Study of selected thermoplastics using dynamic mechanical analysis

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Abstract The aim of given paper is to study selected polymers using dynamic mechanical analysis method (DMA). DMA is one of the most useful techniques for the study of the viscoelastic behaviour of thermoplastic polymers. In relation to DMA, an oscillatory stress and strain is applied to the material at specific frequencies and temperatures and based on this mentioned fact hereinbefore, the resulting changes after the loading in the material are measured. This technique allows detecting the melting temperature and the glass transition temperature of the thermoplastic materials. Furthermore, some spectroscopy techniques, such as energy dispersive X-ray spectroscopy (EDX) and infrared spectroscopy (IR), were also used for the investigation of the thermoplastics. The thermoplastics used for examination, namely polyethylene, polystyrene, polypropylene and polyethylene terephthalate, were gained from the waste of the packaging.

Keywords: Dynamic mechanical analysis, infrared spectroscopy, dispersive X-ray spectroscopy, thermoplastics, packaging

1 Introduction

Dynamic mechanical analysis (DMA) is a method commonly used to measure the viscoelastic behaviour of plastic materials [1]. Plastics are never purely liquid or purely solid, because they exhibit both viscous and elastic characteristics when undergoing deformation. Oscillatory stress and strain is applied to the material at appropriate frequencies and temperatures [2]. Based on this mentioned fact, the resulting changes after the loading in the material are measured. The use of DMA technique along with differential scanning calorimetry (DSC) allows detecting the melting temperature and the glass transition temperature of the thermoplastic materials [3].

Infrared spectroscopy is widely used for characterization of various materials, such as plastics, ceramics or chemicals. Infrared spectrum helps us to identify or characterize an unknown material, and also analyze contaminants or chemical differences in observed material [4]. Infrared spectroscopy can be also used for analyzing additives in plastic compounds. Along with the energy dispersive X-ray spectroscopy (EDX), which provides

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elemental and chemical analysis of the sample, this method has become more important for the identification of potential harmful substances occurring in packaging [5, 6]. Materials for packaging can not contain toxic substances and have to be tough and flexible enough to protect the food from damage and to keep its freshness [7]. Nowadays, the most commonly used thermoplastics in the manufacture of food packaging are polyethylene, polystyrene, polypropylene and polyethylene terephthalate [8, 9].

2 Materials and methods

For the experiment, two types of low density polyethylene (LDPE), polypropylene (PP) and polyethylene terephthalate (PET) were selected as the examined materials. The thermoplastics used for examination were gained from the waste of the packaging.

Dynamic mechanical analysis was performed by the Q800 DMA from TA Instruments. The dimensions of each sample for DMA were 45 x 10 mm. The testing temperature was from -70 °C to 100 °C for LDPE, from -70 °C to 140 °C for PP, from -70 °C to 160 °C for PET at a heating rate of 3 °C per minute and frequency was 10 Hz.

The DSC measurements described in this paper were performed using Mettler Toledo, TGA/DSC 2 instrument. The weight of the samples was about from 10 to 20 mg and they were heated from 0 °C to 350 °C at a heating rate of 10 °C min⁻¹.

In Fourier transform infrared spectroscopy (FTIR), the spectra were measured by ATR technique using a Nicolet iS50 FT-IR spectrometer with a resolution of 4 cm⁻¹ in the range of 4000 - 400 cm⁻¹. The elemental and chemical analysis of the samples, were carried out using energy-dispersive EDX-7000 X-ray fluorescence spectrometer. All measurements were carried out at the CEDITEK (Center for quality testing and diagnostics of materials) workplace at Faculty of Industrial Technologies in Puchov.

3 Results and discussions

3.1 Dynamic mechanical analysis (DMA)

Fig. 1. shows the scan temperature data at frequency of 10 Hz for transparent and red LDPE. There is a decrease in storage modulus at approximately – 46 °C for red and at approximately – 30 °C for transparent sample which can be attributed to the β-relaxation or glass transition temperature (Tg). The loss modulus of red LDPE shows one well-defined transition and two small transitions while at transparent LDPE, all three transitions are well evident. The first peak of red sample at -29 °C may be attributed to the β-relaxation. The second and the third small peaks at approximately 25 °C and 49 °C may correspond with the α-relaxation, but it can also correspond with the melting of the small crystallites.

The possible β-relaxation of transparent sample occurs at approximately 0 °C. The second and the third peaks associated with the α-relaxation and the melting of small crystallites are at approximately 28 °C and at approximately 55 °C. There is increase in tan δ for both samples with peak of red LDPE at approximately 56 °C associated with α-relaxation.
The scan temperature data at frequency of 10 Hz for transparent and yellow PP are shown in Fig. 2. There is a decrease in storage modulus at approximately – 18 °C for yellow and at approximately – 7 °C for transparent PP which corresponds with glass transition temperature of materials. The loss modulus shows two temperature peaks for both samples at approximately - 41 °C and 4 °C for transparent sample and at approximately – 42 °C and -5 °C for yellow sample. These peaks are associated with the α- and β-transitions of polypropylene. The tan δ also shown two peaks associated with these transitions. The peaks of transparent PP are at approximately - 42 °C and 10 °C for transparent sample and at approximately - 43 °C and 2 °C for yellow sample.

The scan temperature data of pink and green PET at frequency of 10 Hz are shown in Fig. 3. The storage modulus of both samples shows well-defined peaks which correspond with glass transition temperature of PET. The peak for pink sample is determined at approximately 73 °C, while for green sample, it is at approximately 94 °C. The glass transition temperature can be also determined from loss modulus and from tan δ. From loss modulus, the T_g for pink sample is at approximately 93 °C and for green sample, it is approximately 103 °C, whereas from tan δ, the T_g for pink sample is at approximately 105 °C and for green sample, it is at approximately 117 °C.
3.2 Differential scanning calorimetry (DSC)

Differential scanning calorimetry was used for detecting the glass transition temperature and melting point of investigated materials. Fig. 4. presents DSC measurements of red and transparent LDPE samples. As can be seen from the graph, the melting peak of red LDPE is less sharp and wider in comparison with transparent LDPE. The melting point of red sample is at approximately 116°C. The second peak is at approximately 164°C. This peak is most likely result of the residual reactants from LDPE synthesis or contamination before analysis. Furthermore, the melting point of transparent LDPE is at approximately 121°C.

DSC measurements of yellow and transparent PP samples are presented in Fig. 5. The melting peak of yellow PP is sharper and narrower in comparison with the transparent one. The melting point of yellow sample is at approximately 173°C while the melting point of transparent sample is at approximately 171°C.
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Fig. 4. DSC measurement of LDPE samples

DSC measurements of yellow and transparent PP samples are presented in Fig. 5. The melting peak of yellow PP is sharper and narrower in comparison with the transparent one. The melting point of yellow sample is at approximately 173°C while the melting point of transparent sample is at approximately 171°C.

Fig. 5. DSC measurement of PP samples

Fig. 6. shows DSC measurements of pink and green PET samples. In contrast to the samples described above, the glass transition temperature is shown in the graph. The glass transition peak of pink and green sample is at approximately 104 °C and at approximately 98 °C, respectively. The melting point of pink sample is at approximately 151°C while the melting point of green sample is at approximately 152°C.

Fig. 6. DSC measurement of PET samples

3.3 Fourier transform infrared spectroscopy (FTIR)

The spectra of transparent and red LDPE samples obtained in absorbance using ATR technique is shown in Fig. 7. As can be seen from data, LDPE samples are very similar, because there are only differences in peak intensity. Characteristic peaks at around 3000 - 2800 cm⁻¹ correspond with methylene group symmetric stretching νₛ(CH₂) and asymmetric stretching νₘ(CH₂). Peak at around 1500 – 1400 cm⁻¹ can be associated with CH₂ due to scissoring. The rocking of methylene group is presented by peaks between 750 and 700 cm⁻¹.
Fig. 7. Measured spectra of LDPE samples

Fig. 8. presents measured spectra of transparent and yellow PP samples obtained in absorbance using ATR technique. From the FTIR spectra scan be seen, that there is similarity between two samples. They are different in 800 – 400 cm\(^{-1}\) IR spectra region. At around 3000 - 2800 cm\(^{-1}\), polypropylene spectral areas present absorption bands which correspond with asymmetric and symmetric stretching vibration of methylene and methyl groups. Different peaks have been also observed in the 1500 - 1350 cm\(^{-1}\) spectral area, where absorption bands can be associated with scissoring vibration of the methylene group (at around 1470 cm\(^{-1}\)) and to symmetric deformation of the methylene group (at around 1380 cm\(^{-1}\)).

Fig. 8. Measured spectra of PP samples

Measured spectra of pink and green PET samples obtained in absorbance using ATR technique are presented in Fig. 8. There is no significant difference between the two PET samples compared with the FTIR spectra scan, because there are differences only in spectra intensity. The weak peak at around 2950 cm\(^{-1}\) represents C-H asymmetric stretching. Spectra of PET also confirm a carbonyl group along with aromatic ring appear as the second strongest peak at around 1720 cm\(^{-1}\). Two peaks at around 1410 cm\(^{-1}\) and 1340 cm\(^{-1}\) represent the deformation of C-H alkane and peak at around 1510 cm\(^{-1}\) belongs to aromatic C-C stretching. The asymmetric C-C-O stretching involving the carbon in aromatic ring appears at around 1230 cm\(^{-1}\) as a strong peak. The strongest peak at around 720 cm\(^{-1}\) is associated with aromatic C-H wagging. The peak at approximately 870 cm\(^{-1}\) corresponds with aromatic C-H bending. The O-C-C asymmetric stretching is at around 1105 cm\(^{-1}\).
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Table 3. EDX analysis of PET samples

| Element | PET pink Contents (weight %) | PET green Contents (weight %) |
|---------|-----------------------------|-----------------------------|
| Sb      | 0.017                       | S                           | 0.019                       |
| S       | 0.016                       | Ca                          | 0.004                       |
| P       | 0.011                       | K                           | 0.002                       |
| Ca      | 0.005                       | Co                          | 0.001                       |
| K       | 0.001                       | Cu                          | 0.001                       |
| Cu      | 0.001                       | Fe                          | 0.001                       |

Conclusions

The measurements were carried out for six samples of thermoplastics, namely LDPE (red and transparent), PP (yellow and transparent) and PET (pink and green). Using DMA and DSC analysis, all the samples were measured and from the resulting graphs, the glass transition temperature and the melting point of each polymer was determined. There are several results of glass transition temperature, because it can be derived from the loss modulus, the storage modulus and the tan δ. It is because each of these results has its own physical essence.

Using the ATR-FTIR spectrometry, the spectra of the individual pairs of polymers were compared. Based on the results obtained in this work, it can be concluded that the ATR-FTIR spectrometry is useful tool for examination of the chemical structure of thermoplastics.

Furthermore, the EDX analysis revealed the presence of some elements and heavy metals in tested polymers.

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Table 3. EDX analysis of PET samples

| Element | Contents (weight %) |
|---------|---------------------|
| Sb      | 0.017               |
| S       | 0.016               |
| Ca      | 0.004               |
| P       | 0.011               |
| K       | 0.002               |
| Cu      | 0.001               |
| Fe      | 0.001               |

Conclusions

The measurements were carried out for six samples of thermoplastics, namely LDPE (red and transparent), PP (yellow and transparent) and PET (pink and green). Using DMA and DSC analysis, all the samples were measured and from the resulting graphs, the glass transition temperature and the melting point of each polymer was determined. There are several results of glass transition temperature, because it can be derived from the loss modulus, the storage modulus and the $\tan \delta$. It is because each of these results has its own physical essence.

Using the ATR-FTIR spectrometry, the spectra of the individual pairs of polymers were compared. Based on the results obtained in this work, it can be concluded that the ATR-FTIR spectrometry is a useful tool for examination of the chemical structure of thermoplastics.

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