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FT-IR studies of the polymeric binder BioCo1 with modified biopolymer – part I

Abstract
This paper presents structural studies of polymeric water soluble compositions consisting of acrylic derivatives/modified biopolymer. The research FT-IR was focused on novel polymer composition poly(acrylic acid)/modified starch (PAA/CMS) designed for application in the foundry industry – a novel group of polymeric BioCo binders. It was found, that the structure changes occur within groups: carboxyl present in poly(acrylic acid) and carboxyl and hydroxyl groups contained in carboxymethyl starch (vibrations: $\nu_s$-C=O, $\nu_s$-COO$^-$ and $\delta$-C–O–H).

Keywords: polymeric composition, polymeric binder, moulding sands, FT-IR

1. Investigations

The polymer composition poly(acrylic acid)/modified starch (PAA/CMS), due to its physical and chemical properties found its application as a new binding agent of moulding
sands (binder BioCo1) [1]. The main loose component of a moulding sand is sand grains of the determined mineralogical, chemical and granular composition (the most often high-silica sand). A component providing a determined mechanical strength and able to bind sand grains is the binding agent (binder). The selection of the proper binder cross-linking moulding sands components, to obtain the binding power sufficient to retain the mould shape – during pouring it with liquid metal – is essential in the foundry technology [2]. Thus, a binder is crucial in the foundry process since it has to bring about obtaining of dimensional castings. In addition, the proper selection of the cross-linking factor has a significant influence on a binder bounding power in moulding sands. Therefore, when designing new polymer binders or modifying already known ones, one should penetrate into its structure and recognize its binding mechanism [3–5]. The results concerning the structure analysis (FT-IR) of the designed binder BioCo1 (polymeric composition PAA/CMS) are presented in the paper.

Structural investigations on the determination of the moulding sand hardening process, i.e. binding agent cross-linking process in the system: binding agent-mineral matrix in relation to its technological properties, will constitute the second part of the publication.

1.1. Materials

The following samples BioCo1 were used as a new polymer binding agent in the form of a two-component polymer composition formed by mixing in water a synthetic polymer (poly(acrylic acid), PAA, BASF) and a modified biopolymer (modified starch: carboxymethyl starch, CMS, Xenon, DS 0.2) in the weight ratio 1:1 was used in the performed investigations. After mixing components the mixture in the form of the water homogeneous polymer composition of pH = 2.5, viscosity 850 mPa·s and 78% of water content, was obtained.

1.2. Structural investigations

Spectroscopic FT-IR tests were carried out by means of the spectrometer Digilab Excalibur FTS 3000 Mx with detector DTGS, electrically cooled. This spectrometer is equipped with two attachments: ATR with ZnSe crystal for multiple reflection and the transmission attachment. Transmission spectra of the investigated samples were recorded in a range: 4000–400 cm⁻¹, at the resolution of 4 cm⁻¹. Number of scans was selected in dependence on the level of the obtained signal (within 32–64 scans). The correction of the obtained spectra base line was performed by means of the Merlin (FTS 3000) software.

1.3. The microscopic studies

Then the formed samples of aqueous composition polymer of PAA/CMS were dried by vacuum and after converting to microscopic examination were conducted of the solid
form using ultra-high resolution scanning electron microscope NOVA Nano SEM 200 (by FEI EUROPE COMPANY) cooperating with the analyzer EDAX EDS company.

2. The obtained results

Thermodynamic solubility of polymers is their ability to formation of stable, homogeneous polymer solutions in the whole range of the components quantitative fraction. The chemical affinity of a solvent to a substance being dissolved and their similar polarity constitutes the condition for solving. The solvent role consists in overcoming the van der Waals forces occurring in between individual macroparticles [6].

Mixtures of two or more polymers by their solving in the same solvent can form complex systems of a single phase or of multiphases (the so-called multi-component polymer compositions) in dependence on the structure and properties of polymers and solvents, their concentrations and mixing temperature.

Taking into consideration the general theory of mixing polymers, the verification of the polymer composition was performed taking into account physical and chemical properties of polymer components (molecular mass, polarity, viscosity, pH, concentration) and their chemical affinity to the solvent (water) [7].

As the first component of the composition the synthetic homopolymer from the group of acrylic polymers: poly(acrylic acid) (Fig. 1) was selected.

![Fig. 1. Structure of poly(acrylic acid)](image)

The second component of the composition was the natural polymer: starch. In order to improve the starch miscibility and solubility in water its chemical modification was performed. The obtained modified carboxymethyl starch (Fig. 2) exhibits better polarity, which means better solubility in water and better miscibility with hydrophilic polymers [8].

![Fig. 2. General formula of carboxymethyl starch](image)
Spectroscopic FT-IR investigations for the initial components (polymers) and after their mixing were performed, which enabled to determine structure changes occurring during the polymer composition formation (Fig. 3).

Three IR spectra were recorded and the occurrence of a broad band – in the wave number range 3700–3000 cm⁻¹ – corresponding to stretching vibrations of the hydroxyl group (band of free –OH group [water] and hydrogen bond).

The spectrum analysis of the poly(acrylic acid) sample confirmed the presence of groups characteristic for carboxylic acids (1713 cm⁻¹) corresponding to the stretching vibrations of carbonyl group: –C=O (Fig. 4, spectrum 1). The band corresponding to stretching vibrations of C–O bond and deforming vibrations of C–O–H was observed in a range of 1246 cm⁻¹. In the case of the carboxymethyl starch spectrum the band characteristic for the asymmetric vibrations of the carboxylate COO⁻ group and deforming vibrations of C–O–H appears in the wave number range: 1632 cm⁻¹ (Fig. 3, spectrum 2).

During polymer mixing the absorption band of the poly(acrylic acid) from 1713 cm⁻¹ (Fig. 3, spectrum 1) corresponding to asymmetric vibrations of C=O shifts in the direction of higher wave numbers: 1718 cm⁻¹ (Fig. 4, spectrum 3), while bands at 1632 cm⁻¹ and 1246 cm⁻¹ characteristic for vibrations: C–OH and C–O in carboxymethyl starch (Fig. 3, spectrum 2) are also shifted in the direction of higher wave numbers: 1637 cm⁻¹ and 1258 cm⁻¹ (Fig. 3, spectrum 3). These bands can be the result of overlapping vibrations related to the formation – during the composition formation – of new ester type bonds with a participation of carbonyl, carboxyl and hydroxyl groups.
Characteristic absorption bands observed in the IR spectra of the investigated system are given in Table 1.

Table 1. Characteristic bands in the IR spectra of polymers and the polymer composition [9–11]

<table>
<thead>
<tr>
<th>PAA [cm⁻¹]</th>
<th>CMS [cm⁻¹]</th>
<th>PAA/CMS [cm⁻¹]</th>
<th>Assignment</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>3439</td>
<td>3463</td>
<td>3457</td>
<td>ν-OH</td>
<td>ν-OH...O-H O–H O–H O=C</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Band of free OH group (water) Hydrogen bond</td>
<td></td>
</tr>
<tr>
<td>2959</td>
<td>–</td>
<td>2962</td>
<td>ν-C−H</td>
<td>Bands of stretching vibrations asymmetric and symmetric</td>
</tr>
<tr>
<td>1713</td>
<td>–</td>
<td>1718</td>
<td>ν s-C=O</td>
<td>Stretching vibrations of carbonyl group in carboxylic acids</td>
</tr>
<tr>
<td>–</td>
<td>1632</td>
<td>1637</td>
<td>δ-C−O−H ν s-COO−</td>
<td>Deforming vibrations Asymmetric vibration</td>
</tr>
<tr>
<td>1450</td>
<td>–</td>
<td>1450</td>
<td>δ (CH₂) n</td>
<td>Shearing symmetric vibrations</td>
</tr>
<tr>
<td>1407</td>
<td>–</td>
<td>–</td>
<td>δ (CH₂) n</td>
<td>Shearing symmetric vibrations</td>
</tr>
<tr>
<td>1246</td>
<td>–</td>
<td>1258</td>
<td>C-O−H C−O</td>
<td>Deforming vibrations Stretching vibrations in a plane</td>
</tr>
<tr>
<td>–</td>
<td>1023</td>
<td>–</td>
<td>C−CH₂</td>
<td>Stretching vibrations</td>
</tr>
</tbody>
</table>

On the grounds of the obtained IR spectra it was found, that there is a possibility of occurrence of the reaction of poly(acrylic acid) and carboxymethyl starch – in the water environment – which reveals itself by changes in a position and intensity of absorption bands in a range of wave numbers: 1700–1600 cm⁻¹ (Fig. 3). Thus, the structure changes occur within groups: carboxyl (−COOH), present in poly(acrylic acid) and carboxyl and hydroxyl groups (−COO⁻ and −OH), contained in carboxymethyl starch.

On the basis of microscopic images, it was found that the composition of PAA/CMS, after drying, are shown separated fractions (starch granules). The obtained results confirm that the polymers PAA and CMS due to the presence of starch grains form a colloidal system (Fig. 4).

Fig. 4. SEM picture of a polymer composition PAA / CMS.
3. Conclusions

On the grounds of the analysis of the IR spectra the structural analysis of the polymer composition PAA/CMS (BioCo1 binder). Spectroscopic investigations for the initial components and after their mixing was performed, which enabled to determine structure changes occurring during the polymer composition formation. On the grounds of the obtained IR spectra it was found, that there is a possibility of occurrence of the reaction in the water environment of poly(acrylic acid) and carboxymethyl starch.

Acknowledgements

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References