Study of Chemical Composition and Some Material Properties for a Hybrid Resin Based on Dammar

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Abstract: The chemical composition of a hybrid resin made of natural Dammar resin and epoxy resin (along with the related hardener), in proportion of volume of 60% and 40%, respectively, was studied based on EDS (Energy Dispersive Spectroscopy) analysis. The presence of epoxy resin was necessary to generate a rapid polymerization process. The material details of the hybrid resin were investigated using SEM (Scanning Electron Microscopy). A high-resolution image of a hybrid resin sample was obtained. The weight loss of the hybrid resin at certain temperature values using thermo gravimetric analysis was measured. In addition, the surface roughness of the hybrid resin was characterized using two methods. By the first method, the classical one, the roughness was measured using two profile parameters and five analysis lines, namely the functional dependence between the roughness height and the gray level of the analyzed surface image. By the second method, the average roughness method, the surface texture of a hybrid resin specimen was measured and the roughness profile was obtained as the difference between the primary profile and the middle line of the filter determined with the Gaussian profile filter. Finally, several possible areas of use of this type of resin have been proposed based on the properties obtained.

Keywords: hybrid resin, chemical composition, thermal stability, roughness

1. Introduction

Natural resins are a mixture of volatile aromatic chemicals and are produced by plants or animals. Natural resins are not soluble in water, but are easily soluble in oil, alcohol and partially in gasoline. In combination with certain organic solvents (turpentine, rosin) these resins form solutions that can be used as coating varnishes [1-4]. A disadvantage for natural lakes is that they cannot form thick resins [1-2]. One solution to eliminate this drawback is to use hybrid resins obtained by combining several constituents, of which at least one is organic and at least one is synthetic [5].

One of the most used natural resins is Dammar. It is harvested from trees of the Dipterocarpaceae family in India and East Asia. The soil where these trees grow influences the appearance of macro and microelements in the chemical structure of Dammar resin, i.e. additional elements may appear such as: iron, silicon, zirconium, cadmium and so on.

Dammar films, in the form of varnish, are used to protect paintings [6]. The effect of long exposure to acetic acid of paintings coated with Dammar-based varnish, which generates an accelerated degradation of them, was investigated in the paper [7-8].

Using a multi analytical approach based on thermo gravimetric analysis coupled with infrared spectroscopy (TGA-FTIR), in the paper [9] was clarified the nature of the macromolecular components of Dammar resin.

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The structure and chemical composition of Dammar resin have been studied in detail in [10] and [11]. Dammar in combination with silicone was used to obtain "ecological" types of binders. In the work [12] a study was carried out on the optimization of the quantities of Dammar and silicone in the composition of "ecological" binders, used to cover aluminum panels, so that they have good properties for impact, hardness, traction and adhesion. The influence of adding Dammar to improving the rigidity, modulus of elasticity and hardness of "ecological" binders was studied in [13]. Paper [14] aimed to expand knowledge about the bio microbiological deterioration of Dammar-based lakes in works of art subject to unsuitable protective conditions, especially long exposures to high levels of relative humidity, suggesting that some microorganisms favor processes of natural damage to the films of Lake Dammar.

The effect of polymethylmethacrylate (PMMA) on the physical properties of Dammar for the application of coating paints was investigated in [15]. It has been found that Dammar when mixed with PMMA can be used for cold coating of rolled steel.

The interdependence between the surface roughness and its image is based, in principle, on the functional dependence between the roughness height and the gray level of the surface image, in the sense that the higher parts of the roughness correspond to the higher intensity pixels [16-17].

2. Materials and methods
2.1. Manufacture of test specimens
It should be noted that the temperature of the medium in which the casting was performed was 21-23°C. All specimens were cut 10 days after pouring the plate.

A hybrid resin plate was poured, in which Dammar was used in a volume proportion of 60%. The difference up to 100% consisted of Resoltech 1050 type epoxy resin together with the corresponding Resoltech 1055 type hardener. The synthetic component (epoxy resin and hardener) were required to generate rapid activation points of the polymerization process.

The volume ratio of the combination of the two synthetic components Resoltech 1050 / Resoltech 1055 was 7/3 (according to the manufacturer's instructions). The manufacturer in [18] gives the thermo-mechanical properties of Resoltech 1050 epoxy resins together with the Resoltech 1055 hardener.

A set of twenty specimens marked with Dammar 1.1-20 was cut from the cast plate. The dimensions of the specimens were length 250 mm, width 25 mm and thickness 6.2 mm. The density of the specimens was between 1.04-1.05 g/cm³. Figure 1 shows a specimen from this set.

Figure 1. Hybrid resin specimen with a volume ratio of 60% Dammar

2.2. Methods of analysis and apparatus used in the tests
The chemical composition of a hybrid resin sample was determined based on EDS analysis. This analysis was performed using a PHENOM PURE PRO X scanning electron microscope [19], with integrated EDS system, with conventional cathode for microscopic study of the structures and surface of different materials, with the possibility of determining the chemical composition and phases in the structure. The possibility of magnification is between 80-130000 times. Among the types of materials for which are used are composite materials with polymer or metal matrix.

The roughness analysis for a hybrid resin sample was performed with the PHENOM PURE PRO X microscope because it is equipped with lighting through objective In addition, the Taylor Hobson Surtronic 3+ surface roughness tester [20] was used to obtain a numerical roughness assessment for a hybrid resin specimen, by what is known as the Ra method (roughness average method).
SEM analysis was performed with an electron microscope - Hitachi model S3400N / type II [21], with the following characteristics:

- SE image resolution: minimum 3.0 nm at 30 kV (x100,000, WD = 5mm, high vacuum mode); minimum 10 nm at 3 kV (x30,000, WD = 5mm, high vacuum mode);
- BSE image resolution: minimum 4.0 nm at 30 kV (x60,000, WD = 5 mm, low vacuum mode);
- magnification: between x5 - x300,000;
- acceleration voltage: 0.3 kV - 30 kV;
- electron gun voltage: in steps with own field and with fixed field with continuously adjustable voltage;
- beam alignment: two-stage electromagnetic deflector;
- objective aperture: 4 holes with click-stop system with a diameter of 30, 50, 80 and 150 microns;
- image displacement: minimum ± 50 microns at WD = 10 mm.

EDS and SEM analyzes were performed in accordance with the provisions of [22].

Based on the thermo gravimetric analysis (TGA), the thermal stability of a hybrid resin sample was determined. We used a Thermo gravimetric Q50 Analyzer connected to a desktop (which can heat a sample up to 1000ºC) and software for experimental analysis of Universal Analysis 2000 data, which offers weight loss at certain temperature values. TGA analysis was performed according to the provisions of [23].

The functional groups and chemical composition of the hybrid resin was determined based on FTIR (Fourier Transformed Infrared Spectroscopy) analysis.

FTIR analysis was performed using a portable IdentifyIR system (equipped with ChemAssist software) [24, 25] have been complied with. The sample size used was less than 5 µL or 100 μm. In addition, the equipment was presented with the Aldrich / Smiths ATR spectral libraries. All the technical characteristics of this system can be found on the manufacturer's website [26].

3. Results and discussions

Figure 2 shows the diagram of the EDS analysis of the chemical composition for a hybrid resin sample, obtained at an intensity of 15 keV.

Figure 2. Diagram of the EDS analysis of the chemical composition for a hybrid resin sample, obtained at an intensity of 15 keV

Based on the EDS analysis, Table 1 shows the chemical composition of a hybrid resin sample. The weight concentration (Atomic Concentration) and the number of atoms in each element express this composition.

To determine the number of atoms in the weight concentration of an element the formula was used:

\[
\text{Atoms Numbers} = \frac{\text{Weight concentration} \cdot \text{Avogadro constant}}{\text{Atomic weight}}
\]
A total number of atoms equal to $\approx 45 \times 10^{23}$ (obtained as the sum of all numbers on the number of atoms column) was used.

**Table 1. Chemical composition of a hybrid resin sample**

| Element Number | Element Symbol | Element Name | Atomic Weight | Atomic Conc. Wt. [%] | Weight Conc. At. [%] | Atoms Numbers |
|----------------|----------------|--------------|---------------|----------------------|---------------------|---------------|
| 6              | C              | Carbon       | 12            | 77.74                | 69.26               | $34.7569 \times 10^{23}$ |
| 8              | O              | Oxygen       | 16            | 20.47                | 24.30               | $9.1459 \times 10^{23}$ |
| 49             | In             | Indium       | 115           | 0.40                 | 3.37                | $0.1764 \times 10^{23}$ |
| 17             | Cl             | Chlorine     | 35.5          | 0.59                 | 1.56                | $0.2646 \times 10^{23}$ |
| 11             | Na             | Sodium       | 23            | 0.59                 | 1.01                | $0.2644 \times 10^{23}$ |
| 16             | S              | Sulphur      | 32            | 0.21                 | 0.50                | $0.0940 \times 10^{23}$ |

The material details of the hybrid resin were investigated using SEM. Figure 3 shows a high-resolution image of a sample of this type of resin.

For the thermo gravimetric analysis, a hybrid resin sample was heated to a chosen temperature of 800ºC (because it was considered that no significant changes will occur above this value) and the weight change was determined.

After the experiment, the air cooled in 20 min. The gas used in this experiment was nitrogen with a purity of 5.0. A sample with a mass of 0.63 mg was taken from the samples made from hybrid resin. This sample was loaded into the platinum vessel at the appliance (the sample was crushed before loading).

The weight loss of the hybrid resin, obtained from the experimental data, compared to the temperature is shown in Figure 4 a, and the thermo gravimetric curve is shown in Figure 4 b.

**Figure 3. SEM analysis for a hybrid resin sample**

**Figure 4.**

a) Deriv. weight change of hybrid resin at various temperatures;  
b) The weight loss (%) of hybrid resin at various temperatures
From the previous figure, five distinct stages of weight loss of the hybrid resin can be observed, as follows:

- weight loss of 0.5534% ∙ min/ºC from 49.85ºC to 150ºC, with a peak value up to 99.44 ºC; this case occurs mainly due to the evaporation of the parts of water that make up the resin;
- a weight loss of 0.1749% ∙ min/ºC from 210.31ºC to 300ºC, with a maximum value of 250.49ºC; residual values of epoxy resins and Dammar remain;
- a weight loss of 3,185% ∙ min/ºC from 309.67ºC to approximately 450ºC, with a maximum value at 363.090 C; residual values of epoxy resins and Dammar remain;
- a weight loss of 0.8127% ∙ min/ºC from about 535.20ºC to about 785ºC, with a peak value of 649.44ºC;
- no final changes are observed from 785 to 800ºC.

Figure 5a shows the roughness analysis for a hybrid resin sample. The roughness value was calculated on the profile, using five lines of analysis. Figure 5b shows the roughness analysis of the same resin monster using the functional dependence between the roughness height and the gray level of the surface images.

The roughness analysis was performed in accordance with [27, 28].

To characterize the roughness, it is used two profile parameters $R_z$ and $R_a$. The parameter $R_z$ represents the average of all the maximum differences between the maximum and the minimum point at the profile within the evaluation length, and $R_a$ is the arithmetic mean of the absolute deviations of the roughness profile from the midline.

![Figure 5. Roughness analysis for a hybrid resin sample: a) on the 5 lines of analysis; b) using the functional dependence between the height of the roughness and the gray level of the surface images](image)

The values of the roughness profile parameters are presented in Table 2.

| No. | $R_z$       | $R_a$    |
|-----|-------------|----------|
| 1   | 9.53 μm     | 1.75 μm  |
| 2   | 9.39 μm     | 2.14 μm  |
| 3   | 10.10 μm    | 1.66 μm  |
| 4   | 5.44 μm     | 1.61 μm  |
| 5   | 13.28 μm    | 2.23 μm  |

The surface texture of a hybrid resin specimen was measured using the medium roughness method.
The roughness profile was obtained as the difference between the primary profile and the middle line of the filter determined with the Gaussian profile filter. Figure 6 shows the tester used by this method of measuring surface roughness.

Figure 6. Taylor Hobson Surtronic 3+ surface roughness tester

Figure 7 shows the general roughness profile, and Figure 8 shows the roughness profile determined using the Gaussian filter. This type of filter was introduced in [29] (for profiles) and with its help the mean line can be determined in surface metrology. Its weighting function is given by the expression:

$$h(t) = \frac{1}{\alpha \lambda_c} e^{-\left(\frac{t}{\alpha \lambda_c}\right)^2}, \quad \text{where: } \alpha = 0.4697,$$

$$\lambda_c$$ is the cut-off wavelength of the filter (in units of $$t$$).

If it used $$x(t)$$ for the position of the unfiltered primary profile, $$m(t)$$ for the Gaussian filtered mean line and $$r(t)$$ for the roughness profile, then:

$$m(t) = x(t) \ast h(t) \quad \text{and} \quad r(t) = x(t) - m(t).$$

The amount of data removed by the Gaussian filter is smaller than that of filters obtained by other methods and thus the roughness profile is more accurate.

Figure 7. General roughness profile

$$\begin{align*}
\int_{-\infty}^{+\infty} x(\tau) h(t - \tau) d\tau.
\end{align*}$$
Figure 8. Roughness profile determined using a Gaussian filter

The data obtained are $R_a=1.76\mu m$, $R_t=13.10\mu m$ and $R_z=9.83\mu m$.

Figure 9 shows the FTIR analysis diagram for a hybrid resin sample.

Figure 9. FTIR analysis of a hybrid resin sample

Based on the FTIR analysis of the spectrum of a hybrid resin sample, the following structural identification was obtained:

- $3371\text{ cm}^{-1}$ indicates the presence of N-H groups in the structure of an aliphatic primary amine;
- $3050-2800\text{ cm}^{-1}$ there is the stretching of the aromatic and aliphatic C-H bonds;
- $1608\text{ cm}^{-1}$ the extension of the C = C bonds of the aromatic rings occurs;
- $1507\text{ cm}^{-1}$ there is an extension of the aromatic C-C bonds;
- $1454\text{ cm}^{-1}$ indicates the presence of methylene (CH2) groups;
- the additional presence of the $1376\text{ cm}^{-1}$ peak is caused by a methyl (CH3) group;
- $1033\text{ cm}^{-1}$ the C-O-C stretch of the ethers occurs;
- $824\text{ cm}^{-1}$ the C-O-C extension of the oxirane groups occurs;
- $763\text{ cm}^{-1}$ the oscillating movement (rocking) of CH2 occurs.

4. Conclusions

From the diagram of the EDS analysis of a hybrid resin sample, it can be seen that oxygen has a significant share in its composition. This fact, is also certified by the SEM analysis, which shows the existence of a large number of air gaps in the structure of the hybrid resin. The explanation may be that the curing time and implicitly the duration of the polymerization process is much longer for the hybrid resin than for the epoxy resin and thus some of the air bubbles formed during the curing process remain trapped inside the resin.
The presence of air bubbles in the structure of the hybrid resin increases the roughness of the analyzed surface. This can be seen in Figure 5b, by the extended presence of shades of gray.

The values of the roughness profile parameters, presented in Table 2 and functional dependence between the roughness height and the gray level of the surface images for the hybrid resin sample (Figure 5b), indicate that we have an average roughness.

From the two experimental ways of determining the roughness parameters, it is observed that there are insignificant differences between their values.

IR spectroscopy is an extremely effective method for determining the presence or absence of a wide variety of functional groups in a molecule. IR spectra can be used to identify the molecule by recording the spectrum for an unknown and comparing this information with the library or a database of spectra of known compounds.

Two functional groups can be distinguished from the FTIR analysis diagram of a hybrid resin sample:
1. cumulative Ketene type double bonds (CH2 = C = O), which are substances with a combined carbonyl and carbon-carbon double bonds;
2. Nitrogen double bond, known as the Nitrite ion (NO2) and it contains nitrogen in a relatively unstable oxidation state.

From the analysis of the thermo gravimetric curve (Figure 4b) it can be seen that the highest percentage weight loss of the analyzed hybrid resin sample occurs in the temperature range between 300 and 450ºC.

When we obtained this hybrid resin, we thought about the possibility of using a matrix in the manufacture of "environmentally friendly" composite materials, by reinforcing them with various types of natural reinforcers.

These composite materials can be used in:
- the field of civil and industrial constructions, making paneling and parquet boards;
- the medical field, for the manufacture of elements to replace gypsum dressings using the immobilization of fractures of the tibia, arm, forearm, etc.

In a subsequent study we will investigate, how the roughness and composition of material will influence the wetting properties by measuring the contact angle (ex. water).

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Manuscript received: 2.11.2020