Mechanical Characterization of two blends PP+PA6+EPDM. Part 2. Results of Tensile Tests

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Abstract: This paper presents the values of several mechanical characteristics for two blends: H (60% PP + 12% PA6 + 8% EPDM + 20% Polybond 3200) and G (20% PP + 42% PA6 + 28% EPDM + 10% Polybond 3200) (%wt), comparing them to those of PP (polypropylene) and PA6 (polyamide 6). The adding of EPDM (ethylene propylene diene monomer rubber) and Polybond 3200 help reducing the disadvantages of simple blends made of PP+PA6, when the PA6 concentration allows for having a PA6 matrix with droplets of PP. SEM analysis helps for understanding the performance of material G as compared to the neat polymers. EDX analysis proved that there was a matrix inversion, material H having a PP matrix and material G a matrix made of PA6 with droplets of PP. Strain at break for PP and material H were proved to be insensitive to test speed, but materials G and PA6 had large value for strain at break and energy at break for v = 10 mm/min. Taking at basis the values for polyamide 6 (PA6), material G has greater values for energy at break: with 97.8% for v = 10 mm/min, with 29.5% for v = 250 mm/min and with 98% for 1000 mm/min, without exhibiting the micro and macro cavitation of PA6. This means that the recipe, the technology and the mechanical characteristics make material G a potential candidate for applications where a low and moderate impact resistant material is required.

Keywords: polyamide 6, polypropylene, blend, EPDM, mechanical characteristic, energy at break

1. Introduction

The polymer blend segment is growing three times faster than the entire plastics industry. Polymer blending is the most suitable method to produce materials that meet requirements of the market. The aspiration is to produce a blend that can be treated like any other polymer; therefore, the processing of the blend must be consistent with that of simple polymers, but with better performance [1]. The main objectives of polymer blending are [2]:

- improving processing and productivity efficiency,
- producing materials with better mechanical performance, lighter parts,
- mixing of scrap reduction,
- mixing reduces the number of assortments to be stored and manufactured.

The method of compatibilizing immiscible polymers (like PP + PA) may include several alternatives [2-6]: the addition of a coating-type copolymer, also called impact modifier because it improves impact qualities, the addition of the third component to the polymer blend, miscible with both polymers, the addition of a reactive element that develops a local miscibility, additives having mechanical and/or chemical influence in the blend, etc.

2. Blends based on PP and PA6

DSM, Atochem and Mitsubishi introduced PA + PP blends due to their lower water absorption than polyamides, dimensional stability (affected by PP concentration), low
density, low permeability of liquids and vapors and moderate impact strength, induced by PA and good resistance to alcohols [7].

PP (polypropylene) is a cheap general-purpose polymer, with high purity and chemical stability, but low impact resistance [8]. It has a crystallinity degree of 50-60% [9]. The set of properties includes low density, high melting point ($T_m = 166 \, ^\circ C$) and good processability. The low temperature impact resistance of unmodified PP and the Izod impact resistance give lower values than those obtained for PE. Fragility is related to spherulitic morphology and the tendency of cracking by spinning PP, which increases in the presence of stress concentrators, especially at negative temperatures [4], [10]. PP has a high wear resistance, does not crack in corrosive environments and it is resistant to ultraviolet rays and oxygen. When stabilizers are added, this performance is improved. When PP is in contact with organic solvents, only a slight swelling occurs. One of the main problems faced by PP components is the low impact resistance. It follows that the blend of PP with other polymers and/or elastomers would improve this property, while maintaining to some extent the qualities of the polymer [11]. The softening of PP begins at 140 °C, the melting temperature being in the range from 164 °C to 170 °C, and from -8 °C down the polymer becomes brittle. Trademarks usually contain antioxidant stabilizers and other additives [12].

All types of polyamides (PA6; PA6.6; PA6.10; PA6.12; PA11, PA12) have many applications because they can be processed much easier as compared to other polymers, have good wear resistance and high melting temperatures, but, as a disadvantage, they have a relatively high cost and absorb water very easily [13].

There are three methods to achieve PP-PA compatibility: physical compatibility by high stress field in processing, heat treatment, irradiation, etc. chemical compatibilization, by incorporating a compatibilizing agent, usually either a copolymer or multipolymer and reactive compatibility during processing by extrusion or injection molding [14], [15].

Blends PP and PA6 have been tested for improving mechanical properties, where PP ensures good processability and moisture insensitivity, while PA6 contributes with mechanical and thermal properties. Researchers have described nanocomposites based on unique polymer matrix [16], [17]. A new approach is represented by composites based on polymer blends of two or more polymeric materials. Nanocomposites based on PA6/PP blends have been analyzed [5]. Blends PP+PA have immiscible components, usually with highly asymmetrical compositions (90/10, 80/20 or 70/30), insular morphologies, with large droplets and a lack of adhesion between phases, as a result of high values of interfacial tensions between immiscible phases [18], [19]. A simple FE model [20] pointed out that a bonded droplet behaves very different as concern the stress and strain distributions, when comparing to a matrix that could slide along the dispersed phases. Immiscible blends have poor mechanical properties if no compatibilizing agents are added. In the recent decades, positive results have been reported for these blends, but with different compatibility agents [21-26].

For PP + PA blends, the researchers investigated the influence of adding elastomers: styrene-ethylene-butylene-styrene maleate (SEBS) [27-29], ethylene-polypropylene-diene copolymer (EPDM) [26], [30-34]; acryl-butadiene-styrene elastomer (ABS), copolymer block styrene-ethylene-butylene-styrene maleate (SEBS) [21], [35].

Impact resistance is usually improved to the detriment of strength and dimensional stability. The rigidity of polyamides with the addition of elastomer can be preserved to some extent by inorganic additions (glass micro spheres [36], short glass fibers [37], minerals and clays [37], carbon nanotubes [38], [39] and organic (short aramid fibers [40]). Chow et al. [19] and Wahit et al. [41] also reported that the ductility and resilience of PA6 + PP + clay blends were improved by the addition of an elastomer.

EPDM was used as a compatibilizer in PP/LDPE blends and their mechanical properties and Izod impact resistance improved due to its addition, especially for LDPE-rich ones and those with a high EPDM content. The compatibilizer did not have an effect of improving the tensile strength, except for the PP/LDPE blend (80/20). Interfacial adhesion was also improved [42].
Here are some examples of elastomers added to PA and PA + PP blends: 
PA6/maleated methylene diylene ethylene propylene monomer (mEPDM)/nano clay [43], 
PA6/metallocene ethylene-propylene-diene terpolymer (EPDM)/sepiolite [44]; PA6/ 
ethylene propylene propylene diene methylene (EPDM-g-MA)/organic clay [45], PA6/ 
ethylene-polypropylene-diene copolymer metallocene + ethylene-polypropylene- 
diene copolymer/ethylene-polypropylene-diene copolymer maleate (EPDM-g-MA) + clay [46].

In polymeric blends, cavitation occurs at micro and/or macro levels. The cavities are 
generated in the blend volume, either during cooling after injection of the melt into the 
mold or due to stresses induced under load [21], [47], [48].

Bai et al. [22] developed a class of blends, composed of three phases: PP, PA6 and 
polyethylene-octene elastomer (POE), crosslinked with maleic anhydride. The mass 
fraction of PA6 was adjusted from 0% to 40%, in steps of 10%, and the mass fraction POE 
was kept at half that of PA6. Their morphology was mainly from PA6 particles dispersed 
in the PP matrix. The POE modifier was observed as a thin interface (less than 100 nm 
 thickness) at the PP/PA6 interface, and as isolated, but few, particles. The modulus of elasticity 
and the tensile strength at break are almost constant for PP and elaborated blends, but 
Izod impact resistance increases greatly with the content of the alloying components. This 
effect is due to POE reducing the interface cavitation, to higher resistance of PA6 drops to 
crack propagation and to the larger plastic deformation.

PA+PP blends were considered interesting materials because both components are 
relatively cheap, with advantageous properties and can be processed by melt mixing. The 
compatibility of polymers in blends can be achieved by the addition of reticulating copol-
ymer, segments that have physical or chemical affinity with immiscible homopolymers. 
In this study, the authors used PP reticulated with maleic anhydride (PP-g-MA). Graphite 
and graphene oxide were added in the blends. The family of blends had the objective of 
 railway components (seals) that simultaneously meet impact strength, wear durability 
and dimensional stability [25].

To reduce the sensitivity of polyamide to crack initiation and its development on low 
velocity impact, the polymer is mixed with 10-25% elastomer (including EDPM that was 
used for these two original recipes) [50].

The mechanical properties, phase morphology and breaking behavior of the ternary 
blends PP/PA6/(EPDM:EPDM-g-MA) (70/15/15), with different concentrations of 
EPDM:EPDM-g-MA, were presented in a recent study [32]. With the addition of EPDM- 
g-MA, the strain at break increases. PP has a semi-ductile behavior in tensile tests and PA6 
does a ductile fracture as compared to PP, with a higher tensile strain and higher tensile 
strength.

3. Materials and Methods

The properties of polymer blends depend on their morphology, the quality of the 
interfacial adhesion, the surface tension and the package of properties with which each 
component contributes [51-55]. As in the case of PA + PP blends, a polymer blend may 
have in its composition several stabilizing agents and nanoparticles that can improve the 
interfacial adhesion and morphological stability of immiscible polymer blends [25].

PA6 brings in a blend its mechanical and thermal properties and PP brings processa-
bility, water insensitivity and dimensional stability [21], [23], [25].

Polybond 3200 as a miscibility agent is a chemically modified polypropylene. Physi-
cal properties of blends could be modified by using lower levels of addition; it is a chem-
ical coupling agent for glass fibers and PP-reinforced mineral filler, which improves the 
physical and thermal properties; it is a compatibilizer for polypropylene/polyamide and 
polypropylene blends for improving processing and mechanical properties.

Chow et al. [19] used MAH-g-PP (Polybond 3200), having 1.2 wt% maleic anhydride 
(MA). Tensile tests were performed at a test rate of 50 mm/min. The modulus of elasticity 
was less sensitive to the addition of MAH-g-PP in blends with PA6, but considerably in-
creased the toughness of a composite, largely due to clay reinforcement.
EPDM (ethylene-propylene-diene maleate monomer) is a synthetic rubber monolayer membrane, sometimes mixed with carbon black, oils, curing agents and other additives, chemically stable, resistant to ultraviolet rays and ozone, practically unlimited. Elongation of over 300% gives EPDM the advantage of withstanding mechanical stress that other materials cannot. At -40 °C, EPDM keeps its properties, while thermoplastics lose their elasticity and become hard and brittle and at +150 °C, EPDM also keeps its properties, while thermoplastics liquefy. After repeated stretching, EPDM retains its properties, while thermoplastics become thin, elongated or breaks [56].

The recipes of the blends were established by agreement with the Monofil SA Săvinești (Romania), which also made the specimens by mold injection. The two polymeric blends consist of four elements, namely: PA6, PP, EPDM and Polybond 3200, in the concentrations shown in Table 1.

| Material | PA6 | PP | EPDM | Polybond 3200 |
|----------|-----|----|------|---------------|
| PP       | 100 | -  | -    | -             |
| H        | 12  | 60 | 8    | 20            |
| G        | 42  | 20 | 28   | 10            |
| PA6      | 100 | -  | -    | -             |

Figure 1a schematically shows the laboratory technology for obtaining the materials developed by the authors, with the involvement of the company Monofil SA Săvinești, Figure 1b presenting the injection molding equipment.

The technological flow consists of the pre-mixing of granules of PA6, PP and Polybond 3200 in a mixer (high speed mixer), with a capacity of 200 l, mixing speed 475/950 rpm, equipped with a heating system with electrical resistances of 11 kW and pneumatic discharge system. It is essential to pre-mix the components in a mixer, before introducing them into the extruder, because the raw materials have different densities. This methodology will guarantee a uniform dispersion of the components in the mass of molten polymer and, implicitly, an extremely low degree of agglomeration [26].

Blends of polymers and additives, resulting from the high-speed mixer, will be introduced at a temperature of 80-100 °C in a drying hopper. The dryer is equipped with an automatic system for loading and unloading, with a capacity of 1500 l and a working flow of 200 kg/h. The pre-mixtures in the drying hopper are automatically loaded into the primary dosing system. The primary dispenser has the following technical data: dosing flow = 150 kg/h, dosing speed = maximum 100 rpm, dosing system with double screw; feed hopper volume = 150 l.

The compounding of polymer and additives was performed on a granulating extruder type EC 52, with double screw (diameter 51.4 mm), with simultaneous rotation, injection pressure being 60 bar. The extruder is equipped with a cooling system of the zones using softened water jacket, respectively a hydraulic system for continuous filtration of the melt, equipped with a 1.5 kW motor and a maximum working pressure of 20 MPa [26]. The injection molding parameters are given in Table 2.
Figure 1. Technological flow chart for obtaining the specimens made of PA6, PP and the two blends

Table 2. Optimal compound processing parameters.

| No. | Technological parameters                              | UM | Values       |
|-----|--------------------------------------------------------|----|--------------|
| 1   | Working temperature on the 3 zones (from 9 available on the equipment): | °C | 130-150      |
|     | Zone I                                                 |    | 240-260      |
|     | Zone V                                                 |    | 220-230      |
| 2   | Screw diameter                                         | mm | 51.4         |
| 3   | Ratio L/D                                              |    | 40:1         |
| 4   | Maximum rotary screw speed                             | rpm| 600          |
| 5   | Modular screw structure, with 5 types of sections of different lengths | mm | 16-48        |
| 6   | Axial pressure                                         | kN | 4.5-5.5      |
| 7   | Melt pressure                                          | bar| 50-60        |
| 8   | Melt temperature                                       | °C | 90-100       |
| 9   | Supply current intensity                               | A  | 24           |
| 10  | Main engine power                                      | kW | 55           |

4. Tensile test results

For each material, 5 tests were performed (according to the standard SR EN ISO 527-2: 2012) for each material and each test speed, namely 10 mm/min, 250 mm/min and 1000 mm/min. The following parameters were analyzed: modulus of elasticity, tensile strength at break, strain at break, energy at break. Each characteristic was represented by the average values, the maximum and minimum values, as obtained from engineering stress-strain curves.

Following the use of INSTRON 2736-004 tensile test machine (INCAS, Bucharest) and its dedicated software, the true stress-strain graphs were represented in Figures 2.

According to the literature, the strain rate, $\dot{\varepsilon}$, can be defined approximately by the relation (1) and the values obtained are given in Table 3.1:

$$\dot{\varepsilon} = \frac{v}{L_0} \left[ s^{-1} \right]$$

where $v$ is the test rate, in mm/s and $L_0$ is the length of constant cross section, supposed to be deformed during the tensile test.

Table 3. Values of the strain rate
Table 1: Tabulated values of the strain rate for different test speeds.

| Test Speed [mm/min] | Strain Rate \( \dot{\epsilon} = \frac{v}{L_0} \) [s\(^{-1}\)] |
|---------------------|--------------------------------------------------|
| 10                  | 3.32 \times 10^{-3}                              |
| 250                 | 83.2 \times 10^{-3}                              |
| 1000                | 32.2 \times 10^{-3}                              |

Figure 2 presents the true stress-strain curves for the analyzed materials for all 5 tested specimens, till their break. The true stress and the true strain were calculated with the help of the following relations:

- True tensile stress
  \[ \sigma = \frac{F}{A} \]  

- Tensile stress at traction (engineering)
  \[ \sigma_{\text{eng}} = \frac{F}{A_0} \]  

- Strain rate in traction (engineering)
  \[ \dot{\epsilon}_{\text{eng}} = \frac{\Delta L}{L_0} \]  

where \( F \) is the force applied at time \( t \), \( A \) - cross area of the specimen at time \( t \), \( A_0 \) - initial cross area of the specimen (as measured), \( L_0 \) - initial length of specimen between marks, \( L \) - the length between marks, at moment \( t \).

If the engineering values are known, the relations used in this paper to calculate the true values are:

- True strain
  \[ \dot{\epsilon}_{\text{true}} = \ln (1 + \dot{\epsilon}_{\text{eng}}) \]  

- True tensile stress in traction
  \[ \sigma_{\text{true}} = \sigma_{\text{eng}} (1 + \dot{\epsilon}_{\text{true}}) \]  

The material PP (Fig. 2, first line), by its tendency of overlapping curves, can be considered a fairly predictable polymer, except for the strain at break, which varies quite a lot at the lowest test speed and much less at higher test speeds. The slower the load is applied to the test piece, the bigger the strain at break as compared to the strain at higher test speeds, where the strain at break is small in value. Strapasson et al. [57] performed tests at 5 mm/min and the results of mechanical characteristics for PP, injected at 170 °C, 180 °C, 190 °C and 200 °C, show the importance of the injection temperature: strain at break at 190 °C, decreases from 650% to 10%, indicating severe degradation. The stress-strain curves after sample injection at 170 °C are approximate to those obtained by the author for 10 mm/min, the range of strain at break values is smaller than that obtained at 5 mm/min (as it is normal), but the values of the elasticity modulus are close.

In material H (Fig. 2, second line), the lowest concentration of polyamide is found, namely 12% PA6. There is a tendency to maintain the shape of the stress-strain curves in the elastic zone, up to about 20-25 MPa, after which the curves are spread in the plastic domain, over a range of about 25 MPa, 40 MPa, respectively. One can note the break occurred at very different values of strain. Therefore, the material has a predictability on which the design engineer can rely only in the range of 0-20 MPa, after this value, the break takes place in a wide range of stress, from 25 MPa to 40 MPa. If parts made of this material are designed, it should not exceed 15-20 MPa, in the maximum values, in order to remain in the elastic zone. Thus, from the tests performed for material H, it is found the unpredictable behavior of this material after stress values of 25 MPa.

Material G has a concentration of 42%wt PA6 and the scattering of the results obtained for strain at break falls within quite wide ranges (Fig. 2, third line). It is possible to observe the overlap of the true stress-strain curves in the elastic domain and the
Figure 2. True stress-strain curves for: (a) PP and (b) PA6, for three test rates (10 mm/min, 250 mm/min and 1000 mm/min)

nonlinear elasto-plastic zone, then a slight distance appears in the area of plastic flow. On all curves, two domains can be observed, except for those obtained at speed $v = 10$ mm/min, where three domains appear, namely the first domain in the elastic zone (almost
linear slope where the upper value and inclination are slightly sensitive to the stress speed, then a zone with a yielding plateau (plastic deformation visibly more pronounced for lower test speeds, ie \( v = 10 \) mm/min), and finally the zone with a steeper slope. The very close behavior of the parts on demand is highly sought after by design engineers, being a defining selection criterion. For test speeds 250-1000 mm/min, the shape and characteristics of the curves are quite similar and the values of stress at break show a slight upward trend (see Fig. 5).

Figures 3 and 4 present the sets of 5 specimens tested for each material, for the selected three test speeds. Local neckings appear on the PA6 material (Fig. 3d, e and f), but not on the PP (Fig. 3a, b and c). At speed \( v = 10 \) mm/min, PP has an almost transverse breaking surface, with the exception of one of the 5 specimens with breaking fibers and a slightly pronounced neck.

In Fig. 3d (last line), from the set of 5 tested specimens made of PA6 material, at speed \( v = 10 \) mm/min, two specimens have a continuous flow over a longer interval until breaking, and the other three have a shorter one. The shape of the stress-strain curves up to rupture, for the other test speeds, also consists of two regions: the first almost linear, in the elastic zone framed in a narrow value band, up to about 30 MPa at \( v = 10 \) mm/min , and 35 ... 40 MPa for the other speeds. In the second region appears the area of the elasto-plastic curves that connect with the flow bearings. The yield level at \( v = 10 \) mm/min has ripples, which means that the material is a polymer formed of crystalline and amorphous micro-volumes where large local flows are highlighted.

Figure 3. Samples made of PP after been tested in traction: (a) 10 mm/min; (b) 250 mm/min; (c) 1000 mm/min and samples made of PA6 after been tested in traction: (d) 10 mm/min; (e) 250 mm/min; (f) 1000 mm/min.

Specimens made of material H (Fig. 4a, b and c) present brittle fracture, the aspect being similar to PP, this being explicable as the matrix of the blend H is made of PP.

In material G (Fig. 4d), at the test speed \( v = 10 \) mm/min there are bottlenecks on the test specimens and large elongations until breaking, this being characteristic of ductile materials and this makes the values of the tensile breaking energy have high values. However, the appearance of local necks can also be observed at the other test speeds, which may mean changes in the morphology and/or crystallinity of the polymer, possibly due to the force of stress.
Figure 4. Samples made of material H after been tested in traction: (a) 10 mm/min; (b) 250 mm/min; (c) 1000 mm/min and samples made of material G after been tested in traction: (d) 10 mm/min; (e) 250 mm/min; (f) 1000 mm/min

Average values of mechanical characteristics and their spread intervals for each tested material are given in Fig. 5.

For PP (Fig. 5, first column), the mechanical properties have a low sensitivity to the test speed and the spread intervals are small. The average strain at break for the other two test speeds varies between 8.6% and 9.9% as comparing to the average values. The values obtained for the strength at break fall within a narrow range, so the highest tensile test speed (1000 mm/min) causes the values for strength at break to increase with 59.14% for PP as compared to the value for $v = 10$ mm/min. In the case of the strain at break, the values are slightly sensitive to the test speed, the samples made of PP broke at a strain of 9 ... 13%. The graphs of energy at break and strain at break follow the same trend, namely, the values of the parameters decrease with increasing test speed, as reported in other research [23]. The modulus of elasticity is slightly influenced by the test speed and the obtained values increase with the increase of the latter, so that the lowest value for this parameter (1623 MPa) was obtained at the lowest test speed (10 mm/min), and for the other two speeds, the values increased by about 13% and 10%, respectively.

Material H (with a content of 12% PA6, Fig. 5, second column) presents the following conclusions regarding the influence of the test speed on the mechanical parameters:

- the modulus of elasticity seems to be less sensitive to the test speed, the increasing values over the test speed range being only 3.8% as compared to value at the lowest test speed,
- the strength at break is clearly dependent on test speed, slightly increasing, almost linear, with a small slope,
- the average values, as well as the maximum strain at break are between about 3.3% and 4.8%, therefore the dependence on the test speed is low, but the spread intervals are quite large,
- the values of the energy at break increase linearly with increasing test speed, so that the lowest value was obtained for $v = 10$ mm/min (2,6 J), after which the values increased by 35% for 250 mm/min and by 107, 7% at the highest test speed (1000 mm/min), while maintaining values lower than those obtained for PP; this is an atypical behavior.

For material H, the average values of the four discussed parameters (strength at break, strain at break, modulus of elasticity and energy at break) show an increasing trend with the test speed (from 10 mm/min to 1000 mm/min) and, yet the values are lower than those of simple polypropylene.
The variation of the values obtained for the breaking strength parameter is insignificant for material G (Fig. 5, third column), but still retains the upward trend with increasing test speed, so for the lowest test speed (10 mm/min) the lowest was obtained average value for this parameter (34.17 MPa) and highest (35.7 MPa) for the highest test speed (1000 mm/min). The capping of the average breaking stress value for a test speed range (around 34 MPa) is not characteristic of many materials [58], [59]. The same tendency is maintained by the modulus of elasticity, so for the lowest test speed, \(E = 1594\) MPa, for \(v = 250\) mm/min \(E = 1623\) MPa) and for \(v = 1000\) mm/min \(E = 1625\) MPa) the increase being only 1.94% compared to the lowest test speed, therefore the fact that this parameter is so little influenced by the test speed makes G look quite particular for a plastic material [60], even for a blend.

The other two parameters (strain at break and energy at break) show the same tendency of the values, namely the highest values (118.7 % and 178.4 J, respectively) were obtained for the lowest test speed (10 mm/min), the scattering ranges obtained are also large, after which the values of these parameters have decreased substantially, which makes material G behave atypically compared to other plastics, because normally the values of these parameters should be to decrease with increasing test speed [58], [61].
The decrease of the deformation at break, from the lowest speed (10 mm/min) to the speed \(v = 250\) mm/min is characteristic of polymers, but the behavior of this characteristic in the range 250 ... 1000 mm/min is atypical, the possible cause being the behavior the EPDM constituent and the cohesion between it, PP and PA6 [26].

For material G, the values of the strength at and elastic modulus show an increasing trend with the test speed (250 ... 1000 mm/min) and yet the values obtained are lower than those of plain polyamide. This means that the elaborated blends have intermediate values for this characteristic, which would confirm the rule of mixtures.

For PA6 (Fig. 5, fourth column), the values obtained for the strength at break fall within a narrow range but with a tendency to increase the values, so the highest tensile test speed (1000 mm/min) makes the values for the breaking strength increases by 20.1% for PA6 as compared to the value of the same parameter at \(v = 10\) mm/min. Some specimens of PA6 were broken at 24 ... 30% of its strain, but others were broken after about 90%, however the scattering interval of the latter is very long, possibly due to the gaps obtained during processing, such of material having a slightly unpredictable behavior.

The Young modulus depends on the test speed, the obtained values increase with the increase of the latter, so, at the lowest test speed (10 mm/min), the lowest value was obtained for this parameter (1444 MPa), and for the others two speeds, the values show insignificant differences, namely 1835 MPa (250 mm/min) and 1839 MPa (1000 mm/min).

Each characteristic is analyzed based on the following formula, written here only for strength at break:

\[
\frac{\sigma_r(1000) - \sigma_r(10)}{\sigma_r(10)} \cdot 100 \% \tag{3.2}
\]

where \(\sigma_r(10)\) is the value for strength at break at test speed \(v = 10\) mm/min and \(\sigma_r(1000)\) is the value for strength at break for the test speed \(v = 1000\) mm/min. The results of the comparative data are given in Table 4.

| Test | PP  | G   | H   |
|------|-----|-----|-----|
| \(\frac{E_r(1000) - E_r(10)}{E_r(10)} \cdot 100 \%\) | 10.32 | 1.97 | 3.82 |
| \(\frac{\sigma_r(1000) - \sigma_r(10)}{\sigma_r(10)} \cdot 100 \%\) | 59.25 | 4.51 | 27.74 |
| \(\frac{E_r(1000) - E_r(10)}{E_r(10)} \cdot 100 \%\) | -31.88 | -75.37 | 44.23 |
| \(\frac{E_r(1000) - E_r(10)}{E_r(10)} \cdot 100 \%\) | -16.10 | -73.29 | 107.97 |

The interface of the constituents influences the development of the continuous phase but also of the co-continuous morphologies. The morphology of the dispersed phase is the one that has a decisive role in establishing the mechanical properties of immiscible polymers. During shearing, the droplet size is a direct result of the fragmentation of the initial droplets and the coalescence process (merging/reuniting of several droplets).

Among the morphologies resulting from blends with immiscible constituents, co-continuous structures are more promising because they give a material in which the components intervene synergistically in establishing the set of properties [26].

The addition of small amounts of adhesives, elastomers can influence the morphology of immiscible blends, either by reducing the droplets of the dispersed phase, or by promoting the formation of co-continuous morphologies. These improve the breaking behavior, having an effect of increasing the strength at break and the energy at break. A stable co-continuous morphology balances or counterbalances the resistance that
would have been created in the absence of particles between the immiscible phases of the blend [62], [26].

5. Analysis of the blend morphologies

The break surfaces of specimens were coated with gold in order to increase the local electrical conductivity, the result being a better resolution of the SEM images.

In blends with PP matrix, the deformation results from a combination of amorphous hyperelastic phase and crystalline plasticity [21]. The PA6 phase is also capable of deforming plastically, but its yield strength falls within the plastic domain and has a much higher value than that of PP [24]. Consequently, in PP/PA6 blends, the isolated particles of PA6 show a lower deformation than the PP matrix, which leads to surface tension concentrations. In contrast, insulated POE nodules deform slightly due to the compatibility properties of the rubber.

The immiscibility of PP + PA6 blends can lead to a reduction of strain at break, this being also reported by other authors [17], [25], [35]. This behavior can be explained by a process of delamination, detachment of the phases from the cooling phase and then on loading, due to the reduced interfacial adhesion between the constituents [26].

Specimens made of PA6 had large central elongated ellipsoidal cavities that were obtained during processing and not under load. Pure polyamide is difficult to process by injection into the mold taking into account the parameters chosen in this paper [60]. Macro-scale gaps may occur due to inadequate processing parameters. Cavities also occur if the volumes on the surface of the injected elements solidify too quickly and the contraction is located inwards. Therefore, the cavities are associated with molds that are too cold to melt and specific recommendations of the injected polymer. Causes of cavities can be: insufficient melting volume in the machine, incorrect processing conditions (pressure, temperature, time and speed of injection and maintenance), wet materials, so their appearance can be avoided by redesigning the injection process. Specialists [63], [64], recommend: drying the constituents, increasing the melting temperature and more precise control over the processing areas, increase of the mold temperature, increase of the injection and maintenance pressure, increase of the injection time, increase of the supply in the melt distribution system, especially if the buffer zone is small, decrease of the supply area if the buffer zone is too big, relocation of the intake paths of melting, removal of thick-walled sections [26].

Figure 6 shows typical details by comparing the aspect of broken surfaces of PP and PA6 specimens, at the lowest test speed, with a significant difference of the break aspects, especially when comparing images at high magnification.
PA6 has a very ductile nature for low test rate (10 mm/min) (Fig. 6d, e and f), but becomes less ductile for higher rates (250 mm/min and 1000 mm/min).

Figure 7 shows typical details by comparison of the rupture of some polypropylene and polyamide specimens, at the test speed of 250 mm/min. The fibrillation process at the level of micro volumes is no longer accentuated at the test speed v = 250 mm/min because it can be seen that the material yields over the entire surface. In contrast, the PP material has a breaking surface specific to a thermoplastic polymer. At low test speed (v = 10 mm/min), the breaking surface of PP shows slightly pronounced local flows (the wavy surface on the right), which means that there are qualitatively different areas at the polymer level (average molecular and degree of mass crystallinity). At higher test speeds, cavities appeared due to the uniaxial stress, highlighting different flows of the polymer. At v = 1000 mm/min, these stress cavities are smaller and rarer, and the breaking surface has a brittle appearance.

Typical details of broken surfaces are given for PP (Fig. 8a) and PA6 (Fig. 8b) specimens, at the test speed of 1000 mm/min. At this test rate, the presence of fibrils may be rare, and they are thin, suggesting a sudden stretching of a more plastic micro-volume.
You can also see on the SEM images very small pores that appeared as a result of the band-type landslides on the broken surface.

Figure 8b shows the breaking of PA6 specimens at the test speed 1000 mm/min, but also the presence of processing cavities in some specimens, which negatively affects the performance of the material. From tested PA6 specimens, approximately 20% were with cavitation. Decohesion (detachment) of particles from the matrix is an important mechanism of deterioration in polymeric blends with ductile matrix (such as PA6) and weakly adherent particles (such as PP when constituting the dispersed phase). This mechanism also occurs in PP + PA + POE blends [22]. The improvement of the behavior of such blends can be done by increasing the adhesion of the intermediate phases between the matrix and the dispersed constituent, by adding compatibilizing agents in PA + PP (clays, fibers, polymer blocks or elastomers).

![SEM images of broken surface](image)

**Figure 8.** Details of broken surface for v = 1000 mm/min, at different magnification scales: (a) PP; (b) PA6

The influence of the compatibilizer concentration on the morphology of blends with PP + PA6 + elastomeric compatibilizer was also evaluated by Li and Xie [65] on blends (PP + PP-g-(MAH-co St)) + PA6 + SEBS (70+ 15 + 15), in which the component (PP + PP-g-(MAH-co St)) had concentrations from 0% to 70%.

The cavitation of isolated elastomer particles and elastomer intermediate layers are the main mechanisms of damage and volume expansion of polymer blends. As the cavitation in PA6 particles was found in the studies of Bai and G’Sell [21-24], but also in other authors [66-69] the deformation damage is controlled by the elastomer cavitation and the interfacial delamination (detachment) [22], [26].

The introduction of an elastomeric agent to increase the resilience in polymeric blends leads to an increase in impact resistance due to the decrease in stiffness and strength. On the contrary, if a compatibility is desired to increase the strength of a PA6 + elastomer blend, Kelnar et al. [70] found that the morphology of the dispersed phase must be small. TEM images showed the formation of hard shell-like particles (elastomer surrounded by clay micro-platelets), which improved the resilience of the elastomer. The result was a simultaneous increase in strength, resilience and dimensional stability.
The cavitation of isolated elastomer particles and elastomer intermediate layers are the main mechanisms of damage and volume expansion in polymer blends. As the cavitation in PA6 particles was found in the studies of Bai and G’Sell [22], [23], [24], but also in [71], [72], the deformation damage is controlled by the elastomer cavitation and the interfacial delamination (detachment) [26]. For low-content blends, intermediate POE layers are very thin and PA6 particles are almost spherical. The crack develops easily and leaves a smooth surface. In contrast, for blends with high-alloy content, the intermediate layer of POE is thick and PA6 particles are much elongated, resulting in a rough surface. Therefore, to improve the impact strength of polyamides, an impact strength modifier, EPDM was included in the blend (8% EPDM in material H and 28% EPDM in material G).

Figures 9 and 10 present SEM images with the magnification aspects of x100 and x2000, respectively, for the blends H (12% PA6) and G (40% PA6), from where the fragile break for material H is pointed out. For material G, the appearance micro-volumes of material strongly deformed can be observed; the break is initiated in small spots, with elongated and thinner shape towards the break point. This aspect may be considered a positive one because the material hardens locally and deforms more until break, also incorporating higher energy till break.

Material H (Fig. 9) shows brittle ruptures for all test speeds: (a) v=10 mm/min, (b) v=250 mm/min and (c) v=1000 mm/min. PA6 droplets could be observed in the broken surfaces, with dimensions of the order of microns, the largest having 10-20 microns, particles partially attached to the PP matrix with EPDM. The presence of EPDM, even at a low percentage of 8% wt, improved the adhesion between the PP + EPDM matrix and the PA6 droplets. In the case of droplets that do not adhere well to the matrix, due to the stress, fixing fibrils, slightly elongated, can be observed. Figure 9b shows typical drops of PA6, partially trapped in the PP matrix (having a more fragile nature).

**Figure 9.** Details of broken surfaces for specimens made of material H tested at different test speed: (a) 10 mm/min; (b) 250 mm/min and (c) 1000 mm/min, with two different magnifications.
For the blend G (Fig. 10), a phase change occurs, so the dispersed phase is PP and its droplet size seems to be quite uniform. The break of the specimen after the tensile test takes place after elongation in the matrix. Second line of SEM images in Fig. 10 shows much larger details (x20000) of tensile fracture surfaces for material G, showing a particular structure, the micromorphology of the fracture surface being with many and longer fibrils at low test speed (Fig. 10a), but being slightly influenced by the higher test speeds (Fig. 10b and c), with shorter fibrils and less visible deformation of the matrix. On the breaking surface of material G, the multitude of droplets (with dimensions of about 10 microns) can be observed and these droplets seem to be trapped in the PA6 matrix through fibrils generated from the matrix.

After performing an EDX analysis (Fig. 11), the constituents of the blends could be identified, so the droplets or islands could be identified as having PA6 if there is a concentration of nitrogen (N) (specific for polyamide) or not (specific for PP and EPDM). In Figure 12, the droplets in the blend H have in their composition the element nitrogen (12.11% wt), this being characteristic for polyamide (C_{6}H_{11}NO)n.

Figure 10. Details of broken surfaces for specimens made of material G tested at different test speed: (a) 10 mm/min; (b) 250 mm/min and (c) 1000 mm/min, with two different magnifications.

Figure 11. EDX analysis for morphology configuration: (a) confirmation of droplets made of PA6 in a matrix de PP (material H) and (b) confirmation of the droplets made of PP in a matrix of PA6 (material G) [26]
For material G, the yield is due to the higher percentage (42% wt) of PA6, and the study of SEM images with EDX analysis found the reversal of the role of constituents: PA6 becomes matrix and PP is dispersed as droplets. It is observed that the PA6 matrix has large local deformations in the tensile breaking section. It is observed that the droplets do not have nitrogen in their composition and knowing that the PP and PA6 polymers are immiscible, it results that the droplets are made of PP and the (visibly ductile) matrix of PA6.

At higher test speeds, the appearance is much different between PP and H, on one hand, and G and PA6, on the other hand. The first two materials have brittle fractures, with fractured, rough surfaces and the other two have visible necking and local material yielding. The aspect of material failure was also kept for Charpy tests [50], brittle for materials PP and H and ductile for materials G and PA6.

6. Conclusions on tensile characteristics

Figure 12 shows four mechanical characteristics, in the sense of increasing PA6 content (0 for PP, 12% for H, 42% for G and 100% for PA6).

![Mechanical characteristics](image)

Figure 12. Mechanical characteristics for tested materials (average values)

Among the studied parameters, the slightly sensitive to the test speed are the modulus of elasticity and tensile strength at break. As concerning the other two parameters (energy at break and strain at break), their values are well differentiated for \( v = 10 \text{ mm/min} \) and for the other two test speeds \( (v = 250 \text{ mm/min} \text{ and } v = 1000 \text{ mm/min}) \), their values being close.

Analyzing Fig. 12, the atypical behavior of material H as compared the other studied blend (G) can be observed, namely, the obtaining of the lowest values of energy at break, which makes this blend not recommended in applications where the energy at break is a decisive factor. If the values of the energy at break obtained for \( v = 10 \text{ mm/min} \) are ignored because, in practice, failure mechanisms at these rates are not often encountered and not accepted in machine components, the other values are grouped in the following ascending
order for all rates (250...1000 mm/min): H, PP, PA6, G. Material G, as developed in [26], [60], [50] have promising results as moderate impact-resistant material. The tensile strength at break shows a tendency to increase the average values with the increase of polyamide concentration, but yet simple polyamide has the highest average values of the tensile strength at break.

At low test rate (v = 10 mm/min), the materials with high energies at break are ductile materials (G and PA6), with values very different from those of simple polypropylene and PP matrix blend (material H). Except for the average value obtained for material G for the strength at break (which, for every test rate has a value close to that of PA6), it can be seen that the other three studied characteristics have even higher average values than the basic constituent (PA6).

Figure 12 points out the increase of the average value of energy at break with the increase of the test speed for material H, but for the other three materials, the increase of the test speed causes the energy at break to decrease much from the lowest test (10 mm/min), the highest value being obtained for material G (38.4 J).

The strain at break has the same trend as that for energy at break for all tested materials, namely the appearance of high average values at the lowest test speed, which is also normal because the slower a load is applied to the test specimens, the specimen has time to deform and, thus, large strains occurs under low stresses.

The values of tensile strength at break (at v = 250 mm/min) are quite close for all materials, the highest value being obtained for PA6 (38.3 MPa).

The modulus of elasticity is slightly sensitive to the test speed (except for v = 10 mm/min), but the PA6 concentration influences this characteristic quite a lot. At v = 250 mm/min, the highest values of this parameter were obtained for the base polymers.

Lower values for energy at break in traction are also be caused by the formation of cavitations during processing and/or cooling (especially in the case of PA6). Mold injection may be the main cause of cavities in many polymers [69].

As concerning the modulus of elasticity (Fig. 12), it can be seen that the base polymers still have the highest average values and that this characteristic is very little sensitive to the test speed and the scattering range of the values obtained for PP is very small. The values of the blends for this parameter are in line with those obtained for the base polymers.

Of the two blends, material G has values well above the blend H, but the spread range of the obtained values (especially for the energy at break and strain at break) is quite large.

If the average values obtained for energy at break for v = 10 mm/min are not taken into account, it can be seen that, as the test rate increases from 250 mm/min to 1000 mm/min, the values increase very slightly for the blends, by 23.96% for blend G.

The large gradient occurs when increasing the test rate from v = 10 mm/min to v = 250 mm/min, respectively v = 1000 mm/min, so the values of energy at break for the material G decrease from 178.4 J to 38.4 J, respectively 47.6 J.

The decrease in values between the test rates of 10 mm/min and 250 mm/min is specific to plastics. The increase of the energy at break for material G with the increase of the test rate is also atypical. At v = 1000 mm/min, the value of energy at break increased by 23.96% as compared to that obtained at v = 250 mm/min.

Following this documentation from the literature, it can be concluded that the immiscibility of polymer blends (in this case, PP + PA6) can be avoided by using compatibilizing agents, namely the addition of an elastomer that would help to produce a morphology suitable for increasing mechanical properties, especially those related to the resistance at moderate impact.

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