The composition of electrical wires and cables is of critical importance in controlling fire risks. In this respect, polyvinylchloride (PVC) composites are extensively used. Yet, PVC composites are multiple systems in which the final properties depend on the nature and size of the reinforcement or flame-retardant agent and the type of lubricant used for their preparation. Thereby, in this study two series of PVC composites, with stearic acid and calcium stearate as lubricants were prepared in parallel, and additivated with various commercial minerals as reinforcement/flame retardant agents, such as calcium carbonate, alumina trihydrate (ATH) and HMH (a mineral of hydromagnesite and huntite). Following the structure (by FT-infrared), thermal behavior (thermal analyses and differential scanning calorimetry) and mechanical properties (dynamic mechanical analyses, tensile strength and elongation at break) of PVC composites, the flame-retardant effect of the selected minerals was investigated by measuring the limiting oxygen index (LOI). Only ATH and HMH-based composites presented higher flame-resistance, relative to the control samples, making them suitable for the proposed application. The contact angle was evaluated for determining the hydrophobicity of composites when using ATH or HMH, to get an opinion about the stability of the materials in moist environments. Finally, SEM was used to determine the homogeneity of PVC samples.

Keywords: PVC, alumina trihydrate, Ultracarb, stearic acid, calcium stearate, flame retardant

PVC is the third most used thermoplastic, after polyethylene and polypropylene. PVC is a versatile polymer and has been widely used because of its low production cost, electrical protecting properties and good insulation performance, chemical and fire resistance, and also high abrasion resistance.

In this context, flexible PVC is intensively used as covering insulation for electrical and communication cables, while rigid PVC is used in domestic applications, such as window frames, doors, profiles and sidings because of its high level of combustion resistance.

PVC, when heated at temperatures above 140 °C releases smoke and toxic gases like hydrogen chloride (HCl) due to dehydrochlorination reactions. Hence, obtaining PVC composites with flame retardant and smoke suppression properties has received an increasing interest during the past decades. Rigid PVC composites do not contribute to fire propagation due to high contents of chlorine (56.8 %), and thus, passes the flammability test without adding flame retardants additives. For flexible PVC composites on the other hand, flame retardant agents must be used because the plasticizer content reduces the resistance to fire [1-4]. In this respect, reinforcing or flame retardant agents or fillers are also used in PVC systems to reduce formulation costs and improve the electrical insulation properties. Moreover, fillers are carefully chosen due to the significant impact upon the composite system, i.e. increased tensile strength and thermal stability and reduced costs [5]. Common filler materials include calcium carbonate of different sizes, kaolin, aluminium hydroxide (or alumina trihydrate), magnesium hydroxide, mica, silica, talc.

When PVC composites are exposed to fire, they decompose and the chlorine content is converted to hydrogen chloride gas, which can be released in the vapour state. Because this gas is toxic it should better remain in the condensed phase. This can be accomplished by adding calcium carbonate in the system, for example, to form calcium chloride [6]. The calcium carbonate particles can capture the emitted HCl and minimise the corrosivity of the smoke, if used as filler; and, hence, acting like a thermal stabilizer [7-8]. Such action also prevents the early PVC decomposition into conjugated polyenes. This latter process is the result of a zipper dehydrochlorination process catalyzed by hydrochloric acid [3, 6].

Alumina trihydrate (noted ATH) is highly used in the thermoplastics industries as a flame retardant and smoke suppressant. The major interest for ATH refers to the water content, about 34.6 % H2O by weight. This intramolecular water is endothermically released at over 200 °C. This means that the heat need for volatilizing the side-products is used for releasing the intramolecular water form ATH. Hence, less heat is accessible to decomposing the polymer components that feed the combustion. In this case, ATH acts as a heat sink when the composites are exposed to fire. Furthermore, ATH increases the char formation in the polymer and a protective layer made up of aluminum oxide and carbonized products are formed on the surface of the polymer, hindering the smoke release upon combustion. Therefore, it can be noted that ATH works both as flame retardant and as smoke suppressant. As a result, the loading of ATH in flexible PVC formulations usually ranges between 20 to 50 phr (parts per hundred resins, base polymer = PVC) to decrease the heat release rate and delay the time to ignition as much as possible [8-10]. The two synergistical effects also improve the limiting oxygen index (LOI). The relative flammability characteristics of
polymeric materials are determined by LOI. The higher the LOI values the greater the resistance of the material to burning (more fire retardant). In this work the targeted values for LOI are beyond the threshold of 21% from where the composites can be classified as flame resistant [11-13].

Another mineral used as flame retardant is a mixture of two minerals: huntite and hydromagnesite (MMH, or Ultracarb as commercial product). It is used particularly in the wire and cable industries and for rubber applications as an alternative to the more expensive ATH. Each of the two minerals, i.e. hydromagnesite and huntite (MMH), are flame retardants and mixing them leads to new products with special properties for various applications. MMH has a unique 3-stage fire retardant mechanism. While the common endothermic flame-retardant fillers act by releasing water in a single process, MMH works as follows: at temperatures of ~ 220°C begins the thermal decomposition and releases water. At around 330°C, CO₂ is released and at temperatures above 560 °C MMH particles accumulate at the surface of the burning PVC composites and form a stable char. This final step leads to the hindering of the combustion process and implicitly the propagation of the fire [9, 14-17].

Considering the present literature, this paper provides an interesting comparative study upon the influence of calcium carbonate with particle size of 1 and 2 mm, of ATH and MMH as fillers, and the influence of stearic acid and of calcium stearate, as lubricants upon the thermal, the mechanical, the structural and the flame resistance properties of novel PVC composite formulations with potential applications in the industry for electrical wires and cables.

Experimental part

Materials

For this research the used materials included PVC Ongrovil S5070 from BorsodChem (K-value 70.4), diisononyl phthalate (DINP) as plasticizer (LG Chem), Ca-Zn salt as heat stabilizer (Baeropan MC 91446 KA/3, Baerlocher), calcium carbonate particles of 1 and 2 µm (noted VO1 and VO2, respectively) as reinforcing agent (Omya Calcita), Martinal ATH OL-104 LEO (Huber) and MMH (Ultracarb, LKAB Minerals) as flame retardants, stearic acid (Steral Chemicals) and calcium stearate as lubricants (Bridgegim).

Preparation of PVC composites

The test specimens were prepared using an industrial methodology, as follows: first, the samples were prepared by melting and mixing PVC with the plasticizer, the heat stabilizer, lubricant and reinforcing agent/flame retardant. Afterwards, the specimens were blended into a two-roll mill at 160 °C for 20 min, followed by compression at 170 °C to form sheets with dimension 150 × 150 × 1 mm³. The compositions of all of the samples are listed in table 1. All the samples, except samples 2 and 3 which are not filled, have the following formulation: PVC 100 phr, DINP 60 phr, inorganic filler 40 phr, lubricant 0.5 phr and heat stabilizer 4 phr. DINP used as plasticizer provides compound flexibility, which improved viscosity and processability [18-19]. This type of phthalate plasticizer behaves like solvents but is not chemically bounded to PVC. The DINP physically bounds PVC by weak electrostatic interactions as the result of heating process during composites manufacturing.

Sample 1 of stabilized PVC and lubricant alone, prepared as a control sample, degraded during the preparation process having a brown-reddish colour and being very brittle. Therefore, these samples could not be analysed and the results were compared with the control samples 2 and 3, which contain stabilized PVC, lubricant and plasticizer (i.e. sample 2 with stearic acid and sample 3 with calcium stearate).

| Sample code | Inorganic filler | Lubricant |
|-------------|-----------------|-----------|
| 2           | -               | Stearic acid |
| 4           | VO2             | Stearic acid |
| 5           | VO1             | Stearic acid |
| 8           | ATH             | Stearic acid |
| 10          | MMH             | Stearic acid |
| 3           | -               | Calcium stearate |
| 6           | VO2             | Calcium stearate |
| 7           | VO1             | Calcium stearate |
| 9           | ATH             | Calcium stearate |
| 11          | MMH             | Calcium stearate |

*All the samples, except samples 2 and 3, which are not filled, have the following formulation: PVC 100 phr, DINP 60 phr, inorganic filler 40 phr, lubricant 0.5 phr and stabilization agent 4 phr

Infrared spectroscopy analysis (FTIR)

Molecular structure of PVC based composites were analyzed using an infrared spectrometer type JASCO FT/IR 6300 Specac Golden Gate ATR with diamond in transmission mode. The measured range was 400-4000 cm⁻¹.

Thermogravimetric analysis (TGA)

The thermogravimetric analysis were performed on a thermogravimetric analyzer (Q5000IR, TA Instruments) at a heating rate of 10 °C min⁻¹ under a nitrogen atmosphere or air. Samples of approximately 8 - 14 mg were analyzed in the 40-700 °C temperature range.

Differential scanning calorimetric analysis (DSC)

Differential scanning calorimetry (DSC) experiments were carried out using DSC Q2000 from TA Instruments, working in temperature-modulated mode (MTDSC) with dynamic purge gas (helium 99.99% with flow rate 25 mL min⁻¹, modulated scan). Approximately, 25 - 30 mg samples were weighted and heated from -110 to 155 °C with a heating rate of 20 °C min⁻¹.

Dynamic mechanical analysis (DMA)

Dynamic mechanical analysis (DMA) experiments were carried out using DMA Q800 from TA Instruments, module DMA Multi-Frequency-Strain, in the tension mode, obtaining thermal behaviours of the PVC based composites, through the temperature ramp method. The sample size was 13.5 × (6.4-7) × (0.77-0.79) mm and the heating rate was 5 °C min⁻¹ from -75 to 105 °C.

As a result of the experiments, the temperature dependences of the elastic shear modulus (E’), shear losses (E”) and the mechanical loss tangent (δ) of the investigated PVC composites were extracted.
FTIR analysis

The transmission mode was applied to obtain the spectra for PVC composites - samples 2 and 3 considered control samples. The two spectra are similar because the main difference is represented by the lubricant type (stearic acid or calcium stearate). Figure 1. a and b shows the characteristic bands of PVC, plasticizer and the two lubricants. In the both spectra the CH and CH2 bands from the PVC backbone are present at 2851 and 2927 cm⁻¹, PVC characteristic band attributed to C-Cl bond appeared at 600 cm⁻¹. The band from 1730 cm⁻¹, corresponding to C=O vibrations is attributed to the lubricant (aliphatic carboxyl) and phthalate plasticizer (aromatic carboxyl). Carbocatenary long chain (CH and CH2, bonds) of the plasticizer and of the lubricants are overlapping with those of PVC at 2851 and 2927 cm⁻¹ and at 1262 and 1471 cm⁻¹ the vibrations of C-O groups from the lubricant (aliphatic carboxyl) and phthalate plasticizer (aromatic carboxyl) were observed. Figure 1. a and b also depicts the spectra of PVC based composite with filler (i.e. flame-retardant agents). From it, the presence of additional bands corresponding to the incorporated fillers was observed. Samples with the two types of calcium carbonate VO1 and VO2 (i.e. samples 4,5 with stearic acid as lubricant -fig.1. a and samples 6,7 with calcium stearate as lubricant -fig.1. b), display specific bands around 880 cm⁻¹ attributed to Ca-O vibrations, and bands from 1425 cm⁻¹ corresponding to C-O group from carbonate structure.

The presence of ATH in the PVC composites (sample 8 with stearic acid as lubricant -fig.1. a and sample 9 with calcium stearate as lubricant -fig.1. b) is demonstrated by the band corresponding to hydroxyl group at 3635 cm⁻¹. Further on, for the PVC composites with HMH inorganic filler (sample 10 with stearic acid as lubricant -fig.1. a and sample 11 with calcium stearate as lubricant -fig.1. b) the bands observed at 3518 cm⁻¹ belong to -OH group from hydromagnesite. Here as well, the band of anion corresponding to the huntite is displayed at around 1551 cm⁻¹.

Thermogravimetric analysis (TGA)

The TGA curve for PVC alone, measured under inert atmosphere (N₂) is presented in fig. 2, where it is compared to the profiles of composites. The total mass loss of PVC alone is about 62 %. In the temperature range 150 - 260 °C no significant mass losses are observed (approximately 2.4 %). Thermal degradation of PVC takes place in two stages as follows: in the first stage (260-380°C), release of hydrochloric acid, followed by the formation of polyene structures, and in the second stage (380 - 550 °C) cyclization of the polyene structures with the formation of aromatic compounds. Thermal decomposition in the second step is represented by degradation of PVC backbone and, therefore, lower molecular weight compounds (and smoke) are produced [20-22].

After the addition of plasticizer and lubricant, the mass loss is about 14.7 % for sample 2 (fig.2. a) and about 15 % for sample 3 (fig.2. b) over a temperature range of 150-260 °C. This increase in mass loss compared to neat PVC is due to plasticizer desorption. Desorption of a low molecular weight compound comprises two consecutive steps: diffusion of the plasticizer from the mass of the sample to its surface and evaporation from the surface. Further on, elimination of the hydrochloric acid from the composites begins in the range of 260-380 °C and the mass loss increases significantly up to 56 % for sample 2 and 57 % for sample 3. In this temperature range also begins the decomposition of the lubricants: stearic acid or stearate calcium (and also of Ca-Zn stabilizer, which is also a metal carboxylate). When adding reinforcing/fire-retardant agents, the loss of mass decreases and the decomposition temperature also presents higher maximum compared to samples 2 and 3, as summarized in tables 2 and 3.
PVC-based composites reinforced with calcium carbonate of 2 µm (sample 4 lubricated with stearic acid and sample 6 lubricated with calcium stearate) show lower mass losses compared to samples with 1 mm calcium carbonate (5 and 7, respectively) and to control samples 2 and 3 with no reinforcing agent. Calcium carbonate, used as a reinforcing agent, reduces the emissions of hydrochloric upon composites degradation by trapping it in its structure to form CaCl₂. It looks like calcium carbonate with larger particle sizes (VO2) absorbed higher amounts of hydrochloric acid, so the mass loss in the range of 260-380 °C was lower compared to the small particle size of calcium carbonate (VO1). Hence, it was confirmed that calcium carbonate acts as a heat stabilizer by capturing hydrochloric acid, especially compared to the control sample 2, which presented much higher mass losses in this range. Yet, calcium stearate, besides its role as lubricant, has a co-stabilizing role, and hence, the decomposition maximum of PVC is shifted with 2 °C compared to the stearic acid-based PVC composites (fig.2. b). It can also be noted that stearic acid-based PVC composites also presented a shift for the decomposition maximum of PVC with 3 °C relative to the control samples (fig.2. a).

The effect of fire retardants and smoke suppression agents is due to the Lewis acid catalyst. Lewis acids can favor PVC crosslinking to quickly form coal (ash) during burning. Thus, by this mechanism, metals from flame retardants (ATH and HMH) can favor PVC crosslinking during decomposition to increase coal formation [8-10]. Thermogravimetric analysis for samples with ATH and HMH (samples 8 and 10 with stearic acid—fig.2. a, and samples 9 and 11 with calcium stearate—fig.2. b) showed that the rate of polymer decomposition is slowed-down because the heat is absorbed by the decomposing metal hydroxide and the released water vapours dilute the flammable by-products. Further on, the metal oxide residue acts as a thermal barrier and prevents the decomposition of the underlying polymer. From the results obtained for the samples containing ATH and HMH, classified as flame retardants, it seems that the highest residue at 550 °C is obtained when HMH is used. The effect of ATH upon PVC decomposition is similar to the one obtained when calcium carbonate was used as reinforcing/flame retardant agent according to the residues at 550°C. Furthermore, the residues of the samples with ATH and HMH at 700°C, for measurements performed in air indicated that HMH was more efficient for suppressing the smoke emitted during combustion (table 4).

### Table 2

| Sample | 40 - 150°C | 150 - 250°C | 250 - 380°C | 380 - 550°C |
|---|---|---|---|---|
|   | Wt. loss | Wt. loss | Wt. loss | T₂ range | Wt. loss | Tmax |
| PVC | 0.01 | 2.42 | 61.78 | 291.0/330.2 | 27.96 | 453.2 |
| 2  | 0.12 | 15.71 | 58.63 | 268.8/281.0 | 13.09 | 458.6 |
| 4  | 0.10 | 13.52 | 44.14 | 279.8/290.1 | 11.78 | 457.6 |
| 5  | 0.07 | 12.82 | 45.19 | 276.7/278.2 | 12.46 | 459.8 |
| 8  | 0.06 | 11.72 | 52.15 | 293.3/310.8 | 11.03 | 459.8 |
| 10 | 0.13 | 12.20 | 49.91 | 272.3/288.3 | 16.05 | 456.7 |

### Table 3

| Sample | 40 - 150°C | 150 - 250°C | 250 - 380°C | 380 - 550°C |
|---|---|---|---|---|
|   | Wt. loss | Wt. loss | Wt. loss | T₂ range | Wt. loss | Tmax |
| PVC | 0.01 | 2.42 | 61.78 | 291.0/330.2 | 27.96 | 453.2 |
| 3  | 0.09 | 14.12 | 57.82 | 279.8/292.3 | 15.40 | 459.7 |
| 6  | 0.08 | 14.07 | 45.79 | 277.8/289.8 | 11.67 | 458.9 |
| 7  | 0.08 | 11.38 | 46.29 | 278.2/288.9 | 10.81 | 451.8 |
| 9  | 0.08 | 14.58 | 49.16 | 286.3/310.1 | 12.42 | 450.5 |
| 11 | 0.13 | 10.51 | 46.04 | 273.3/287.3 | 17.22 | 457.8 |

Fig. 2. Thermogravimetric analysis for PVC based composites with and without filler: a) PVC composites based on stearic acid as lubricant and b) PVC composites based on calcium stearate.
Differential scanning calorimetric analysis (DSC)

Using DSC, the fluctuations in the glass transition temperature, $T_g$, of obtained composites were evaluated. The need for studying this behavior of PVC composites refers to the fact that decomposition temperature of neat PVC (above 160 °C) is lower than the melting temperature (260 °C). As a result, it is necessary to lower the $T_g$ to prevent degradation of the polymer before processing, but also to establish a maximum temperature of processing and application of the composites [23-26].

In the DSC diagrams (fig.3. a and b), the shift of the baseline following the glass transition for each PVC sample can be observed. The variations in the $T_g$ depend upon the type of filler but also upon the type of used lubricant. Neat PVC, without additives, shows a $T_g$ of 85 °C. According to fig.3. a, the smallest $T_g$ is obtained for the samples with stearic acid, filled with calcium carbonate of 2 µm (sample 4, $T_g = -40.5^\circ$C) and HMH (sample 10, $T_g = -40.2^\circ$C). PVC compositions with calcium stearate (fig.3. b) have slight higher $T_g$ values relative to stearic acid-based composites. The lowest values for $T_g$ in the samples series with calcium stearate were obtained for samples 6 and 7, with calcium carbonate of 2 µm ($T_g = -39.4^\circ$C) and 1 µm ($T_g = -39.1^\circ$C), respectively. Thus, the influence of the lubricant nature is perceivable as the $T_g$ shifts towards lower values when stearic acid is used as lubricant (-40.5°C). One explanation to this $T_g$ decrease for samples with stearic acid can be due to the polarity of stearic acid, which seems to lead to strong interactions with the PVC matrix [26-28].

The effect of lubricants can be better seen from the summarized data on DSC in table 5. The widening of the $T_g$ range (the difference between the Onset and the End point) is narrower for nominated samples 4, 10 and 6, 7 containing either stearic acid or calcium stearate, respectively. It can be mentioned here that the heat capacity ($\Delta C_p$) that accompanies glass transition was also in line with these observations, showing lower values, particularly for sample 4 with stearic acid as lubricant.

Dynamic mechanical analysis (DMA)

The effect of filler loading on the dynamic mechanical properties of PVC composites was studied. DMA was complementary on determining the storage modulus, loss modulus at maximum value and the dynamic loss tangent at maximum value [29-30]. When the storage modulus ($E'$) increases with the filler loading usually indicates that mobility of the polymer chain is decreased in the presence of the reinforcing/flame-retardant agents. In this study, it was observed that filler incorporation into the PVC matrix may decrease the mechanical damping of PVC (fig. 4. a and b). The height (peak intensity) and position of $E'$ peaks for the composite samples varied with type of filler.

The values of $E'$ were determined for each sample at a temperature of -70 °C. The highest $E'$, 7583 MPa, was attained by sample 11 (with HMH as filler and calcium stearate as lubricant) while the lowest $E'$, 4799 MPa, was

| Sample | Residue at 700°C (%) |
|--------|----------------------|
|        |summary for PVC based composites with ATH and HMH |
characteristic of sample 7 (with calcium carbonate of 1 mm and calcium stearate as lubricant). The registered values of E' for the control samples 2 and 3 were 5426 MPa and 4829 MPa, respectively. A clear comparison of characteristics can be extracted from table 6. The values of E' decreased, while the E'' values gradually increased. Hence, at temperatures above -20 °C, all the investigated composites presented high elasticity. The filler incorporation increased the storage and loss modulus, which explained the higher heat dissipation in the PVC composites compared with neat PVC, confirming that the new formulations of PVC are flexible.

**Tensile strength and elongation at break**

To evaluate the quality of flexible PVC composites in industrial application, one of the most important properties is tensile strength. It can be observed from fig. 5. a, b, c and d that elongation at break and tensile strength of test samples decreased with the filler loading for each type of PVC composite. This decrease in mechanical properties is attributed to restriction of polymer chain movements [31-32]. For control sample 2 and 3 with no filler, the highest mechanical properties values are obtained. Yet, depending on the subsequent use of the composites certain properties are required.

Stearic acid composites (fig.5. a and b) exhibit large variations in mechanical properties, due to the size of calcium carbonate and the capacity of the lubricant to allow homogenization during composites preparation. Sample 5 with 1mm calcium carbonate attained the highest tensile strength and the lowest elongation from the series of composites with stearic acid and fillers, which is bizarre but maybe due to the low mobility of the polymer. Calcium carbonate of 2 µm (sample 4) is better embedded in the material and more elastic samples are obtained. For the series with calcium stearate (fig.5. c and d) it seems that the size of the calcium carbonate did not affect the mechanical properties significantly. This means that the lubricant also plays an important role in the compounding of the samples. Calcium stearate ensures more uniform mechanical properties due to better compatibility with calcium carbonate, and hence, the size of this filler does not affect the tensile strength and elongation at break.

When the reinforcing/flame-retardant agent is added to the composites, the tensile strength and elongation at break decrease continuously even more for the PVC composites with stearic acid as lubricant (fig.5. a and b), with exception for sample 5. For the calcium stearate-based composite (fig.5. c and d), adding ATH (sample 9) in the formulation leads to similar values to that of samples 6 and 7 with calcium carbonate, meaning elongation at break 432 % and tensile strength of 16 N/mm². It can be noted that HMH based composites, samples 10 and 11, presented the lowest values for tensile strength and elongation at break (except for sample 5) in both series of composites.

**Limited oxygen index (LOI)**

![Fig. 4. Dynamic mechanical analysis for PVC based composites with and without filler: a) PVC composites based on stearic acid as lubricant and b) PVC composites based on calcium stearate](image)

| Sample | Loss Modulus | Tan Delta | Storage Modulus |
|--------|--------------|-----------|-----------------|
|        | E' Peak      | Tan δ     | E', MPa|Temperature, °C |
|        | Temp. | Height | Temp. | Height | Onset | Midpoint | End | -70 | 0 | 30 | 100 |
| 1      | -33.69 | 536.2 | 35.32 | 0.3674 | -47.41 | -34.44 | 1.98 | 5426 | 1012 | 180.2 | 6.45 |
| 2      | -31.23 | 604.2 | 32.55 | 0.369 | -45.26 | -29.45 | -2.22 | 6512 | 1014 | 172.8 | 8.229 |
| 3      | -29.07 | 541.1 | 31.17 | 0.3748 | -43.66 | -23.06 | 4.29 | 6165 | 1238 | 222.6 | 8.54 |
| 4      | -33.69 | 580.2 | 31.62 | 0.3713 | -51.6 | -40.02 | 2.31 | 6414 | 985.3 | 149.6 | 8.01 |
| 5      | -35.85 | 713.2 | 28.85 | 0.374 | -47.45 | -31.25 | -2.37 | 7293 | 1091 | 144.8 | 8.501 |
| 6      | -31.23 | 462.4 | 37.48 | 0.3714 | -47.15 | -34.94 | 6.82 | 4829 | 1039 | 196.6 | 6.497 |
| 7      | -30 | 571.6 | 32.55 | 0.3682 | -46.29 | -28.72 | -0.85 | 6156 | 1018 | 171.4 | 8.109 |
| 8      | -31.84 | 454.4 | 30.7 | 0.3653 | -42.8 | -36.48 | 2.45 | 4759 | 902.2 | 135.1 | 7.338 |
| 9      | -32.46 | 624.7 | 31.07 | 0.3751 | -51.86 | -24.14 | -1.27 | 7184 | 1054 | 152.4 | 7.92 |
| 10     | -31.84 | 703.1 | 32.86 | 0.3708 | -47.53 | -25.9 | 1.77 | 7383 | 1392 | 194 | 8.849 |
The minimum oxygen concentration in a variable oxygen/nitrogen atmosphere is known as the limiting oxygen index (LOI) that will just support ignition of a sample. This test is generally recognized to be useful for quality control purposes [33-36]. The LOI test is extensively valuable and used to determine the flammability of polymeric materials. Greater fire retardation is given by a higher measured value of LOI. Values of LOI greater than 21 indicate that composites do not burn immediately in air.

The LOI values of tested PVC composites are shown in figure 6. a and b. The results indicated that the minimum amount of oxygen for the ignition of sample had an increasing trend when using flame retardants in the composition. For control samples 2 and 3 with no filler in the composition the LOI value is around 21, so the sample can ignite in air easily. As expected, when adding filler in the composition, the LOI value increase.

Calcium carbonate acts as scavenger during ignition, forming calcium chloride with the hydrochloric acid released during ignition. This process hinders the reaction between oxygen and PVC molecules, and implicitly, leads to high LOI values [6, 37]. But, the flame-retardation effect of calcium carbonate is limited. In a study made by Xu et al., calcium carbonate showed little or no flame-retardant effect on PVC and for improving the efficiency, the authors coated CaCO₃ with SnO₂ [38]. Herein, the LOI values increased for the samples filled with calcium carbonate in both series of composites relative to the control samples 2 and 3. For the 2 mm calcium carbonate composites, i.e. samples 4 and 6, the values were above 23 %, while for the ones with 1 μm calcium carbonate, i.e. samples 5 and 7, between 22.2 and 22.6 %. The results obtained for the two types of calcium carbonate fillers are in agreement with the thermal analyses, which indicated a higher capacity of the 2 μm calcium carbonate to adsorb hydrochloric acid.

LOI values of PVC based composites increased even more when ATH and HMH were used in the PVC composite formulation. ATH and HMH decompose endothermically and release water. The endothermic decomposition of water hijacks the heat needed for polymer burning and retards the flame. This phenomenon is also referred as sink phenomenon. Water released during decomposition of these flame retardants dilutes the gases and forms water vapour barriers that retains the oxygen, which otherwise ignites and supports the flame [8-10, 39-40]. The highest LOI value were achieved for ATH formulations meaning, sample 8 with stearic acid as lubricant (25.8 %) and sample 9 with calcium stearate as lubricant (26 %). It is also important to point out that PVC composites, with calcium stearate as lubricant, presented superior LOI values compared to the ones with stearic acid.

Contact angle (CA)

Contact angle (CA) is the most important parameter used to quantify the wettability of solid surfaces. In this respect, the need to preserve or enhance the hydrophobicity of the composite materials, which are used in out-door applications, refers to stability in moist environment and preservation of flame-retardant properties on long term [41]. Conventionally, the contact angle describes the
behaviour of a liquid droplet on a solid surface in air, and is defined as the angle between the tangent at the three phase point and the solid surface. The degree of wetting when a solid and liquid interact is indicated by wettability studies, which usually involve the measurement of contact angles as the primary data. High wettability corresponds to small contact angles (<<90°), while large contact angles (>>90°) correspond to low wettability or hydrophobicity [42-45].

Table 7 reflects a viable agreement of contact angles with the samples composition in terms of presence or absence of reinforcement material and lubricant. It was observed that the samples without reinforcing or flame retardant material (samples 2 and 3) attained high contact angles, due to the low or complete lack of OH groups (fig. 7). When introducing calcium carbonate (samples 4, 5 and 6, 7 respectively), the wettability increases for both series of composites due to the water adsorption capacity of the filler, while the contact angles of PVC composites with ATH and HMH (samples 8 and 10, and 9 and 11, respectively) are slight higher.

The explanation suggested by some authors is related to the hydrophilic surface of ATH and HMH, which becomes partly hydrophobic after lubrication and compounding in the PVC matrix, leading to lower surface energy [42-45].

Table 7 CONTACT ANGLE VALUES FOR PVC COMPOSITES

| Samples with Stearic acid | Contact angle (°) | Sample with Calcium stearate | Contact angle (°) |
|---------------------------|------------------|-----------------------------|------------------|
| 2                         | 112.90           | 3                           | 114.38           |
| 4                         | 90.87            | 6                           | 98.95            |
| 5                         | 104.17           | 7                           | 108.70           |
| 8                         | 106.39           | 9                           | 114.37           |
| 10                        | 105.37           | 11                          | 114.31           |

Fig. 7. Contact angle values for the two PVC series of composites: with stearic acid (samples 2, 4, 5, 8 and 10) and with calcium stearate (samples 3, 6, 7, 9 and 11)

The final step to establish an optimum formulation for flame-retardant flexible polyvinylchloride composites was scanning electronic microscopy of PVC composites. As it can be seen in figure 8, calcium stearate used as lubricant led to more homogenous composites and the presence of macropores was reduced, compared to the series of samples with stearic acid. The composites with CaCO\(_3\) of 1 µm (samples 5 and 7) presented macropores and agglomerations.

The samples with HMH (sample 10 with stearic acid and sample 11 with calcium stearate) were rather homogenous, presenting some zones with smooth

Scanning electronic microscopy (SEM)

The final step to establish an optimum formulation for flame-retardant flexible polyvinylchloride composites was scanning electronic microscopy of PVC composites. As it can be seen in figure 8, calcium stearate used as lubricant led to more homogenous composites and the presence of macropores was reduced, compared to the series of samples with stearic acid. The composites with CaCO\(_3\) of 1 µm (samples 5 and 7) presented macropores and agglomerations.

The samples with HMH (sample 10 with stearic acid and sample 11 with calcium stearate) were rather homogenous, presenting some zones with smooth
surfaces. Nevertheless, the samples with ATH (sample 8 with stearic acid and sample 9 with calcium stearate) presented a more discrete surface arrangement with a better dispersion of ATH, particularly for sample 9.

**Conclusions**

New formulations of PVC composites with potential applications in the industry for electrical wires and cables were prepared and characterized. Calcium carbonate with particle size of 1 and 2 μm, which is a cheap version of flame-retardant fillers (also called reinforcing agents), has proven to be proper for decreasing the Tg value of PVC and to contribute to the optimization of mechanical properties referring to elasticity, elongation at break and tensile strength of composites. Yet, the flammability of thereof was high compared to that of control samples. HMM-based composites presented medium properties, comparable to those of calcium carbonate, but exhibited higher fire retardation. ATH, seemed to be the most proper filler for preparing flame-retardant PVC composites, leading to low Tg values of PVC, good mechanical properties, flexibility and high limiting oxygen indexes (comparable to that of HMM-based PVC Composites). Comparing the composites from the point of view of the two used lubricants, stearic acid and calcium stearate, it can be concluded that calcium stearate ensures better compatibility of components, enhanced hydrophobicity and leads to more homogenous PVC composites, as confirmed by SEM analysis and also other studies [46-49].

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