Colored Polymeric Microcomposites with Minimal Environmental Impact

MIRCEA RUSE1*, SANDU PERETZ2, ANCA ANGELA ATHANASIU1, LOTI CORNELIA OPROIU1, CATALIN FILIPESCU1, RALUCA SENIN2

1 National Research and Development Institute for Chemistry and Petrochemistry – ICECHIM, 202 Spl. Independentei, 060021, Bucharest, Romania
2 Institute of Physical Chemistry I. Murgulescu – ICF (Academy), Bucharest, 202 Spl. Independentei, 060021, Bucharest, Romania

The paper presents the research carried out by our team on achieving amphiphilic polymeric microcomposites, colored in ruby and violet shades for natural fibers dyeing (cotton, wool). The colored polymeric microcomposites were characterized in terms of their morphological (by optical microscopy and SEM analysis) and physicochemical performances (by FT-IR and UV-Vis spectroscopy, colorimetric analysis). The colored microcapsules were tested by studying the controlled release of the encapsulated dyes in water, to pH variation. This study was necessary to the coloristic characterization of colored polymeric microcomposites to determine the optimal microcapsules dye concentration and dyeing concentration of natural fibers.

Keywords: amphiphilic, polymeric, microcapsules, colorimetric analysis

Microencapsulation is a process in which tiny particles or droplets are surrounded by a shell (membrane) to obtain small capsules with useful properties. Most of the microcapsules have a diameter from a few μ to several mm. The definition was expanded and includes most medicines. There have been encapsulated a series of food ingredients and fragrances, in particular [1-3]. Further the technique of microencapsulation is expected to be applied to the organic and inorganic dyes and pigments class. The purpose of microencapsulation is the isolation of the active substance from the environment, of reactive and metal complex dye in our case, to defend them of the harmful effect of atmospheric oxygen and UV radiations and controlling the speed with which the dye leaves the microcapsule, such as: controlled release of the studied dyes from the microcapsule in water [4, 5]. The paper presents the research carried out by our team on achieving amphiphilic polymeric microcomposites, colored in ruby and violet shades for natural fibers dyeing (cotton, wool). It was developed the laboratory microencapsulation technology for reactive and metal complex dyes in a biodegradable natural polymer matrix (calcium alginate) [6-8]. As coloring materials were used reactive and metal-complex dyes, synthesized by our team, previously analysed in terms of physicochemical (purity, concentration, FT-IR and UV-Vis spectral parameters) and coloristical performances (hue, intensity, dyeing concentration, resistance to light, water, wet and dry friction, acid and alkaline perspiration, solubility and degree of fleet exhaustion). Chemical formulas of the used dyes to achieve microencapsulation are I), (II) and (III). Their performances are comparable to known foreign brands (BASF, ICI, Ciba). The colored polymeric microcomposites were characterized in terms of morphology (optical microscopy, SEM analysis) and physicochemical performances (FT-IR and UV-Vis spectroscopy, colorimetric analysis). The colored microcapsules were tested by studying the controlled release of the encapsulated dyes in water, to pH variation. This study is still necessary to the coloristic characterization of colored polymeric microcomposites, to determine the optimal microcapsules dye concentration and dyeing concentration of natural fibers.

(I) Wool Ruby Metal Complex Dye

(II) Cotton Violet Reactive Dye

(III) Cotton Ruby Reactive Dye

* email: mircearuse1@yahoo.com; Phone: +40 0726787868

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fibers. The technologies used in obtaining and applying of colored microcomposites are green and are made with minimal specific consumption and according to currently existing environmental information, both raw materials (reactive and metal-complex dyes, anionic polymer, precipitating agent) and end products show reduced eco-toxicological risk.

**Experimental part**

**Materials and equipment**

Microencapsulation agent: sodium alginate (NaAlg); Precipitation agent: calcium chloride (CaCl₂); Colouring material: Ruby reactive dye, Violet reactive dye and Ruby metal-complex dye.

IR spectra were measured with a FT-IR – Jasco 6300 spectrophotometer, using KBr pellets UV-Vis spectra were performed in distilled water with a UV/Vis/NIR spectrophotometer, in 190-2500 nm range. Were used cells with 1 cm pathlength, working in the range dye concentrations of 0.3 - 2.3 \times 10^{-2} g/L in distilled water. UV-Vis JASCO V 570 spectrophotometer, in 190-2500 nm range. Optical Microscope – MAGNUM: trinocular head, with fluorescence system and magnification x 20, x40, x100; fine focusing knob, min division: 0.002 mm. SEM - Scanning Electron Microscope with point resolution: 1.2 nm with focused ion beam (FIB).

**Synthesis**

Microcomposites synthesis was made into a capsule with large area with d = 26 cm, which contained a colored solution of 6.2% CaCl₂ conc., equipped with an anchor stirrer and three dropping funnels in which was placed a solution of 1.7% NaAlg conc. NaAlg solution was dropped at ambient temperature within approx. 9 h. The optimum amount of dye taken into work, was considered in relation to the used amount of NaAlg, reaching a value of approx. 0.43%, corresponding to an optimal absorption of the dye inside the microcapsule and a weak colored solution load to the initial white polymeric microcomposites. Their relative concentration to: Ruby reactive dye, 52 g/L (cotton) dyeing. By colorimetric analysis was determined its relative concentration of 70% having λ\text{max} = 515 nm. Monochlorotriazine reactive azo dyes, in liquid form, used to obtain colored microcomposites in ruby and violet shades were synthesized by our team and belong to Cibanicron (Ciba Geigy) dyes type for cellulose fibers (cotton) dyeing. By colorimetric analysis was determined their relative concentration to: Ruby reactive dye, 52 g/L with λ\text{max} = 545 nm; λ\text{2} = 528 nm; Violet reactive dye, 49 g/L with λ\text{max} = 543 nm. Sodium alginate (NaAlg) was purchased from Aldrich company, with 99% conc., 2% solution at 25°C and viscosity is η = 250 cp. Colored polymeric microcapsules were morphologically (by optical microscopy, SEM analysis) and physicochemical (by FT-IR and UV-Vis spectroscopy, colorimetric analysis) characterized. Also tests have been carried out to determine the controlled-release rate of the dye from the microcapsules in water by measuring the dye concentration variation of the solution, in time.

**Results and discussions**

**Morphological characterization**

Optical microscopy (microcapsules size determination) CaAlg, microcapsules were white. Optical microscopy determinations were made on lyophilized samples. We found that white microcapsules are in the quasi-spherical form with a size of 500μm. For samples containing dye is found that colored microcapsules are larger, around 800-1200μm.

SEM (scanning electron microscopy)

White microcapsules have a rough structure with pores in the scales form. This rough surface and their pores indicate that they are suitable for absorption of organic substances, such as dyes (fig. 1). Colored polymeric microcomposites containing reactive or metal-complex dyes are found to have a completely different look compared to the initial white polymeric microcomposites. Their surface is more compact and shows micro-cracks (fig. 2 and fig. 3).

**FT-IR Spectroscopy**

Determining of the interaction type between the reactive dye and white polymeric matrix

FT-IR analysis for ruby and violet reactive dyes adsorbed by white microcapsules, leads to the fact that the maximum absorption bands for the white polymeric matrix, are located on the next frequency values: 2931 cm⁻¹ for \text{Ruby Microcapsules} and 2935 cm⁻¹ for \text{Violet Microcapsules}.
OH groups; - 1593 cm\(^{-1}\) and 1420 cm\(^{-1}\) for COO\(^{-1}\) groups; 1011 cm\(^{-1}\) for -CO- groups. It finds that the maximum absorption peak at 3231 cm\(^{-1}\) of –OH groups of white polymeric matrix, is displaced and affected at 3246 cm\(^{-1}\) in the case of colored microcapsules containing reactive dye, due to formation of weak intermolecular hydrogen bonds and dipole – dipole interactions type, between the -OH groups and -N = N- groups and/or aromatic residues of the reactive dye. We see a movement and an affectation of maximum absorption peaks from 1593 cm\(^{-1}\) and 1420 cm\(^{-1}\) for COO\(^{-1}\) groups of white polymeric matrix to lower wavenumber 1590 cm\(^{-1}\) and 1417 cm\(^{-1}\) in the case of colored microcapsules, due to weak electrostatic interactions with – N=N- groups (having streching vibration to 1586 cm\(^{-1}\) and 1481 cm\(^{-1}\)) of the reactive dye. We can assume that we are dealing with physical forces, which favors a time controlled-release in water of the reactive dye, at ambient temperature.

Determining of the interaction type between the ruby metal complex dye and white polymeric matrix

FT-IR analysis of ruby metal complex dye adsorbed by white microcapsules leads to the fact that the maximum absorption bands for white polymeric matrix and ruby metal complex dye are located on the next frequency values: 3231 cm\(^{-1}\) for –OH groups; 1593 cm\(^{-1}\) and 1420 cm\(^{-1}\) for COO\(^{-1}\) groups; 1011 cm\(^{-1}\) for -CO- groups; 1430 cm\(^{-1}\) for –NO\(_2\) groups, in the ruby dye. It finds that the maximum absorption peak at 1593 cm\(^{-1}\) of the white polymeric matrix for –COO\(^{-1}\) groups is displaced at 1417 cm\(^{-1}\), in the case of colored polymeric microcapsules. It finds that the maximum absorption peak at 1430 cm\(^{-1}\) for polar –NO\(_2\) groups of ruby metal complex dye is displaced at 1453 cm\(^{-1}\), in the case of colored polymeric microcapsules. We can assume that we are dealing with electrostatic interactions between hydrophile groups of polymer matrix and polar – NO\(_2\) groups of ruby metal complex dye. So the ruby metal complex dye remains embedded in the polymeric matrix and it can not be released from the microcapsules at ambient temperature.

UV-Vis spectroscopy (colorimetric analysis)

Relative concentration determination of the reactive dye within the microcapsules

It was performed colorimetric analysis to dyestuffs solutions samples to determine the relative concentrations of reactive dyes in colored microcapsules in ruby and violet shades. For this we’ve determined the maximum absorbance in UV-Vis absorption spectra, for each reactive dye, to:

- standard samples of known concentration: 52 g/L for ruby and 49 g/L for violet dye;
- corresponding filtered samples, resulted after colored microcomposites isolation;
- samples of the primary extract, resulting after the desorption in water of the dye contained by the microcapsules.

It was concluded that the two reactive dyes respect Lambert-Beer law and from the calculations were obtained the following results: Dye concentration in the colored polymeric microcomposites: \(\text{Conc}_R = 0.29\%\); \(\text{Conc}_V = 0.15\%\); Microencapsulation yield: \(\text{Y}_R = 59\%\); \(\text{Y}_V = 32.3\%\).
Time controlled-release of the Violet Reactive Dye
In the test were used 8 microcapsules (dry) containing reactive dye, which was placed in a spectrophotometer cuvette, containing 5 mL of distilled water, at ambient temperature.

Controlled release was followed by measuring the dye concentration variations in time according to the absorbency values at different times by UV-Vis colorimetry analysis: $\lambda_{max} = 543$ nm.

Test conditions:
- pH = 6.5 with continued stirring, to approx. 70 rpm at ambient temperature.
- pH = 8, with continued stirring, to approx. 70 rpm at ambient temperature.

It finds that we have a phenomenon of rapid release of the dye from microcapsules after 45 min. of microcapsules introduction in water (with continuous stirring), the absorbance is 54.6% at pH = 8 to 27.8% in the case of pH = 6.5 (fig. 8). After approx. 225 min, we found that the absorbance value remains virtually constant. We have done research both in weak acid medium at pH = 6.5 and at slightly alkaline medium at pH = 8 and were noted: at pH = 6.5, was released approx. 94.3% of the initial dye amount, the microcapsules were bleached, becoming almost transparent, so it was released most of the encapsulated dye; at pH = 8 was released approx. 96.8% of the initial dye amount.

At a slightly alkaline medium, it is found that the reactive dye release occurred more quickly.

Time controlled-release of the Ruby Metal Complex Dye
In the test were used 8 microcapsules (dry) containing the ruby metal complex dye, that was placed in a spectrophotometer cuvette, containing 5 mL of distilled water, at ambient temperature.

Controlled release was followed by UV-Vis measurements at $\lambda_{max} = 515$ nm. Test conditions:
1) pH = 6.5 and 2) pH = 8, with continued stirring, to approx. 70 rpm at ambient temperature.

After 225 min of placing the colored microcapsules in water (stirring continuously), we found that absorbance of the dye is almost constant at both pH = 6.5 and pH = 8, but reaches a max. of 2-3% (fig. 9). In a slightly alk. medium is found that the dye release is very low, in time.

<table>
<thead>
<tr>
<th>Reactive Dye</th>
<th>Max. abs. $\lambda_{max}$ (nm)</th>
<th>Standard Absorbance</th>
<th>Standard Conc. (g/L)</th>
<th>Extract Absorbance</th>
<th>Extract Conc. (g/L)</th>
<th>Filtrate Absorbance</th>
<th>Filtrate Conc. (g/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ruby</td>
<td>522, 544</td>
<td>2.3187</td>
<td>1.733*10$^{-2}$</td>
<td>1.3131</td>
<td>0.9814*10$^{-2}$</td>
<td>0.4206</td>
<td>0.3144*10$^{-2}$</td>
</tr>
<tr>
<td>Violet</td>
<td>543</td>
<td>0.9360</td>
<td>1.633*10$^{-2}$</td>
<td>0.2907</td>
<td>0.507*10$^{-2}$</td>
<td>0.3193</td>
<td>0.557*10$^{-2}$</td>
</tr>
</tbody>
</table>

Table 1
SPECTRAL PARAMETERS OF THE INVESTIGATED REACTIVE DYES

![](image1)

Fig.6 - UV-Vis spectrum of Ruby reactive dye samples: standard, extract, filtrate

Fig.7 - UV-Vis spectrum of Violet reactive dye samples: standard, filtrate, extract

Coloristic characterisation of the colored polymeric microcomposites
At cotton dyeing were used both colored microcomposites (in ruby and violet shades) and standard unencapsulated reactive dyes. Before dyeing, the colored microcomposites were suspended in water, maintained 24 h with stirring, for reactive dye release. We worked according to the known dyeing technology, at 40-60°C temperature, 1:40 fleet report and finally the decolored
microcomposites were isolated. Further, was used the colored filtrate to dyeing. Were made corresponding dyeings to three different dyeing concentrations: 1%; 1.5%; 2%. Have resulted coloristic dyeing intensities close to those of standard reactive dyes. Coloristic resistences of dyed samples are similar to those given by standard dyes: 1) Xe light resistance: 4-5; 2) Concentration of dyeing: 1-2%; 3) Water resistance: 4-5 / 4-5 / 4-5; 4) Resistance to acid sweat: 4-5 / 4-5 / 4-5; 5) Wet friction resistance: 5; 6) Dry friction resistance: 5.

Conclusions

We have obtained two new colored materials in ruby and violet shades in form of polymeric microcomposites used to dye cellulose fibers (cotton);

We have obtained one new colored material in ruby shade in form of polymeric microcomposites, which cannot be used to dye protein support (wool, leather);

We developed a modernized and ecological laboratory technology of conditioning of investigated coloring materials by microencapsulation in a biodegradable natural polymer;

From FT-IR spectra we can assume that we are dealing only with absorption based on physical forces between polymeric matrix and reactive dyes, which favors a time controlled-release of the dye contained by the microcapsules;

From FT-IR spectra we can assume that we are dealing with electrostatic interaction between hydrofile groups of polymeric matrix and polar –NO₂ groups of ruby metal complex dye, which prevents the time controlled-release of the metal complex dye, contained by the microcapsules.

From the isolation data of colored polymeric microcomposites and colorimetric analysis in the UV-Vis field it was determined the optimal dye concentration in microcapsules, Conc_R = 0.29% and Conc_V = 0.15%, to ensure the optimal dyeing concentration to the cellulose textile support (cotton) at 1, 1.5, 2% values. It was demonstrated the functionality and utility of microencapsulation technology, with microencapsulation efficiency between 30-60%.

Acknowledgements: The work was funded by Executive Agency for Higher Education, Research, Development and Innovation (UEFISCDI) under The Ministry of Education from Romania. We thank Mr. dr. Valentin Raditoiu to colorimetric analysis and absorption spectra in UV-Vis and Mr. Marian Deaconu to residual water discoloration method made.

References

1. FERRES J. M. R., SERRABASA E. P., LIRO M., (1998), ES 2,112,150 (B01, J13/10);
2. CHOKYUN R., SANCHEZ D. R., (1988), US 4,744,933 (A61, K9/16);
3. SHEVCHENKO A.V., BIRJUKOVA L.A.; (1999), RU 2,132,224 (C08, J9/32);
4. MING XU, (2007), WO (C09, B67/00B4F);
5. TAKASHI O., HIRONORD S., KAZUO S., (1998) JP 63252543 (A) (A61, K8/11C);

Manuscript received: 7.11.2016