTIR and ¹H-NMR spectra as listed in Table 1, confirmed the structures illustrated in Schemes 1-4. The interesting absorption bands in the FTIR spectrum of monomer 4, are: 3452 cm⁻¹ and 3369 cm⁻¹ (related to stretching vibration of NH₂) (Figure 1). Aromatic C-H vibration peak appear on 3028 cm⁻¹. Corresponding stretching vibration peaks of C=O are on 1473 and 1382 cm⁻¹ and peaks on 1290 and 1617 cm⁻¹ are assigned to C-N and C=O vibrations.

In ¹H-NMR spectrum monomer 4(LH₄₄), of Figure 2, the peak on 5.2 ppm is related to NH₂. With envisaging the fact that, two NH₂ groups are present in structure and if the peak areas relating to these two NH₂ groups are considered 4 and other peaks evaluated regard to this peak, one can conclude that, the peaks higher than 6 ppm are related to aromatic hydrogen's which integration of these peak areas is equal to 34 verifying the number of aromatic hydrogen's. The peak on 2 ppm is related to methylene hydrogen's which has been split due to diastereotopic effect. Integration of peak areas was 2, proving the number of methylene hydrogen's.

As shown in Figures 3a and 3b, in ¹³C-NMR spectrum, a peak on 30.38 ppm is related to CH₃, and the peaks in region of 113-148 ppm, is related to aromatic carbons.
Figure 3a: $^{13}$C-NMR spectrum of LH$_4$.

Figure 3b: $^{13}$C-NMR spectrum of LH$_4$.

Polyamides synthesis and characterization: As illustrated in Scheme 5, the polyamides were prepared by polycondensation of LH$_4$ with aliphatic and aromatic dicarboxylic acid dichlorides under N$_2$ atmosphere. Elemental analysis of PA4 is presented in Table 2. That is in good agreement with the predicted structure. The inherent viscosity [n] of the polyamides determined at concentration of 0.125 g/dL in NMP and 25°C was in the range of 0.31-0.57 dl/g. FTIR spectra of the polyamides showed characteristic absorption bands in the regions of 3298-3362 cm$^{-1}$, 1629-1661 cm$^{-1}$ and 1269-1285 cm$^{-1}$, related to the N—H, C = O and C—N stretching, respectively. $^1$H-NMR spectra of PA1 and PA2 are shown in Figures 4 and 5. $^1$HNMR spectrum of PA1 resembles to $^1$HNMR of monomer 4 but chemical shift in protons of methyl and amide groups in PA1 can attribute to the polymer formation. Peaks at 1.9 and 2.1 ppm are related to the methyl protons and peak at 588 ppm verifies the amide protons. Peak of aromatic hydrogens appears in region of 7.15 to 7.48 ppm. For PA2, peaks at 0.93 and 1.3 ppm are related to aromatic methylene groups and a peak at 2.3 ppm corresponds to the methylene in amide structure. Peaks at 6.8 and 7.8 pertain to aromatic hydrogens and a peak at 10.7 ppm relates to amide hydrogens. Chemical shift toward down fields is due to hydrogen bond formation of amide group.

Figure 4: $^1$H-NMR spectrum of PA1.

Figure 5: $^1$H-NMR spectrum of PA2.

| Substrate     | IR (KBr, cm$^{-1}$)                      | $^1$HNMR (DMSO-d$_6$, δ ppm)         | $^{13}$CNMR (DMSO-d$_6$, δ ppm) |
|---------------|-----------------------------------------|--------------------------------------|---------------------------------|
| Monomer (1)   | 3088, 2917 (Ar-H), 1593 (C=C), 1344, 1529 (NO$_2$) | 8.24, 8.28, 8.46 (H, phenyl), 4.22 (2H, methylene) | -                               |
| Monomer (2)   | 3137 (NH$_2$), 2950 (Ar-H), 1475, 1560 (C=C) | 7.26, 7.29, 7.33 (6H, phenyl), 3.13 (2H, methylene) | -                               |
Table 1: Spectra data results of monomers.

| Monomer (3)                                                                 | IR (cm⁻¹)                  | Elemental analysis                                                                 |
|-----------------------------------------------------------------------------|-----------------------------|-----------------------------------------------------------------------------------|
| 3088, 2917 (Ar-H), 1344, 1529 (NO₂), 1539 (C=C)                              | -                           | 6.41-7.82 (3H, phenyl), 5.2 (4H, amine), 1.77, 1.93 (2H, methylene)               |

| Monomer (4)                                                                 | IR (cm⁻¹)                  | Elemental analysis                                                                 |
|-----------------------------------------------------------------------------|-----------------------------|-----------------------------------------------------------------------------------|
| 3306, 3347 (NH₂), LH4 2919, 3033 (Ar-H), 1634 (C=C), 1297 (C-N)              | -                           | 113-148 (54C, phenyl), 30.38 (2C, methylene)                                     |

Table 2: Spectra data and elemental analysis results of polyamides.

Solubility of PAs: One of the main goals of this investigation was to prepare Schiff base polyamides with enhanced solubility. For solubility test, 50 mg of polymer was dissolved in the solvents listed in Table 3. PA2 and PA4 are more soluble compared to the others. Easier solubility of these polymers in weaker solvents is due to aliphatic groups existing in polymer structure, as these groups enhance the flexibility of the chains. Comparison of PA1 and PA3 solubility proved more presence of phenyl groups in PA1, due to lower solubility of PA1.

Thermal properties of PAs: Thermal stability of the PA1 and PA3 has been investigated by TGA measurements under N₂ atmosphere at heating rate of 10 K/min. Initial weight loss (T₀) below 150°C is related to the surface water desorption. 10% weight loss occurred in the region of 400-450°C. Mass of the rest polymers at 600°C was in the range of 48-50%. Results are presented in Table 4. As results illustrate, PA1 is more thermal resistant than PA3. While polymer temperature ascends, chemical bonding will start to vibration and by increasing the temperature, the bonding vibrations will increase till the degradation starts. It can be concluded that, phenyl groups in polymer result in higher thermal resistivity and rigidity of the polymer. PA1 has higher thermal resistance compared to the PA2 because it has one phenyl group more than PA2.

Table 3: Solubility of the polyamides where Soluble (+); partially soluble (±); insoluble (-). Solubility tested with 50 mg of polymer in 10 mL of solvent; NMP=N-methylpyrrolidone; DMF=Dimethylformamide; DMSO=Dimethylsulfoxide; THF=Tetrahydrofuran.

Table 4: Data of TGA where a: 10% weight loss, b: 20% weight loss, c: The char yields of the polyamides at 600°C.
Conclusion

Four new Schiff-base polyamides were synthesized from the reaction of a novel diamine monomer (LH$_4$) with dicarboxylic acid and dichlorides. Thermal properties of these polyamides were studied by TGA. The char yields of these polyamides at 600°C were in the range of 48%-50%, showed their excellent thermal resistance. The bulky phenyl groups in the polyamides caused the enhanced thermal resistance and introduction of aliphatic groups into polyamides chain resulted in improved solubility.

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