Synthesis and Characterization of New Aromatic Co-Polyamides

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Six new aromatic Co-polyamides CoP₁-CoP₆ were prepared by direct Yamazaki's polycondensation reaction of various aliphatic and aromatic dicarboxylic acid (adipic acid, phthalic acid, terephthalic acid and 4-phenylenediacrylic acid) with new various types of aromatic diamine monomers: monomers containing flexible linkages (methylene group), Schiff-base diamine monomer and diamine monomer containing pyridine heterocyclic group and bearing bulky aromatic pendant groups in the 4-position of the pyridine ring, in the presence of LiCl in pyridine and triphenyl phosphate as condensing agents in N-methyl-2-pyrrolidinone as solvent. 4-Phenylenediacrylic acid (PPDAA) was prepared by the condensation reaction of terephthalaldehyde with malonic acid in the presence of pyridine. The monomers were characterized by FTIR and ¹H NMR. The resulting polyamides were characterized by FTIR and ¹H NMR and their physical properties including solubility, thermal stability and thermal behaviour were studied as well. All of these new polymers show good solubility in polar aprotic solvents and very good thermal stability.

Keywords: Co-polyamides, Aromatic Co-polyamides, Aromatic diamine monomers.

INTRODUCTION

Thermal characteristics are very important for all polymers processed by moulding, injection or spinning. This applies not only to the melting temperature, but also the temperature of decomposition, the crystallization temperature and other characteristics. In addition, the enthalpies of melting and crystallization are important for semi-crystalline polymers as well. The DSC study of all components of blends (used for any product prepared via melting) therefore gives basic information about the components' future behaviour in the blend and during processing.

Some properties of homopolymers are not always satisfying for their application in certain fields. Copolymer monomer based on a certain monomer with some amount of another, functional co-monomer may have specific characteristics which predetermine this copolymer as raw material for products with better properties, or as an additive for homopolymer and may thus improve the properties of the product. Copolymers are able to improve many properties, including thermal properties.

Copolymer has one great advantage, good compatibility with relevant homopolymers and from this point of view the deterioration of the blend properties is lower. Semi-crystalline (block) copolymers can co-crystallize with homopolymers, several type of crystals with various sizes and levels of perfection can be formed and blocks of copolymer can create their own crystallites.

During the copolyamide formation both mechanisms are applicable, but different to such an extent that there are formed block rather than statistic copolymides.

In the present paper the synthesis of a new types of aromatic Co-polyamides from various aliphatic and aromatic dicarboxylic acid (e.g., adipic acid, phthalic acid, terephthalic acid and 4-phenylenediacrylic acid) with new aromatic diamines, has been reported whose light scattering and dilute solution behavior have been published elsewhere. The polycondensation is carried out in the presence of triphenyl phosphite (TPP) in N-methyl-2-pyrrolidinone (NMP), pyridine mixture, containing 3 wt. % LiCl by the method of Yamazaki et al. Because of the low solubility of Co-polyamides, the polycondensation reaction terminates quickly and results in low molecular weight products. In order to do away with this problem, the polycondensation is carried out in the presence of LiCl.

EXPERIMENTAL

Fourier transform infrared (FTIR) spectra were recorded on a SHIMADZU-FTIR-8400S spectrometer (Japan) with KBr pellets in the optical range of 4000-400 cm⁻¹. ¹H NMR spectra were registered using a Bruker, 250 MHz, spectrometer, at Polymer Laboratories Co. Iran using DMSO as a solvent.
Differential scanning calorimetry (DSC) were measurements by DSC 131 Evo, Setaram, (France) using a heating rate of 10 °C/min in N\textsubscript{2} atmosphere within the temperature range of (30-600 °C). The sample weight used approximately (10 mg) mg. The peaks are used to determine the thermal properties of the samples. The solubility of the polymers was determined with (0.01 g) g of a Co-polymer in (2 mL) of a solvent.

\(N\)-Methyl-1-2-pyrrolydinone (NMP) from (ALFA-PRODUCTS); absolute methanol, acetic acid, lithium chloride, palladium on charcoal 10 \%, from (BDH, England); dichloromethane from (Biosolve); ammonium acetate, malonic acid and adipic acid from (Chem-supply). Hydrochloric acid, acetamide, 1,4-phenylene diamine, phthalic acid, terephthalic acid, piperidine, pyridine, terephthalaldehyde, triphenyl phosphite (TPP), tetrahydrofurane (THF), all from (MERCK); absolute methanol, acetic acid, lithium chloride, methane from (Biosolve); ammonium acetate, malonic acid (PAPP) \([M1]\): 

\[
\text{CHO} + 2 \text{NH}_2 + \text{Cl} \rightarrow \text{H}_2\text{N} - \text{N} - \text{H} + \text{Cl} \tag{60 \% \text{H}_2\text{SO}_4; 40 \% \text{methanol}} \text{120 °C; 10 h,} \tag{Scheme-IV}
\]

\[\text{Synthesis of 4-phenyl-2,6-bis(4-nitrophenyl)pyridine (PNPP): In a round-bottomed flask (150 mL) with a reflux condenser, a mixture of benzaldehyde (1.6 g, 15 mmol), p-nitro acetonitrile (5 g, 30 mmol), ammonium acetate (15 g) and glacial acetic acid (37.5 mL) was refluxed at 140-142 °C for 2 h. Upon cooling, crystals separated, which were filtered and washed first with acetic acid (50 \%) and then with cold ethanol. These dark yellow crystals were recrystallized from absolute ethanol and then dried at 60 °C under vacuum to produce 4.88 g (82 \% wt.) of very-deep yellow crystals: m.p. 240-244 °C (Scheme-I):} \]

\[\text{Synthesis of 1,4-Bis(6-aminopyridin-2-ylimino)-dimethylene benzene [M2]: This monomer was prepared by} \]

\[\text{Synthesis of 2-[bis(4-aminophenyl)methyl]phenol [M3]: To a 0.1 M solution of H}_2\text{SO}_4 \text{ in methanol (60 \%; 40 \%)}, \text{aniline (2 g, 21 mmol) and salicylaldehyde (1.2 g, 10 mmol) were added and the mixture was refluxed at 120 °C for 10 h with constant stirring. Upon completion of the reaction, the solvent was removed under vacuum and to the residue water (2 x 25 mL) and then CH}_2\text{Cl}_2 (50 mL) were added. The organic extract was dried over anhydrous Na}_2\text{SO}_4 to yielding 2.3 g (79 \% wt.) of deep yellow crystals: m.p. 75-95 °C (Scheme-IV).} \]

\[\text{The other diamino compounds were prepared by the same procedure as above using 3-chloro aniline with salicylaldehyde (M5), 2-bromo aniline with salicylaldehyde (M6), 3-chloro aniline with acetone (M7), 2-bromo aniline with acetone (M8), respectively, are shown in (Table-1) (Schemes V to VIII).} \]
A typical procedure was followed as described above. In a three-necked flask equipped with a reflux condenser and N₂ inlet, dicarboxylic acid and diamines (Table-2) were added. Then (2 g) dried lithium chloride, (14 mL) N-methyl-2-pyrrolidinone, (2.8 mL) water, acetic acid and acetone and then dried in a vacuum oven at room temperature to give 5.1 g (90 % wt) of white crystals; m.p. 222-225 °C (Scheme-X and XI).

**Synthesis of 4-phenylenediacrylic acid [PPDAA]:** In a 100 mL round-bottomed flask were added terephthal-aldehyde (3.48 g, 26 mmol), malonic acid (8.27 g, 94 mmol) to 30 mL of pyridine containing small amount of piperidine. The reaction mixture was stirred for 2 h at 45 °C, 4 h at 80 °C and 3 h at 110 °C, respectively. The solution was poured into large amount of distilled water and neutralized with 10 % HCl to obtain white precipitate. The precipitate was filtered, washed with water, acetic acid and acetone and then dried in a vacuum oven at room temperature to give 5.1 g (90 % wt) of white crystals; m.p. 222-225 °C (Scheme-X and XI).

**Synthesis of Co-polyamides [CoP₁-CoP₆]:** A typical procedure was followed as described above. In a three-necked flask equipped with a reflux condenser and N₂ inlet, dicarboxylic acid and diamines (Table-2) were added. Then (2 g) dried lithium chloride, (14 mL) N-methyl-2-pyrrolidinone, (2.8 mL) pyridine and (2.8 mL) triphenyl phosphite were charged. The reaction mixture was stirred for 2 h at 45 °C, 4 h at 80 °C and 3 h at 110 °C, respectively. The solution was poured into large amount of distilled water and neutralized with 10 % HCl to obtain white precipitate. The precipitate was filtered, washed with water, acetic acid and acetone and then dried in a vacuum oven at room temperature to give 5.1 g (90 % wt) of white crystals; m.p. 222-225 °C (Scheme-X and XI).

The FTIR spectrum of (M3) indicated absorption bands at (1676-1622 cm⁻¹) show the presence of the aromatic ring and (1550-1520 cm⁻¹) to heteroaromatic ring (C=N). In the FTIR spectrum of (M1), the characteristic absorptions of the nitro group disappeared and (-NH₂) stretching absorption bands of the amino group appeared at (3380 cm⁻¹).

**RESULTS AND DISCUSSION**

The FTIR spectrum of (PNPP) indicated absorption band at (1500 cm⁻¹), (1340 cm⁻¹) to (-NO₂) asymmetric and symmetric stretching, respectively, absorption bands around (1676-1622 cm⁻¹) show the presence of the aromatic ring and (1550-1520 cm⁻¹) to heteroaromatic ring (C=N).

In the FTIR spectrum of (M1), the characteristic absorptions of the nitro group disappeared and (-NH₂) stretching absorption bands of the amino group appeared at (3380 cm⁻¹).

1H NMR spectrum of (M1), is assigns the following chemical shifts: δ(2.5) ppm for DMSO, 5.6 (s, 4H) for NH₂ group, δ(6.1-8.5) ppm (s, 15H ) for ArH group.

The FTIR spectrum of (M2 Schiff-base) which indicates absorption bands at (3625 cm⁻¹) to (-NH₂ group), (3070 cm⁻¹) to (aromatic -CH stretching), (2916 cm⁻¹) to (aliphatic -CH stretching), absorption bands around (1610-1542 cm⁻¹) show the presence of the aromatic ring and (1519-1504 cm⁻¹) to heteroaromatic ring (C=N).

The FTIR spectrum of (M3) indicated absorption bands at (3440 cm⁻¹) to (-NH₂ group), absorption band at (3370 cm⁻¹) to (OH group), (3108 cm⁻¹) to (aromatic -CH stretching), (2916 cm⁻¹) to (aliphatic -CH stretching), absorption bands around (1606-1540 cm⁻¹) show the presence of the aromatic ring. The characteristic absorption of (C=O aldehyde) disappeared.

The FTIR spectrum of (M4) indicated absorption bands at (3415 cm⁻¹) to (-NH₂ group), absorption band at (3388 cm⁻¹)
Scheme-X: Synthesis of CoP₁-CoP₃
Scheme XI: Synthesis of CoP₆-CoP₄
to (-OH group), 3107 cm\(^{-1}\) to (aromatic -CH stretching), 2918 cm\(^{-1}\) to (aliphatic -CH stretching) and absorption bands around (1608-1510 cm\(^{-1}\)) show the presence of the aromatic ring and (721 cm\(^{-1}\)) to (C=O amide). The characteristic absorption of (C=O aldehyde) disappeared.

1\(^{1}\)H NMR spectrum of (M4), assigns the following chemical shifts; δ (2.5) ppm for DMSO, δ(5.8) ppm (s, 1H) for C-H group, 6.2 (s, 4H) for NH\(_2\) group, δ(6.3-8.6) ppm (s, 10H) for ArH group, 6.01 (s, 1H, OH) δ (2.135) ppm (s, 2H ) for N-OH group.

The FTIR spectrum of (M5) indicated absorption bands at (3425 cm\(^{-1}\)) to (-NH\(_2\) group), absorption band at (3100 cm\(^{-1}\)) to (-OH group), (2932 cm\(^{-1}\)) to (aliphatic -CH stretching) and absorption bands around (1614-1521 cm\(^{-1}\)) show the presence of the aromatic ring and (552 cm\(^{-1}\)) to (C=O amide). The characteristic absorption of (C=O aldehyde) disappeared.

The FTIR spectrum of (M6) indicated absorption bands at (3425 cm\(^{-1}\)) to (-NH\(_2\) group), absorption band at (3100 cm\(^{-1}\)) to (-OH group), (2932 cm\(^{-1}\)) to (aliphatic -CH stretching), (2918 cm\(^{-1}\)) to (aliphatic -CH stretching) and absorption bands around (1608-1510 cm\(^{-1}\)) show the presence of the aromatic ring and (725 cm\(^{-1}\)) to (C-Cl). The characteristic absorption of (C=O aldehyde) disappeared.

The FTIR spectrum of (M7) indicated absorption bands at (3415 cm\(^{-1}\)) to (-NH\(_2\) group), absorption band at (3000 cm\(^{-1}\)) to (-OH group), (3000 cm\(^{-1}\)) to (aromatic -CH stretching), (2932 cm\(^{-1}\)) to (aliphatic -CH stretching) and absorption bands around (1614-1523 cm\(^{-1}\)) show the presence of the aromatic ring and (554 cm\(^{-1}\)) to (C-Br). The characteristic absorption of (C=O aldehyde) disappeared.

The FTIR spectrum of (CoP\(_1\)) indicated absorption bands at (3378 cm\(^{-1}\)) to (-OH group), (3208 cm\(^{-1}\)) to (aromatic -CH stretching), (2901 cm\(^{-1}\)) to (aliphatic -CH stretching), absorption bands around (1610-1536 cm\(^{-1}\)) show the presence of the aromatic ring, the sharp band at (1616 cm\(^{-1}\)) to (C=O amide) and (1544 cm\(^{-1}\)) to vinyl segment.

1\(^{1}\)H NMR spectrum of (CoP\(_2\)), assigns the following chemical shifts; δ (2.5) ppm for DMSO, δ (8.5) ppm for HC=CH δ (6.3-8.6) ppm (s, 10H ) for ArH group, 5.7 (s, N-H) group.

The FTIR spectrum of (CoP\(_3\)) which indicated absorption bands at (3335 cm\(^{-1}\)) to (-OH group), (3123 cm\(^{-1}\)) to (aromatic -CH stretching), (2915 cm\(^{-1}\)) to (aliphatic -CH stretching), absorption bands around (1603-1511 cm\(^{-1}\)) show the presence of the aromatic ring, the sharp band at (1621 cm\(^{-1}\)) to (C=O amide), (1524 cm\(^{-1}\)) to vinyl segment and (808 cm\(^{-1}\)) to (C=Cl).

1\(^{1}\)H NMR spectrum of (CoP\(_4\)), assigns the following chemical shifts; δ (2.5) ppm for DMSO, δ (6.2) ppm (s, 1H) for C-H group, 6.0 (s, 4H) for NH\(_2\) group, δ (6.2-8.9) ppm (s, 10H) for ArH group, 5.7 (s, 1H, OH) group.

The FTIR spectrum of (CoP\(_5\)) which indicated absorption bands at (3200 cm\(^{-1}\)) to (aromatic -CH stretching), (2923 cm\(^{-1}\)) to (aliphatic -CH stretching), absorption bands around (1610-1549 cm\(^{-1}\)) show the presence of the aromatic ring, the sharp band at (1619 cm\(^{-1}\)) to (C=O amide), (1537 cm\(^{-1}\)) to vinyl segment and (748 cm\(^{-1}\)) to (C-Br).

The FTIR spectrum of (CoP\(_6\)) which indicated absorption band at (3210 cm\(^{-1}\)) to (aromatic-CH stretching), absorption bands around (1654-1633 cm\(^{-1}\)) show the presence of the aromatic ring, (1570-1511 cm\(^{-1}\)) to heteroaromatic ring (C=N) and the sharp band at (1588 cm\(^{-1}\)) to (C=O amide).

1\(^{1}\)H NMR spectrum of (CoP\(_7\)), assigns the following chemical shifts; δ (2.5) ppm for DMSO, δ (6.1) ppm (s, 1H) for N-H group, δ (6.7 -8.5) ppm for Ar-H group.

The FTIR spectrum of (CoP\(_8\)) which indicated absorption bands at (3213 cm\(^{-1}\)) to (aromatic -CH stretching), (2943 cm\(^{-1}\)) to (aliphatic -CH stretching), absorption bands around (1608-1538 cm\(^{-1}\)) show the presence of the aromatic ring, the sharp band at (1630 cm\(^{-1}\)) to (C=O amide) and (818 cm\(^{-1}\)) to (C-Cl).

The FTIR spectrum of (CoP\(_9\)) which indicated absorption band at (3315 cm\(^{-1}\)) to (-OH group), (3207 cm\(^{-1}\)) to (aromatic -CH stretching), absorption bands around (1645-1627 cm\(^{-1}\)) show the presence of the aromatic ring, (1547-1498 cm\(^{-1}\)) to heteroaromatic ring (C=N), the band at (1634 cm\(^{-1}\)) to (C=O amide) and (687 cm\(^{-1}\)) to (C-Br).

**Differential scanning calorimetry study:** From the DSC curves, the glass transition temperature (T\(_g\)), the crystallization temperature (T\(_c\)) and the melting temperature (T\(_m\)) were measured. The endothermic peaks of copolyamides are related to melting temperature. This increased in T\(_m\) is assigned to the linear terephthalic acid and p-phenylenediacrylic acid moieties incorporated into the polymer back bone during synthesis of co-polyamide. Hence, T\(_m\) is higher for CoP\(_1\), CoP\(_2\), CoP\(_3\) and CoP\(_4\) as the polymer chain more rigid parts and/or a polymer contains less free volume. Samples usually showed multiple endotherms which are explained as due to the fusion of different population of crystallites with different sizes. As expected , T\(_m\) values are lower for CoP\(_5\) and CoP\(_6\) due to the higher chain flexibility of the formers afforded with increasing the amount of methylene of polymeric back bone and steric hindrance of terminal functional groups can influence T\(_m\) of resultant polymers. In addition, the polymer CoP\(_6\) has the lowest thermal stability than the other polymers containing rigid pyridine and phenylene moieties. This behaviour can be explained by the presence of methylene units which are more vulnerable to thermo-oxidative processes\(^{21-23}\).

Glass transition temperature (T\(_g\)) of all co-polyamides were in the range (236-254 °C), in case of copolyamides (CoP\(_1\), CoP\(_2\), CoP\(_3\), CoP\(_4\), CoP\(_5\) and CoP\(_6\)) it is interesting to note that the change of the diacids had a noticeable T\(_g\) difference. In addition, endothermic peaks above their glass transition temperatures were observed in DSC scans, which may be attributed to the crystalline molecular structure for all copolymides. The higher in T\(_g\) of (CoP\(_1\), CoP\(_2\), CoP\(_3\) and CoP\(_4\)) compared to that analogous co-polyamides might be endorsed to the existence of heteroaromatic pyridine rings and phenylene moieties. This increased in T\(_g\) is assigned to the linear terephthalic and p-phenylenediacrylic acids moieties incorporated into the polymer back bone during synthesis of co-polyamide. Particularly, the end group contribution becomes significant\(^{24}\). Thus it will be interesting to introduce different chain ends using substitution reaction since end group modification in (CoP\(_1\),
Co-P, Co-P, and Co-P had a strong effect on Tg as observed in previous studies. Usually, amine polar functional group causes Tg to be higher. Tg of co-polyamides may be strongly dependent on the inter-association of functional groups with the neighboring molecule as well as the volume fraction occupied by terminal groups. It seems that several factors such as the rigidity, polarity, length and steric hindrance of terminal functional groups can influence Tg of resultant polymers. DSC data indicated that the Co-P, containing symmetric dimethyl substituted resulted in lower Tg value than that of unsubstituted co-polyamides. The consideration of segmental symmetry is useful in explaining this behavior. The dimethyl substitution leads to an symmetric segment which can result in less efficient chain packing and hence more free volume that led to relatively easier chain mobility of polymer segments in comparison with asymmetrical substituted and unsubstituted Co-polyamides, ultimately leading to decrease in glass transition temperature. The Tg, Tm, and Tc obtained from DSC are reported in Table-3.

### Table 3

| Samples | Tm (°C) | Tg (°C) | Tc (°C) |
|---------|---------|---------|---------|
| Co-P1   | 510     | 248     | 272     |
| Co-P2   | 531     | 254     | 288     |
| Co-P3   | 461     | 236     | 272     |
| Co-P4   | 502     | 240     | 264     |
| Co-P5   | 510     | 250     | 290     |
| Co-P6   | 416     | 238     | 290     |

Solubility of Co-polyamides: Solubility of Co-polyamides Co-P,-Co-P was qualitatively tested in organic solvents and the results are summarized in (Table-4). The method that attempt to enhance their processabilities and solubilities were either by introducing bulky groups, flexible linkages, or molecular asymmetry into the polymer backbones. In this work, the attachment of bulky pendant groups in polymer backbone not only could provide an enhanced solubility because of decreased packing density and crystallinity, but also could impart an increase in Tg by restricting the segmental mobility.

One of the major objectives of this work was producing Co-polyamides with improved solubility. The solubility was investigated as 0.02 g of polymeric sample in 5 mL of solvent. All of the newly synthesized Co-polyamides have good soluble in common polar and dipolar aprotic solvents without need for heating.

Conclusion

Six new aromatic Co-polyamides have been synthesized. For this purpose, a typical polycondensation process has been followed under mild conditions. The reaction variables, such as temperature, time, monomer concentration, solvent system etc. have been found to have strong influence on the molecular weight of the polymers. Synthesis of Co-polyamides containing methylene unit, Schiff-base linkages and pyridine hetero cyclic ring with adipic acid, phthalic acid, terephthalic acid and 4-phenylenediacrylic acid, respectively at 120 °C for 2 h under the nitrogen atmosphere. The polymers were characterized by proton nuclear magnetic resonance spectroscopy and infrared spectroscopy confirmed the molecular composition of six strictly alternating, highly ordered Co-polyamides having methylene chains as flexible spacers and phenylene or pyridine heterocyclic groups as the rigid segments. From the DSC curves, the glass transition temperature, the crystallization temperature and the melting temperature were measured. All the polymers showed a glass transition temperature, in the range of 236-256 °C. The higher in Tg of (Co-P, Co-P, Co-P, and Co-P) compared to that analogous Co-polyamides might be endorsed to the existence of heteroaromatic pyridine rings and phenylene moieties. This increased in Tg is assigned to the linear terephthalic and p-phenylenediacylic acids moieties incorporated into the polymer back bone during synthesis of Co-polyamide. In the same time, the presence of symmetric dimethyl substituted together with ether linkages brings much more flexibility to the macromolecular chain and decreases the glass transition of the polymers Co-P, and Co-P. The melting temperatures of the Co-P, and Co-P, decrease with increasing the amount of methylene of polymeric backbone and steric hindrance of terminal functional groups. The Co-P has the lowest melting point among the series of the result and Co-polyamides. Tm is higher for Co-P, Co-P, Co-P, and Co-P as the polymer chain more rigid parts and/or a polymer contains less free volume. All of these new polymers show good solubility in polar aprotic solvents and good thermal stability.

### Table 4

| Solvent     | Co-P1 | Co-P2 | Co-P3 | Co-P4 | Co-P5 | Co-P6 |
|-------------|-------|-------|-------|-------|-------|-------|
| DMAc        | ++    | ++    | ++    | ++    | -     | -     |
| DMF         | ++    | ++    | ++    | +     | ++    | -     |
| NMP         | +     | -     | ++    | ++    | ++    | ++    |
| Pyridine    | -     | ++    | ++    | ++    | ++    | ++    |
| n-Cresol    | +++   | +++   | +++   | ++    | +     | -     |
| THF         | ++    | ++    | ++    | +     | -     | -     |
| CHCl3       | -     | -     | ++    | ++    | ++    | ++    |
| CHCl2       | ++    | +++   | +++   | ++    | +     | -     |
| DMSO        | +++   | +++   | +++   | ++    | ++    | ++    |
| Conc. H2SO4 | ++    | ++    | ++    | -     | -     | -     |

Full soluble +++; Soluble at room temp. ++; Partially soluble +.
14. S. Mehdipour-Ataei, M. Hatami and M. Hossein Mosslemin, *Chinese J. Polym. Sci.*, **27**, 781 (2009).
15. S. Mehdipour-Ataei, Y. Sarrafi, M. Hatami and L. Akbarian-Feizi, *Eur. Polym. J.*, **41**, 491 (2005).
16. I. Kaya, A. Bilici and M. Sacak, *J. Inorg. Organomet. Polym.*, **19**, 443 (2009).
17. R.J. Sarma and J.B. Baruah, *Dyes Pigments*, **61**, 39 (2004).
18. K. Faghihi, *J. Sci. I.A.U.*, **18**, 56 (2008).
19. T. Chen, M. Zhang and X. Tang, *Chinese J. Polym. Sci.*, **26**, 793 (2008).
20. S. Mehdipour-Ataei and H. Heidari, *Macromol. Symp.*, **193**, 159 (2003).
21. D. Wilson, H.D. Stenzemberger and P.M. Hergenrother, *Polymides*, Glasgow Black and Sons (1990).
22. S.-H. Hsiao and L.-M. Chang, *High Perform. Polym.*, **12**, 285 (2000).
23. R. Hariharan, G. Anuradha, S. Bhuvana, M. Saroja Devi and M. Anitha, *J. Polym. Res.*, **11**, 239 (2004).
24. S. Mehdipour-Ataei, M. Hatami and M. Hossein Mosslemin, *Macromolecules*, **27**, 1968 (1994).
25. K.L. Wooley, C.J. Hawker, J.M. Pochan and J.M.J. Frechet, *Macromolecules*, **26**, 1514 (1993).
26. B.I. Voit, *J. Polym. Sci. A Polym. Chem.*, **38**, 2505 (2000).
27. X. Zhao, C. Wang, L. Chen and M. Zhu, *Colloid Polym. Sci.*, **287**, 1331 (2009).
28. Asian J. Chem.