Nanohardness of electron beam irradiated polyamide 11

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Abstract — The submitted paper presents the assessment of mechanical properties of the surface layer of modified polyamide 11. Hard surface layer was made using the technology of radiation crosslinking, which enables modification of polymer materials and hence the change of their end-use properties. The process of mechanical stress is applied by nanohardness test. The surface layer of polymer material such as polyamide 11 is modified by β – radiation. After the polyamide 11 is subjected to radiation, changes of the surface layer at applied load are observed. Material properties of the created surface layer are measured by nanohardness test using the DSI method (Depth Sensing Indentation).

Keywords — Crosslinking, irradiated, nano-hardness, PA 11.

I. INTRODUCTION

Polyamide belongs to the group of synthetic thermoplastics polymers. This group consists of aliphatic polyamides and aromatic polyamides. Family of aliphatic polyamides is well known as Nylon. The Nylon is one of used the most polymers.

As Polymers belong to constructive materials which find use at the most industry branches. The advantage is a low weight together with the excellent mechanical properties, very good chemical resistance and other properties, which assign them for various applications. Disadvantage is mainly low temperature stability which significantly reduces usage of these polymers.

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Every properties improvement especially temperature stability helps to increase application possibilities. In addition, properties modification of standard polymers, which are relatively cheap products, gives them advantage for another usage. One of the possibilities of polymers improvement is their radiation cross-linking.

The irradiation cross-linking of thermoplastic materials via electron beam or cobalt 60 (gamma rays) is performed separately, after processing. Generally, ionizing radiation includes accelerated electrons, gamma rays and X-rays.1

Radiation processing with an electron beam offers several distinct advantages when compared with other radiation sources, particularly γ-rays and x-rays. The process is very fast, clean and can be controlled with much precision. There is no permanent radioactivity since the machine can be switched off. In contrast to γ-rays and x-rays, the electron beam can be steered relatively easily, thus allowing irradiation of a variety of physical shapes. The electron beam radiation process is practically free of waste products and therefore is no serious environmental hazard. The main difference between beta and gamma rays is in their different abilities to penetrate the irradiated material. Gamma rays have a high penetration capacity. The penetration capacity of electron rays depends on the energy of the accelerated electrons. Due to electron accelerators, the required dosage can be applied within seconds, whereas several hours are required in the gamma radiation plant. (Fig. 1). [1,2]

![Fig. 1 Design of gamma rays (a) and electron rays (b), 1 – Penetration depth of an electron, 2 – Primary electron, 3 – Secondary electron, 4 – Irradiated material, 5 – Encapsulated Co – 60 Radiation source, 6 – Gamma rays [2]](image)

Beta and gamma rays can be used for the irradiation of polyolefines, polyesters, halogen polymers and polyamides from the thermoplastics group, elastomers and thermoplastic elastomers. Some of them need the addition of a cross-linking agent. [1,6,7,8]
Radiation cross-linking usually improves strength, reduces creep, contributes to chemical resistance improvement and in many cases improves tribological properties. Effect of radiation cross-linking significantly improves temperature stability. Because of that, materials which belong to group of standard polymers can be used in applications which would be in term of temperature stability intended only to constructive thermoplastic polymers.

II. EXPERIMENTAL

For this experiment polyamide 11 (PA11) PTS – CREAMID - 11T * M600/13 transparent; PTS Plastics Technologie Service, Germany was used. The material already contained a special cross-linking agent TAIC - triallylisocyanurate (5 volume %), which should enable subsequent cross-linking by ionizing β- radiation. The prepared specimens were irradiated with doses of 33, 66 and 99 kGy at BGS Beta-Gamma Service GmbH & Co. KG, Germany [1-4].

The samples were made using the injection molding technology on the injection moulding machine ArburgAllrounder 420C. Processing temperature 280–310 °C, mold temperature 60 °C, injection pressure 80 MPa, injection rate 50 mm/s.

Instrumented nanohardness tests were done using a Nanoindentation Tester (NHT2) – Opx/Cpx, CSM Instruments (Switzerland) according to the CSN EN ISO 6507-1. Load and unload speed was 100 mN/min. After a holding time of 90 s at maximum load 50 mN the specimens were unloaded. The indentation hardness HIT was calculated as maximum load to the projected area of the hardness impression according to:

\[
H_{IT} = \frac{F_{\text{max}}}{A_p} \quad \text{with} \quad h_c = h_{\text{max}} - \varepsilon \frac{F_{\text{max}}}{S}
\]

(1)

Where \( h_{\text{max}} \) is the indentation depth at \( F_{\text{max}} \), \( h_c \) is contact depth. In this study the Oliver and Pharr method was used calculate the initial stiffness (S), contact depth (\( h_c \)). The specimens were glued on metallic sample holders.

The indentation modulus is calculated from the Plane Strain modulus using an estimated sample Poisson’s ratio:

\[
E_{IT} = E^* \cdot (1 - v_i^2)
\]

(2)

The deduced modulus is calculated from the following equation:

\[
E^* = \frac{\sqrt{\pi \cdot S}}{2 \cdot \beta \cdot \sqrt{A_p(h_c)}}
\]

(3)

The Plane Strain Modulus \( E^* \) is calculated from the following equation:

\[
E^* = \frac{1}{\frac{1}{E_i} - \frac{1 - v_i^2}{E_i}}
\]

(4)

Where \( E_i \) is the Elastic modulus of the indenter, \( v_i \) is the reduced modulus of the indentation contact, \( v_i \) is the Poisson’s ratio of the indenter. [8] [12] [41].

Determination of indentation creep \( C_{IT} \):

\[
C_{IT} = \frac{h_2 - h_1}{h_1} \cdot 100
\]

(5)

Where \( h_1 \) is representing the indentation depth at time \( t_1 \) of reaching the test force (which is kept constant), \( h_2 \) is the indentation depth at time \( t_2 \) of holding the constant test force (Fig. 2) [1] [7] [15].

![Fig. 2 Illustration of indentation creep parameters](image)

Fig. 2 Illustration of indentation creep parameters

Elastic part of the indentation work \( \eta_{IT} \) (Fig. 3):

\[
\eta_{IT} = \frac{W_{\text{plast}}}{W_{\text{total}}} \cdot 100 \quad \text{with} \quad W_{\text{total}} = W_{\text{elast}} + W_{\text{plast}}
\]

(6)

Plastic part \( W_{\text{plast}} / W_{\text{total}} \) follows as 100% - \( \eta_{IT} \)

(7)

![Fig. 3 Illustration of coefficient of back deformation](image)

Fig. 3 Illustration of coefficient of back deformation

III. RESULTS AND DISCUSSION

For Instrumented nanohardness teset was used three different loads

A. Indentation load 10 mN

The values measured during the nanohardness test showed that the lowest values of indentation hardness were found for
the non-irradiated PA11. On the contrary, the highest values of indentation hardness were obtained for PA11 irradiated by a dose of 99 kGy (by 26% higher in comparison with the nonirradiated PA11), as can be seen at Fig. 4.

![Fig. 4 Hardness HIT of PA11 vs. irradiation doses](image)

Higher radiation dose does not influence significantly the nanohardness value. An indentation hardness increase of the surface layer is caused by irradiation cross-linking of the tested specimen.

According to the results of measurements of nanohardness, it was found that the highest values of indentation modulus of elasticity were achieved at the PA11 irradiated with dose of 33 kGy (by 10% higher than compared with non-irradiated PA11). On the contrary, the lowest values of the indentation modulus of elasticity were found for nonirradiated PA11 as is seen at Fig. 5.

![Fig. 5 Elastic modulus EIT of PA11 vs. irradiation doses](image)

The lowest values of hardness Vickers were found for the non-irradiated PA11. On the contrary, the highest values of hardness Vickers were obtained for PA11 irradiated by a dose of 99 kGy (by 26% higher in comparison with the nonirradiated PA11), as can be seen at Fig. 6.

![Fig. 6 Hardness Vickers of PA11 vs. irradiation doses](image)

Other important material parameters obtained during the nanohardness test were elastic and plastic deformation work. From the elastic deformation work we determine the reaction of material to applied (multiaxial) load with reversible deformation. The plastic part of the deformation work $W_{pl}$ defines toughness of the tested material (surface layer) and its resistance to plastic deformation (Fig. 7).

The greatest values of elastic deformation work were obtained for PA11 irradiated with dose of 99 kGy. The greatest values of plastic and elastic deformation work were obtained for non-irradiated PA11. The lowest values of elastic deformation work were obtained for PA11 irradiated with dose of 33 kGy. The lowest values of plastic deformation work were obtained for PA11 irradiated with dose of 99 kGy. Radiation of specimens caused lower values of plastic and higher values of elastic deformation work which is apparent in Fig. 7.

![Fig. 7 Elastic and plastic deformation work of PA11 vs. irradiation doses](image)

The greatest values of indentation creep were obtained for PA11 irradiated with dose of 66 kGy. The lowest values of indentation creep were obtained for non-irradiated PA11.
Radiation of specimens caused increase of indentation creep and subsequent decrease of indentation creep which is apparent in Fig. 8.

![Graph showing Indentation Creep of PA11 vs. irradiation doses](image)

**Fig. 8 Indentation Creep of PA11 vs. irradiation doses**

The lowest values of back deformation coefficient $n_{IT}$ were found for the PA11 irradiated with dose of 66 kGy. On the contrary, the highest values of back deformation coefficient $n_{IT}$ were obtained for PA11 irradiated by a dose of 99 kGy (by 12% higher in comparison with the PA11 irradiated with dose of 66 kGy), as can be seen at Fig. 9.

![Graph showing Coefficient of back deformation $n_{IT}$ vs. irradiation doses](image)

**Fig. 9 Coefficient of back deformation $n_{IT}$ vs. irradiation doses**

**B. Indentation load 50 mN**

The values measured during the nanohardness test showed that the lowest values of indentation hardness were found for the non-irradiated PA11.

On the contrary, the highest values of indentation hardness were obtained for PA11 irradiated by a dose of 99 kGy (by 18% higher in comparison with the nonirradiated PA11), as can be seen at Fig. 10.

![Graph showing Hardness HIT of PA11 vs. irradiation doses](image)

**Fig. 10 Hardness HIT of PA11 vs. irradiation doses**

Higher radiation dose does not influence significantly the nanohardness value. An indentation hardness increase of the surface layer is caused by irradiation cross-linking of the tested specimen.

According to the results of measurements of nanohardness, it was found that the highest values of indentation modulus of elasticity were achieved at the PA11 irradiated with dose of 33 kGy (by 8% higher than compared with non-irradiated PA11). On the contrary, the lowest values of the indentation modulus of elasticity were found for nonirradiated PA11 as is seen at Fig. 11.

![Graph showing Elastic modulus EIT of PA11 vs. irradiation doses](image)

**Fig. 11 Elastic modulus EIT of PA11 vs. irradiation doses**

The lowest values of hardness Vickers were found for the non-irradiated PA11. On the contrary, the highest values of hardness Vickers were obtained for PA11 irradiated by a dose of 99 kGy (by 26% higher in comparison with the nonirradiated PA11), as can be seen at Fig. 12.
The greatest values of plastic deformation work were obtained for PA11 irradiated with dose of 66 kGy. The greatest values of elastic and elastic deformation work were obtained for PA11 irradiated with dose of 99 kGy. The lowest values of elastic deformation work were obtained for PA11 irradiated with dose of 66 kGy. The lowest values of plastic deformation work were obtained for PA11 irradiated with dose of 99 kGy. Radiation of specimens caused lower values of plastic a higher values of elastic deformation work which is apparent in Fig. 13.

The greatest values of indentation creep were obtained for PA11 irradiated with dose of 33 kGy. The lowest values of indentation creep were obtained for non-irradiated PA11. Radiation of specimens caused increase of indentation creep and subsequent decrease of indentation creep which is apparent in Fig. 14.

The lowest values of back deformation coefficient $n_{IT}$ were found for the PA11 irradiated with dose of 66 kGy. On the contrary, the highest values of back deformation coefficient $n_{IT}$ were obtained for PA11 irradiated by a dose of 99 kGy (by 9% higher in comparison with the PA11 irradiated with dose of 66 kGy), as can be seen at Fig. 15.

The values measured during the nanohardness test showed that the lowest values of indentation hardness were found for the non-irradiated PA11. On the contrary, the highest values of indentation hardness were obtained for PA11 irradiated by a dose of 33 kGy (by 10% higher in comparison with the nonirradiated PA11), as can be seen at Fig. 16.
Higher radiation dose does not influence significantly the nanohardness value. An indentation hardness increase of the surface layer is caused by irradiation cross-linking of the tested specimen.

According to the results of measurements of nanohardness, it was found that the highest values of indentation modulus of elasticity were achieved at the PA11 irradiated with dose of 33 kGy (by 7% higher than compared with non-irradiated PA11). On the contrary, the lowest values of the indentation modulus of elasticity were found for nonirradiated PA11 as is seen at Fig. 17.

The lowest values of hardness Vickers were found for the non-irradiated PA11. On the contrary, the highest values of hardness Vickers were obtained for PA11 irradiated by a dose of 99 kGy (by 11% higher in comparison with the nonirradiated PA11), as can be seen at Fig. 18.

The greatest values of elastic deformation work were obtained for PA11 irradiated with dose of 99 kGy. The greatest values of plastic and elastic deformation work were obtained for non-irradiated PA11. The lowest values of elastic deformation work were obtained for non-irradiated PA11. The lowest values of plastic deformation work were obtained for PA11 irradiated with dose of 33 kGy. Radiation of specimens caused lower values of plastic a higher values of elastic deformation work which is apparent in Fig. 19.

The greatest values of indentation creep were obtained for PA11 irradiated with dose of 33 kGy. The lowest values of indentation creep were obtained for non-irradiated PA11. Radiation of specimens caused increase of indentation creep and subsequent decrease of indentation creep which is apparent in Fig. 20.
The lowest values of back deformation coefficient $n_{\text{IT}}$ were found for the PA11 irradiated with dose of 66 kGy. On the contrary, the highest values of back deformation coefficient $n_{\text{IT}}$ were obtained for PA11 irradiated by a dose of 99 kGy (by 9% higher in comparison with the PA11 irradiated with dose of 66 kGy), as can be seen at Fig. 21.

IV. CONCLUSION
For measurement with load of 10mN we obtained lowest indentation depth for PA11 irradiated with dose of 99 kGy. The greatest values were obtained for non-irradiated PA11, as can be seen at Fig. 22. For measurement with load of 50mN (Fig. 23) we obtained similar results as can be seen at Fig. 22.

The properties of surface layer of polyamide 11 modified by beta radiation improved significantly. The nanohardness values increased about 20%. Stiffness of surface layer increased significantly by 18% as a result of radiation. Changes of behavior in the surface layer were confirmed by final values of plastic deformation work whose values decreased in correlation with the increasing radiation dose. Elastic deformation was increasing with radiation dose. The highest values of micromechanical properties were reached at
radiation dose of 99 kGy. The results of nanomechanical properties of surface layer of modified polyamide 11 show that it can be used in more difficult applications in some industrial fields, in particular where there are high requirements for strength, stiffness and hardness of surface layer which appears to be the most suitable area of application. The resistance of surface layer to wear makes its use suitable for the production of gears, friction parts of machinery and as alternative to some metal materials. Thanks to its low weight polyamide 11 modified by beta radiation is a suitable alternative to commonly used materials in the car and electrical industry.

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