VERA OBRADOVIĆ, DUŠICA STOJANOVIĆ, ALEKSANDAR KOJOVIĆ, IRENA ŽIVKOVIĆ, VESNA RADOJEVIĆ, PETAR USKOKOVIĆ, RADOSLAV ALEKSIĆ Scientific paper UDC:620.197.7(n)

# Aramid composites impregnated with different reinforcement: nanofibers, nanoparticles and nanotubes

In this study, the electrospun nanofibers, nanoparticles and nanotubes were used as reinforcement in order to improve mechanical properties of materials for ballistic protection. The samples of polyurethane/p-aramid multiaxial fabric forms (Twaron fabrics and Colon fabrics) were impregnated with poly (vinyl butyral) (PVB) ethanol solution with the different content of reinforcement. Due to the improvement in mechanical, ballistic and impact properties, the surface modification of the fabrics with γ-aminopropyltriethoxysilane (AMEO silane)/ethanol solution was used and silica nanoparticles were modified with AMEO silane. This study is divided into three parts of research: electrospinning process, ballistic testing of the p-aramid composites and impact tester analysis of the Colon composites. The mechanical properties of all the composites were tested by the dynamic mechanical analysis (DMA).

**Keywords:** p-aramid composites, Silica nanoparticles, Carbon nanotubes, AMEO silane, PVB/SiO₂ nanofibers, Ballistic testing, Impact testing.

#### INTRODUCTION

Electrospinning is a simple and unique technology for producing polymer fibers from polymer solutions and polymer melts. The produced fibers are in the micrometre and nanometre scale diameters [1]. The process of electrospinning is driven by a high voltage source to charge the polymer solution. These free charges are on the surface or inside a polymer liquid. The liquid is then accelerated toward a grounded metal collector of opposite polarity. The fiber jet travels allowing the solvent to evaporate from it, thus leading to the deposition of the solid polymer fibers on the collector. In fact, as the jet moves towards the collector, the electrostatic forces accelerate and stretch the jet. Stretching and evaporation of the solvent cause the jet diameter to become smaller. The basic electrospinning set-up consists of a syringe filled with the polymer solution, a needle, a high voltage source and a grounded conductive collector [2]. An electric field is initiated between the tip of a syringe and the collector and thus the electrostatic forces are generated. Under the influence of an electrostatic field, a pendant droplet of the polymer solution at the end of the syringe is deformed into the conical shape called Taylor cone. When these forces overcome the surface tension of the polymer solution, a jet is ejected from the tip of the syringe. Then, the solvent from the jet evaporates leading to the randomly oriented fibers on the collector [3-5].

Author'address: University of Belgrade, Faculty of Technology and Metallurgy, Karnegijeva 4, Belgrade, Serbia

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The schematic drawing of the electrospinning process is illustrated in Figure 1.

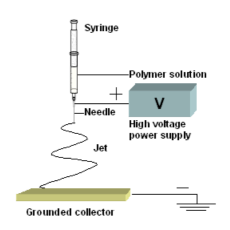


Figure 1- Basic electrospinning set-up

In this work, the effect of the electrospinning process parameters on the morphology of PVB/  $SiO_2$  nanocomposite fibers was analysed.

Woven fabrics produced from high tenacity fibres are used in ballistic protection due to their low density, high strength and high energy absorption capacity. These properties are extremely important for a bullet-fabric interaction which is a complex phenomenon.

The higher kinetic energy of bullet mass and speed during ballistic impact generates more deformation of the fabric. When the bullet speed remains within the limits, the bullet is stopped inside the fabric and the deformation formed on the fabric spreads outwards with the degree of deformation

which depends on the fabric structure. In contrary, when the bullet speed exceeds a certain maximum, the bullet passes through the fabric. Fabric layers have a restricted energy absorption capacity and it is necessary to develop different structures of them to block the bullet when its kinetic energy exceeds fabric energy absorption limits [6,7]. In the work, the *p*-aramid/PVB/AMEO-30 wt.% SiO<sub>2</sub> hybrid composite system was made for this function.

Ceramics have usually been used as reinforcement in the body armour owing to its properties, such as introducing flexibility and light weight of the overall fabric structure [8]. The addition of the nanoscale ceramic reinforcement to polymer matrices improves the mechanical properties, notably compared to the pure matrix material in the body armour [9]. Among this ceramics, nanosilica is one of the primary nanofillers used in polymer composites. The silica nanoparticles are defined by their high specific surface area and due to their collective agglomeration, silane coupling agents modify nanosilica surface for their successful dispersion and deagglomeration, causing the formation of chemical bonds between them and organic components [9,10].

Fiber reinforced composites present impressive engineering materials nowadays. They have their common use in aerospace, automobile, building and military industries. Despite their versatility and great advantages, composites possess certain imperfections and failure properties. Fiber or matrix crackings, as well as delaminations, are common accidents which do not appear in uniform materials [11]. Composite structures are exposed to different impact loading conditions and when they are not fabricated properly, they can undergo a great failure.

It is assumed that impact phenomenon could be divided into four categories according to the impact velocity. Thus, there exist low (< 0.25 km/s), medium (0.25-2 km/s), ballistic (2-12 km/s) and hyper velocity impacts (> 12 km/s) [12].

There are two forms of impact damages on composite structures - complete penetration of the laminate by high-velocity ballistic strikes and nonpenetrating impacts by low-velocity strikes [13]. With a low velocity impact loading conditions, the time of contact between a projectile and a target is considerably long and the load history can be detected with important information about damage progression [14]. During the impact process, the effect of transverse shear and thickness deformations are significant. Delamination appears among the layers, because of insufficient interlaminar bonding which causes a drop in the shear strain that is greater than the bending strain as a result of deformations in the layers [15]. A low speed impact produces local, micro cracks of a matrix material.

This does not consequently lead to a fracture appearance; though it will produce redistribution of the stress and energy concentrations at the interlaminar sections. In these zones, there is a large diversity in material stiffness which generates a fracture growth due to inter-ply delamination [16, 17]. The total impact energy represents a sum of crack initiation energy and crack propagation energy. The fragile materials possess high values of the crack initiation energy and low values of the crack propagation energy, while with the toughest materials it is completely opposite [18].

Carbon nanotubes (CNTs) possess light weight and excellent energy absorption capacity and thus are applied in producing antiballistic materials for bulletproof vests, body and vehicle armor and body armor which can withstand high bullet speed [19, 20]. Pristine CNTs are usually not well dispersed in polymer matrices owing to the strong Van der Waals attractive forces between the nanotubes which tend to their aggregation. A small quantity of CNTs in polymer matrices is more easily dispersed and it is assumed that the modulus is enhanced. On the contrary, when the amount of CNTs is large, the nanotubes may interact and form small aggregates, which reduce the interfacial contact area for stress transfer in the polymer matrix [21]. In order to prevent from the agglomeration due to the presence of the Van der Waals forces between the nanotubes, compatibility with the polymer matrix can be achieved by chemical modification of CNTs with solvents, surfactants, strong acids, noncovalent and covalent bonding and other chemical methods [22-24].

In the current study, the mechanical and impact properties of hybrid thermoplastic composites were tested. The samples of Colon fabrics were impregnated with poly (vinyl butyral)/ethanol solution, and the AMEO silane surface modification of one sample improved the properties besides the addition of multiwalled carbon nanotubes.

# **EXPERIMENTAL**

#### Materials

For preparation of PVB solution in all three cases, a polymer powder, poly (vinyl butyral) (Mowital B60H, Kuraray Specialities Europe) and absolute ethanol (Zorka Pharma, Šabac) were used. The silica nanoparticles with an average particle diameter of about 7 nm (Evonik-Degussa, Aerosil 380) were introduced into the PVB solution. The modification of silica nanoparticles was managed with γ-aminopropyl triethoxysilane (AMEO silane). Multiaxial aramid fabrics (Martin Ballistic Mat, Ultratex, Serbia) were used with *p*-aramid fiber type Twaron (Teijin Aramid). For the impact testing, the carbon nanotubes, MWCNT (Cheap Tubes Inc, USA), were put into the PVB solution. The dimensions of

the nanotubes were from 3  $\mu m$  to 30  $\mu m$  in their length and from 13 nm to 18 nm in their outer diameter. Multiaxial aramid fabrics (Martin Ballistic Mat, Ultratex, Serbia) were used with p-aramid fiber type Colon (Heracron, Kolon Industries, Inc, Korea) which were impregnated with polyurethane (Desmopan, Bayer) [8]. The dimensions of the fabric sample were 15 cm  $\times$  15 cm, and their thickness was around 2 mm.

#### PREPARATION OF THE SAMPLES

The silica nanoparticles were put into the PVB solution (10 wt.%) and stirred continuously for 24 hours and afterwards ultrasonically dispersed in the solution for 15 minutes afterwards. The PVB solutions without the silica particles as well as with them (all with different contents) were tested by changing applied voltages and flow rates in the electrospinning process. A set of experiments was driven with the applied voltages of 16, 20, 24, 28 and 30 kV while the flow rate was kept constant Q = 1 mL/h. Another experimental set was performed with flow rates valued 0.2, 0.4, 0.6 and 0.8 mL/h while the voltage was held at V = 20 kV. During the electrospinning process, the formed fibers were deposited onto the flat aluminium foil which served as a collector. The distance between the needle tip and the collector was h = 10 cm in all cases.

Three pieces of *p*-aramid fabrics (fiber type Twaron) were coated with the PVB solution and the produced PVB-SiO<sub>2</sub> fibers were deposited continuously onto the fabrics during the electrospinning process (Figure 2). The dimensions of each fabric were 29.7 cm x 21 cm, weighing 40 g approximately. The fibers contained 18 g of PVB powder and 9 g of silica nanoparticles (both of them were in 10 wt.% PVB/ethanol solution) and they were deposited onto the aramid fabrics. The flow rate of the PVB-SiO<sub>2</sub> solution for this process was Q=50 mL/h and the voltage was held at V=30 kV, Figure 2.



Figure 2 - The coated aramid fabrics with deposited PVB-SiO2 fibers

The aramid fabrics with the deposited PVB- $SiO_2$  fibers were hot compressed using N 840 D Hix Digital Press (Hix, Corp., USA) at a temperature of 170 °C for 30 minutes under pressure of 4 bar. The fabrics were additionally pressed with P-125 press (170 °C, 10.6 MPa) for 2 hours and then brought under pressure of 20.6 MPa for 1 hour.

For the ballistic properties, the multiaxial aramid fabrics (Martin Ballistic Mat, Ultratex, Serbia) with *p*-aramid fiber type Twaron (Teijin Aramid) were tested. The modification of silica nanoparticles with y-aminopropyl triethoxysilane (AMEO silane,  $(C_2H_5O)_3SiC_3H_6NH_2$ ) began with adding the silane into 95 wt.% ethanol-5 wt.% water mixture in order to reach a 2 wt.% final AMEO concentration. The process of hydrolysis and silanol formation needed about 10 minutes. The solution was homogenized with a magnetic stirrer for 30 minutes after the addition of the nanoparticles. After three hours of sonication, the solution was centrifuged for 30 minutes at a speed of 3000 rpm. After the centrifugation, the particles were placed at the bottom of the beaker and the supernatant was decanted. In the end, the particles were rinsed with ethanol and dried in an oven at 110 °C [8].

The modified silica nanoparticles were with an average particle diameter of about 7 nm and they were added into 10 wt.% PVB/ethanol solution. All the fabrics were coated with  $\gamma$ -aminopropyl triethoxysilane (AMEO silane)/ethanol solution, and then left to stand for about 24 hours for ethanol evaporation. Then, two of the fabrics were coated with poly (vinyl butyral) (PVB)/ethanol solution while the other two were coated with the same solution but with the addition of 30 wt.% AMEO silane modified silica nanoparticles and all of them were left to stay for 24 hours for the ethanol evaporates.

The hard composite samples were hot-pressed using N 840 D Hix Digital Press (Hix, Corp., USA) at a temperature of 170 °C and under pressure of 4 bars for 60 minutes. The fabrics were further pressed with P-125 press (170 °C, 8.2 MPa) for 30 minutes. Within the flexible samples, the flexible composite fabrics were stitched together with the lightweight polyester thread (Korteks, Turkey). The two hard pressed samples contained 17 pieces of fabrics, while the flexible ones consisted of 16 layers of fabrics.

For the impact testing, the 10 wt.% PVB/ethanol solution was used for the experiments. The MWCNT were firstly added into ethanol with concentration of 0.5 and 1.0 wt.%, where MWCNT/ PVB ratio was 0.005 and 0.01, respectively. This ethanol-MWCNT solution was ultrasonicated for 30 minutes separately in order to provide satisfying dispersion of the carbon nanotubes. After this process, the ethanol-MWCNT mixture was added into the 20 wt.% PVB/ethanol solution due to producing a 10 wt.% solution which was mixed on/in a magnetic stirrer overnight. The good dispersion of MWCNT in PVB/ethanol solution was achieved and thus the modification of the nanotubes was not needed while the surface modification of Colon fabrics was applied for some samples.

There were three series of composites which consisted of four pieces of fabrics. The compositions of the samples are presented in Table 1. For the third sample (Colon/AMEO/PVB/1 MWCNT), the fabric was firstly impregnated with AMEO silane/ethanol solution owing to the surface modification. The modification of this composite was managed by impregnation with 2 wt.% AMEO silane/ethanol solution. After the impregnation, the fabrics were left to dry for 24 hours and the appropriate PVB/ethanol solution was coated on the fabrics. The solution consisted of 72 g of ethanol and 8 g of PVB and appropriate MWCNT concentration (0, 0.5 or 1.0 wt.%). Both sides of all the three samples were impregnated with the appropriate solutions, after which they were left to stay for 24 hours for ethanol evaporation. The samples were processed in the compress machine (N 840 D Hix Digital Press) under a pressure of 3 bars for 30 minutes and at a temperature of 170 °C.

Table 1 - The composition of the samples for the impact testing

Sample no.	Composition	
1	Colon/PVB	
2	Colon/PVB/1 wt.% MWCNT	
3	Colon/AMEO/PVB/1 wt.% MWCNT	

#### Characterisation

The electrospinning apparatus used for the experiments in this work is the Electrospinner CH-01 (Linari Engineering) which is located at the Faculty of Technology and Metallurgy in Belgrade, Figure 3



Figure 3 - The electrospinning apparatus at the Faculty of Technology and Metallurgy, Belgrade

The resistance test of the ballistic impact was performed in accordance with the modified ballistic resistance of the personal body armor NIJ standard 0101.04, by applying integrated range instrument system (IRIS) by SABRE ballistics UK [25].

Two types of bullets were used for the ballistic testing: the 357 Magnum FMJ 10.2 g and the 44 Rem. Magnum JHP 15.6 g, with nominal masses of 10.2 g and 15.6 g, correspondingly. The first four shots for each sample were fired with the 357 Magnum and the 44 Rem Magnum was used for the fifth one, all of them being shot at a reference velocity of 436 m/s for each sample as a target. There were fifteen shots altogether during the ballistic test: twelve with the 357 Magnum and three with the 44 Rem Magnum.

The digital ruler with nonius PRO-Max (Fowler) was used for the measuring of the shots' penetration depths. The print and penetration depth image analysis of the shots were performed in Image Pro-Plus software. The images were converted to the grayscale mode, and further subjected to the bitmap analyses. The created images were calculated in pixels and converted to millimeters in order to complete 3D image analysis of the penetration depth.

The specimens for the impact tests had a square shape, with the dimensions of 6 cm × 6 cm and the thickness of about 2 mm. The impact tests were conducted on a high speed puncture impact testing machine Hydroshot HITS-P10 Shimadzu, Japan. The diameter of the striker of the impact testing device was 12.7 mm while the puncture velocity was 2 m/s, and the experiment was carried out at room temperature. The impact tests were run due to the determination of impact force and impact absorbed energy. The impact testing equipment used in this work is presented in Figure 4.



Figure 4 - Overall equipment for the high speed impact tester

## RESULTS AND DISCUSSION

#### Electrospinning results

The average diameters of the electrospun fibers from the optical microscopy images ranged from 1.14 to 2.93  $\mu m$ . It was observed that some

fibers had a diameter smaller than 500 nm and they were visible only by the SEM images. The images showed that, in some cases, the fibers tended to stay close to each other. Probably, the main reason for this was the insufficient working distance h. All the formed fibers were mainly narrow at lower voltages of 16 kV and 20 kV. Oppositely, the electrospun fibers started to remain whipped while increasing the applied voltages from 24 to 30 kV. The whipped fiber structure was also obtained by reducing the flow rate.

The fibers of the PVB solution showed an expected tendency of decreasing their diameters at increasing the applied voltages (Table 2). The maximal value of the average fiber diameter is produced with Q = 1 mL/h, V = 20 kV, and the minimal one is obtained by Q = 0.8 mL/h, V = 20 kV process parameters [26].

Table 2 - Average fiber diameters of the PVB solution with different process parameters

Flow rate	Applied voltage	Average diameter
Q, mL/h	<i>V</i> , kV	d <sub>avg</sub> , μm
0.2	20	1.67
0.4	20	1.68
0.6	20	1.54
0.8	20	1.46
1	16	2.13
1	20	2.94
1	24	2.08
1	28	1.88
1	30	1.77

According to the studies [27-29], the PVB composite fibers with a great content of 50 wt.% SiO<sub>2</sub> nanoparticles were deposited on p-aramid fabrics which had been impregnated with the PVB solution. Figure 5a shows the temperature dependence of the storage modulus for the p-aramid fabric with PVB film and the p-aramid fabric with PVB/PVB-50 wt.% SiO<sub>2</sub> fibers. The composite paramid/PVB showed a lower value of the storage modulus at the temperatures of 40 °C and 70 °C. The difference in the two storage moduli was greater when the temperature was increased (Table 3). The glass transition temperature of the p-aramid/PVB/PVB-50 wt.% SiO<sub>2</sub> fibers composite was 76.4 °C, about 8 °C higher that the p-aramid/PVB composite, Table 3. Tan Delta of the paramid/PVB/PVB-50 wt.% SiO<sub>2</sub> fibers composite had a lower value which indicated that the addition of the PVB fibers with silica nanoparticles promote enhanced thermal properties of the p-aramid fabrics, Table 3, Figure 6b, which was the aim of the addition of the composite fibers. The mechanical

properties were slightly improved, but insufficiently due to the weak bonds among *p*-aramid fabric, PVB and silica nanoparticles.

Table 3 - DMA results for p-aramid fabrics

Sample	E'(MPa) 40 °C	E'(MPa) 70 °C	T <sub>g</sub> (°C)	tan δ
<i>p</i> -aramid	2230	567	68.6	0.37
<i>p</i> -aramid/ 50 wt. % SiO₂ fibers	2336	1355	76.4	0.36

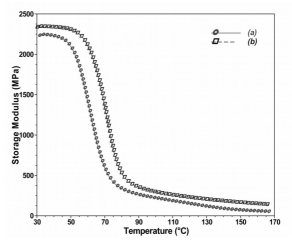


Figure 5 - Storage modulus of: (a) p-aramid/PVB fabrics and (b) p-aramid/PVB fabrics with PVB/SiO<sub>2</sub> nanofibers

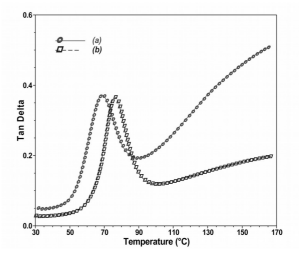


Figure 6 - Tan Delta (tan δ) of: (a) p-aramid/PVB fabrics and (b) p-aramid/PVB fabrics with PVB/SiO<sub>2</sub> nanofibers

The  $SiO_2$  nanoparticles were then modified with AMEO silane. Based on the images of the electrospun fibers with unmodified silica nanoparticles, it was evident that these nanoparticles fused into big aggregates, Figure 7, scale bar 10  $\mu$ m. Unlike them, the modified nanoparticles were well-distributed inside the fibers, Figure 8, scale bar 10  $\mu$ m.

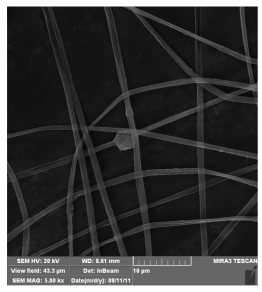


Figure 7 - SEM image of the PVB fibers with 5 wt.% unmodified  $SiO_2$  nanoparticles (Q = 1 mL/h, V = 30 kV)

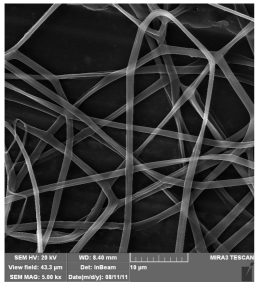


Figure 8 - SEM image of the PVB fibers with 5 wt.% modified  $SiO_2$  nanoparticles ( Q = 1 mL/h, V = 30 kV )

### Ballistic properties results

Dynamic mechanical properties of the three hybrid composites, *p*-aramid/PVB, *p*-aramid/PVB (5 wt. % SiO<sub>2</sub>) and *p*-aramid/PVB/AMEO (5 wt. % SiO<sub>2</sub>) samples were investigated. The samples consisted of four pieces of *p*-aramid fabrics, fiber type Twaron. The storage modulus vs. temperature dependencies are shown in Figure 9. The addition of AMEO-modified silica nanoparticles maximized the storage modulus of the *p*-aramid/PVB/AMEO-5 wt. % SiO<sub>2</sub> composite, where the value of the storage modulus was 2983 MPa, compared with the value of the *p*-aramid/PVB composite which was 1916 MPa. The modification of the nano-

particles with AMEO silane made the chemical bonds between the  $SiO_2$  nanoparticles, polymers and p-aramid fibers, producing significant increase in the mechanical properties of the p-aramid//PVB/AMEO-5 wt. %  $SiO_2$  composite. This modification also led to improved dispersion of the silica nanoparticles and therefore enhanced the mechanical properties of the composite [8].

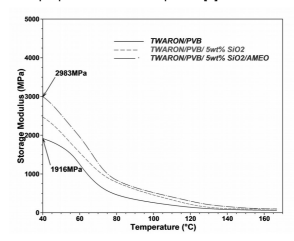


Figure 9 - Storage modulus of the samples

Due to the previous results, the two samples which were used for the bullet shooting tests contained only AMEO modified silica nanoparticles. The ballistic results are presented in Table 4. The samples are denoted as follows: No. 1- 17 pieces of fabrics with PVB - hard option; No. 2- 17 pieces of fabrics with PVB and 30 wt.% AMEO silane modified silica nanoparticles-hard option; No. 3- 16 pieces of fabrics with PVB-flexible option; No. 4- 16 pieces of fabrics with PVB and 30 wt.% AMEO silane modified silica nanoparticles-flexible option. The sample No.1 is removed after the first shot due to the complete penetration of the bullet no.1. The provided number of shots is four instead of five for the sample No. 3 because there was no registered velocity for one bullet.

From the Table 4 it is obvious that the 44 Rem. Magnum bullets made a larger penetration depth than the 357 Magnum ones in each sample. This was typical due to the greater mass of the 44 Rem. Magnum bullets.

Due to the bullet shooting test it is confirmed that the composites of *p*-aramid/PVB without silica reinforcement could not always serve as full protection against bullets. This is evident from the complete penetration of bullet no.1 in sample No. 1 with 17 pieces of fabrics, Table 4, Figure 10. For example, shot no. 2 is realised with the 357 Magnum bullet which partially penetrated 17 pieces of fabrics with PVB and 30 wt.% AMEO silane modified silica nanoparticles, making the indentation depth of 11.65 mm, Table 4). Also, all the other shots produced partial penetrations of the bullets. The introduction of AMEO silane modified silica

nanoparticles in the composite of *p*-aramid/poly(vinyl butyral) provided significant improvement in the mechanical properties, producing hybrid ballistic system of higher protection, Figure 11.

Table 4 - Results of bullet-shooting test

No. of sample	Shot No.	Type of bullet	Indentation depth (mm)
1	1	357 Magnum	complete penetration
2	2	357 Magnum	11.65
2	3	357 Magnum	12.9
2	4	357 Magnum	21.5
2	5	357 Magnum	15.7
2	6	44 Rem. Magnum	21.85
3	7	357 Magnum	7.9
3	8	357 Magnum	10
3	9	357 Magnum	11.3
3	10	44 Rem. Magnum	16.6
4	11	357 Magnum	11.5
4	12	357 Magnum	7.2
4	13	357 Magnum	9.4
4	14	357 Magnum	11
4	15	44 Rem. Magnum	19.9

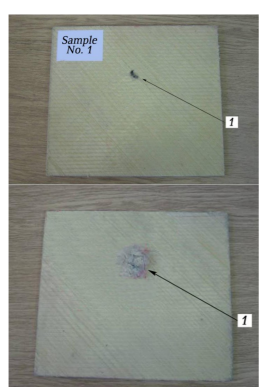


Figure 10 - Front and back side of shot sample No. 1

The magnum bullet no.1 completely penetrated the sample No.1, with the thickness of 8.2 mm. The millimeter scales analysis of the entrance print of this shot is illustrated in Figure 12. A proper 3D graph image of a penetration depth is depicted in Figure 13.

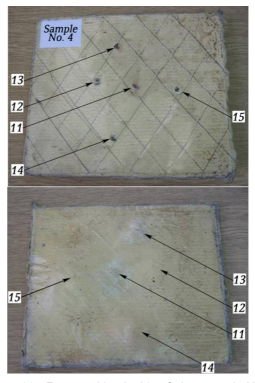


Figure 11 - Front and back side of shot sample No. 4

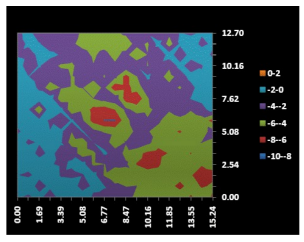


Figure 12 -Print analysis of the shot No. 1

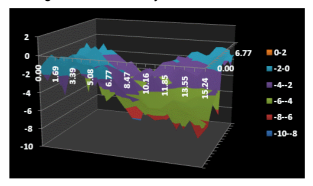


Figure 13 - Indentation depth analysis of the shot No. 1

The sample No.2-hard option had a lower value of the storage modulus (368.4 MPa) compared to the sample No.4 (the flexible option) which was

769.3 MPa due to a process of delamination in the hard sample. Tan  $\delta$  of the sample No.2 had a lower value compared to the sample No.4, owing to the weak bonds among *p*-aramid, PVB and SiO<sub>2</sub> nanoparticles. With glass transition temperature of T<sub>g,2</sub> = 68.94 °C which originates in PVB, the hard sample (No. 2) exposed better thermo mechanical properties than the flexible one (No. 4, T<sub>g,4</sub> = =64.77°C), due to the compression in the hard option.

## Impact tester analysis

The samples were deformed after the impact process, without the complete penetration, Figures 14 and 15. The absorbed energy is accountable for the energy spent in producing damage.

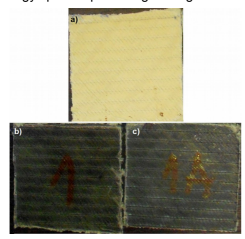


Figure 14 - Samples before the impact test:
a) Colon/PVB; b) Colon/PVB/1 wt.% MWCNT;
c) Colon/AMEO/PVB/ 1 wt.% MWCNT

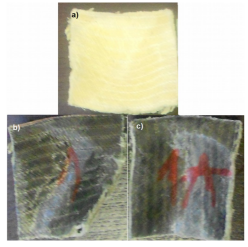


Figure 15 - Samples after the impact test:
a) Colon/PVB; b) Colon/PVB/1 wt.% MWCNT;
c) Colon/AMEO/PVB/ 1 wt.% MWCNT

The force curve in the impact process performs linearly in the beginning, and this part is defined as an elastic zone with small elastic deformations of the composite. After this segment, the change of

the force curve slope appears and with the increase of impact absorbed energy of a striker, the composite begins to endure plastic deformations. Then, the maximum of the impact force occurs ( $F_{\text{max}}$ ), with the appropriate absorbed energy ( $E_{\text{Fmax}}$ ). After this point, there is a sudden drop in force and rise of absorbed energy when a process of delamination develops with a failure. At the end of the impact process, the force returns to zero and relevant absorbed energy at this point presents the total absorbed energy ( $E_{\text{tot}}$ ) which takes a constant value [12].

The results of DMA analysis exposed that the modulus (E') of the Colon/PVB sample increased with the addition of multiwalled carbon nanotubes and surface modification of the fabric (Figure 16). The introduction of MWCNT to the Colon/PVB fabrics increased the storage modulus by 21.2 % for the Colon/PVB/1 wt.% MWCNT and by 60.1% for the Colon/AMEO/PVB/1 wt.% MWCNT sample in comparison with the Colon/PVB respectively, Figure 16, Table 5. These values demonstrate that a good dispersion of MWCNT was achieved and the impregnation of the fabric with AMEO silane maximized the storage modulus for the third sample as a result of the strong bonds between AMEO silane and Colon/PVB surface.

Table 5 - DMA test results for all the samples

Sample no.	E' (MPa, 40 °C)	tan δ1	Tg (°C)
1	1377	0.2934	65.53
2	1669	0.3029	66.35
3	2204	0.3012	69.21

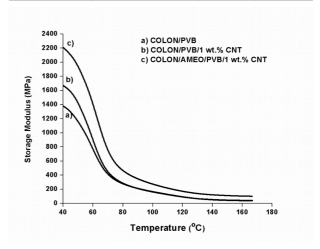


Figure 16 - Storage modulus of the samples

The first peak of Tan Delta shows the glass transition temperature,  $T_{\text{g}}$ , of PVB in the impregnated Colon fabrics. The  $T_{\text{g}}$  value for the Colon/PVB sample was 65.53 °C, and the value for

the Colon/PVB/ 1 wt.% MWCNT sample was  $66.35^{\circ}$ C, Figure 17, Table 5. For the third sample, after modification with AMEO silane, the storage modulus increased considerably and  $T_g$  progressed slightly. The value of  $T_g$  for the Colon/AMEO/PVB/1 wt.% MWCNT was  $69.21^{\circ}$ C. Hence, there was a light increase in thermal stability with the addition of MWCNT and modification with AMEO silane which diminished the overall mobility of the polymer chains.

Tan Delta value of these fabrics was almost the same for all the samples at the  $T_{\rm g}$  temperature, Table 5, Figure 17. With the rise of temperature, the Tan Delta value of the Colon/AMEO/PVB/1 wt. % MWCNT sample was decreasing due to the strong bonds among AMEO silane, PVB, MWCNT and polyurethane. The glass transition temperature of polyurethane is about 150 °C.

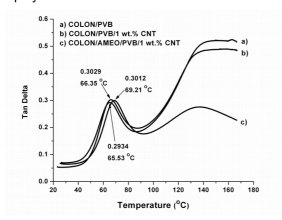


Figure 17 - Tan Delta of the samples

The results of the impact tester analysis are given in Tables 6 and 7. The maximum impact force ( $F_{\text{max}}$ ) of the Colon/PVB samples increased with the addition of pure multiwalled carbon nanotubes. According to Table 3, the AMEO modified sample demonstrated the greatest impact force (4.55 kN) compared to the Colon/PVB sample (3.20 kN). The result for the Colon/AMEO/PVB/ 1 wt.% MWCNT yielded around 42% improvement in the maximum impact force compared to the Colon/PVB sample.

All the samples had very similar time and displacement for their peak force values, Figures 18 and 19, Table 6.

Table 6 - Impact force test results for all the samples

Sample no.	Time, (msec)	Displacement (mm)	Fmax (kN)
1	13.66	12.11	3.20
2	10.47	16.11	3.64
3	12.43	18.75	4.55

The slope of the linear part of the force-time curve is proportional to the storage modulus [30]. It increased from the first sample to the third one, with the addition of MWCNT, and AMEO silane surface modification, Figure 18.

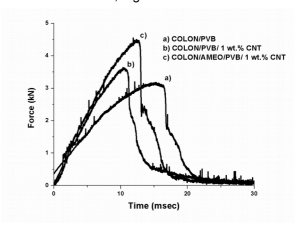


Figure 18 - Force versus time for all the samples

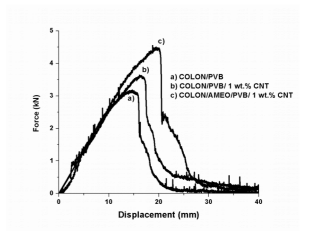


Figure 19 - Force versus displacement for different Colon/PVB fabrics

Similarly to the results for the impact forces, the total absorbed energy of the Colon/PVB samples increased with the addition of pure multiwalled carbon nanotubes, Figures 20 and 21. Also, from Table 7, the AMEO modified sample displayed greater absorbed energy compared to the other two samples. The third sample produced around a 73% improvement in the total impact energy absorption (65.69 J) compared to the Colon/PVB sample (37.97 J).

Table 7 - Impact energy test results for all the samples

Sample no.	$E_{Fmax},(J)$	E <sub>tot</sub> , (J)
1	18.84	37.97
2	31.86	49.94
3	44.32	65.69

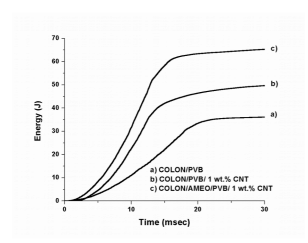


Figure 20 - Energy versus time for all the samples

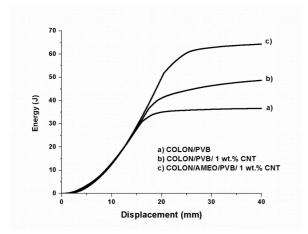


Figure 21 - Energy versus displacement for different Colon/PVB fabrics

The results of the samples from the impact tester analysis were in coherence with the results from DMA analysis and they indicated the effect of the AMEO modification and MWCNT content on the mechanical and impact properties of the composites.

# CONCLUSION

The structure and morphology of the electrospun  $PVB/SiO_2$  composite fibers was investigated by changing two process parameters - the flow rate of the solution Q and the applied voltage V, as well as the nanoparticle content and treatment. The produced nanofibers were merely narrow at low voltages of 16 kV and 20 kV. However, the fibers began to whip at the voltage of 24 kV and by further increasing, up to 30 kV, they were completely coiled.

The addition of the PVB fibers with silica nanoparticles promote enhanced thermal properties of the *p*-aramid fabrics comparing to ones coated with PVB film, producing hybrid ballistic system of higher protection. The penetration of the samples during the impact test did not occur. The Colon/AMEO/PVB/1 wt.% MWCNT sample exhibited the increase of storage modulus, impact force and absorbed energy compared to the Colon/PVB sample. The results from DMA analysis were consonant with the ones from impact tester analysis and they demonstrated that the best results for the properties of Colon/PVB sample were achieved with the addition of 1% wt. concentration of MWCNT and the surface modification with AMEO silane.

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## **IZVOD**

# ARAMIDNI KOMPOZITI IMPREGNISANI SA RAZLIČITIM OJAČANJIMA: NANOVLAKNA. NANOČESTICE I NANOCEVI

U ovom istraživanju su se elektrospinovana vlakna, nanočestice i nanocevi koristili kao ojačanja koja bi unapredila mehanička svojstva materijala za balističku zaštitu. Uzorci poliuretan/p-aramidnih multiaksijalnih lamina (Twaron i Kolon lamine) su impregnisani sa poli(vinil butiral) (PVB)/etanol rastvorom sa različitim sadržajem ojačanja. Radi poboljšanja mehaničkih, balističkih i impakt svojstava, modifikovala se površina lamina sa γ-aminopropiltrietoksisilanom (AMEO silan)/etanol rastvorom i koristile su se silika nanočestice modifikovane AMEO silanom. Ovaj rad je podeljen na tri dela istraživanja: proces elektrospininga, balističko testiranje p-aramidnih kompozita i impakt tester analiza Kolon kompozita. Mehanička svojstva svih kompozita su bila testirana pomoću dinamičko mehaničke analize (DMA).

**Ključne reči**: p-aramidni kompoziti, silika nanočestice, ugljenične nanocevi, AMEO silan, PVB/SiO<sub>2</sub> nanovlakna, balističko testiranje, impakt testiranje.

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