EDYTA HEBDA, JAN PIELICHOWSKI*

POLY(ASPARTIC ACID) – BASED CATALYSTS FOR THE OXIDATION REACTIONS

ZASTOSOWANIE KATALIZATORÓW NA BAZIE POLI(KWASU ASPARAGINOWEGO) W REAKCJACH UTLENIANIA

Abstract

In the current work we describe the synthesis of novel poly(D, L-aspartic acid)-supported cobalt or copper(II) catalysts for oxidation reactions. Oxidation reactions of some compounds were carried out on synthesized catalysts at atmospheric pressure in the presence of molecular oxygen. As the main products epoxides and ketone were obtained with very high yield and selectivity.

Keywords: poly(aspartic acid), polymer catalysts, oxidation reactions

Streszczenie

W artykule przedstawiono syntezę nowych katalizatorów kobaltowych i miedziowych na bazie poli(D, L-kwasu asparaginowego) z przeznaczeniem do reakcji utleniania. Reakcje oksydacji prowadzono pod ciśnieniem atmosferycznym i w obecności tlenu cząsteczkowego. Jako główne produkty otrzymano epoksydy i ketony z wysoką wydajnością i selektywnością.

Słowa kluczowe: poli(kwas asparaginowy), katalizatory polimerowe, utlenianie

1. Introduction

The aerobic epoxidation of alkenes with a transition metal catalyst has been widely studied over the past decade. One of the well-known methods of alkene epoxidation in homogeneous system is the Mukaiyama procedure, where substrate is epoxidized using a transition metal complex as catalyst, molecular oxygen as oxidant and an aliphatic aldehyde as co-reactant. Transition metal complexes, such as Schiff’s base complex, are also suitable for the aerobic epoxidation of alkenes with a co-reacting aldehyde. For example, cobalt(II) Schiff’s bases give good results; however, these catalysts are rather not selective for epoxidation.

It has been reported in literature that the epoxidation of alkenes by various oxidants can proceed very efficiently when a polymer-supported catalyst is used. It offers several advantages in the preparation procedures. The polymer is very stable (in the meaning of thermal properties) even in the oxidative atmosphere. As an example, a polyaniline-supported cobalt(II) catalysts were prepared first by Pielichowski and Iqbal and used for the aerobic epoxidation of alkenes. For example, a polyaniline-supported catalyst has been used for epoxidation of trans-stilbene with very high yield and selectivity [1–5].

In the last decade biodegradable polymers have deserved a special attention as their decomposition products are non-toxic. To this group of polymers – due to its advantages – belongs e.g. poly (aspartic acid) (here: PAA), which is a biologically inert polyamino acid. Poly (aspartic acid) has been recently very intensely examined over the possibility of the utilization of PAA and its derivatives in medicine and agriculture [6–8]. Because poly(D, L-aspartic acid) have functional groups to coordinate, thus, in this paper, we report on novel poly(aspartic acid) supported cobalt(II) or copper(II) catalysts, which were tested in the oxidation processes of hydrocarbons – it is hoped that the catalysts chemically linked with polymeric support will be stable and able to be separated easily after the reaction.

These catalysts, based on poly(aspartic acid), have been obtained and tested in trans-stilbene, indene and 1-decene oxidation reactions.

2. Experimental

In the Department of Chemistry and Technology of Polymers, Cracow University of Technology has used new, original method synthesis of poly(aspartic acid) under microwave irradiation. The application of the microwave irradiation has permitted shortening of polymerisation time to several minutes, increase in efficiency and elimination of the catalyst – this fact is very important in synthesis compounds, which are used for the preparation of catalysts.

The general method for poly(aspartic acid) sodium salt (PAA-Na) synthesis is illustrated in Figure 1.

In our experiments the catalyst were prepared by a three-step method.

2.1. Polycondensation of aspartic acid

The synthesis of cyclic PAA was carried out in a microwave reactor “Milestone” of 1000 W capacity. Polycondensation of cyclic PAA was realized at the temperature range
from 176°C to 230°C, carbonate propylene was used as solution. Water was removed from the system by an azeotropic distillation under normal pressure. Cyclic PAA was precipitated with methanol, washed in water and dried. The reaction efficiency was above 93%.

![Chemical structure](image)

**Fig. 1. Synthesis of PAA sodium salt**

Rys. 1. Synteza soli sodowej poli(kwasu asparaginowego)

### 2.2. Hydrolysis of cyclic forms of PAA

PAA sodium salt has been obtained during the reaction of hydrolysis of cyclic forms of PAA, water solutions of sodium. The process of hydrolysis was carried out in the solution with pH = 8.5 to 13.0 [4].

Analysis **FT-IR** (KBr): characteristic absorption band of stretching vibrations of C = O group is about 1610 and 1660 cm\(^{-1}\), strong absorption band of OH stretching vibrations is in the range of 3400–3500 cm\(^{-1}\).

Analysis **\(^1\)H NMR** (D\(_2\)O): the methane proton is seen as two resonances at about 4.7 and 4.5 ppm, the methylene protons are observed as three resonances at about 2.8, 2.7 and 2.55 ppm.

### 2.3. Synthesis of complex of PAA sodium salt with cobalt(II) or copper(II) acetate

The complex of PAA sodium salt with cobalt(II) or copper(II) acetate was obtained by mixing of 1.00g of PAA-Na with the corresponding amount of metal acetate.

In this way were received two catalysts:
- PAA-Na + cobalt acetate – PAA-Co,
- PAA-Na + copper acetate – PAA-Cu.

The two catalysts – PAA-Co and PAA-Cu – were examined by SEM/EDXS. The specimen analyses were carried out using a HITACHI S-4700 field emission scanning electron microscope (SEM) equipped with a NORAN Vantage energy dispersive X-ray spectrometer (EDS). A focused electron beam was restored across the sample surface. The
Fig. 2. SEM images of a) PAA-Co, b) PAA-Cu and metals distribution on PAA cobalt (a) and copper (b) respectively

Rys. 2. Obraz SEM i rozkład metalu odpowiednio w katalizatorach PAA-Co (a) i PAA-Cu (b)

Fig. 3. Positive secondary ion mass spectrum (a) and negative secondary ion mass spectrum (b) of PAA-Na

Rys. 3. Widma (+) i (−) masowe PAA-Na
secondary or backscattered electrons produced were detected and used to map the surface topography and compositional contrast based on the density differences. The X-rays emitted when the electron beam struck a sample yielded information about the chemical composition of the sample surface (EDS). The analysis depth of EDS varied between 0.3 and 4.5 μm depending on the material analyzed and the primary beam energy. Prior to the analysis the catalyst samples were coated with conductive carbon films to a thickness of 20–30 nm. The surface images of the catalysts are shown in Fig. 2. It can be seen that these heterogeneous catalysts created agglomerates with the size of the order of 20–40 μm and smaller, of about 5–10 μm. These agglomerates consist of many smaller spherical grains in case of PAA-Co and flaky-lamellar ones for PAA-Cu.

The application of EDS spectroscopy coupled with SEM let us test the surface distribution of chemical elements, for example metals, in the form of a surface map – Fig. 2. We observed that metal was spaced exactly on the catalyst surface. We can suggest that a chemical reaction took place between the polymer and cobalt or copper(II) salt, and only one compound was obtained.

The received catalysts were tested for the presence of metals using time-of-flight secondary ion mass spectrometry (ToF-SIMS). ToF-SIMS measurements were performed using a PHI TRIFT 2100 time-of-flight secondary ion mass spectrometer equipped with gallium liquid metal ion gun (LMIG). This instrument allows spectroscopy, for the characterization of surface chemical composition. The system uses a pulsed primary ion beam to desorbs and ionize species from a sample surface [9–13].

The results for PAA-Na and its cobalt and copper derivatives, assigned as PAA-Co and PAA-Cu respectively, are reported here. Positive mass spectrum of PAA-Na is dominated by sodium (Fig. 3), which reflects low ionization potential of this metal under the measurement conditions. The secondary fragment ions, despite their low intensities, support the structure of PAA-Na. The characteristic fragments of PAA-Na ions present in positive and negative mass spectra are shown in Tab. 1. The lines are of low intensities and 93% of the total positive ion emission originates from sodium.

<table>
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<th>Table 1</th>
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### Characteristic fragments of PAA-Na ions present in positive and negative mass spectra

<table>
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<tr>
<th>Characteristic Fragments of Ions</th>
<th>Positive</th>
<th>Negative</th>
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<tbody>
<tr>
<td>Formula</td>
<td>m/z, amu</td>
<td>Formula</td>
</tr>
<tr>
<td>[CH$_2$N]$^+$</td>
<td>28.02</td>
<td>[CN]$^-$</td>
</tr>
<tr>
<td>[CH$_2$N]$^+$</td>
<td>30.03</td>
<td>[CNO]$^-$</td>
</tr>
<tr>
<td>[CONH$_2$]$^+$</td>
<td>44.01</td>
<td>[COOH]$^-$</td>
</tr>
<tr>
<td>[C$_3$H$_4$N]$^+$</td>
<td>44.05</td>
<td>[H$_2$O]$^-$</td>
</tr>
<tr>
<td>[CH$_2$O$_2$]$^+$</td>
<td>46.01</td>
<td>[C$_3$H$_2$O$_2$]$^-$</td>
</tr>
<tr>
<td>[C$_2$H$_2$O$_2$]$^-$</td>
<td>71.01</td>
<td>[C$_2$H$_2$NO$_2$]$^-$</td>
</tr>
</tbody>
</table>
The process of ion exchange resulted in the replacement of mobile sodium ions with cobalt ion, which is shown in Fig. 4. The spectra clearly indicate that most of the sodium was substituted during the process. The spectrum for PAA-Cu is similar for PAA-Co.

Figure 4 shows the presence of the [COCo]+ fragment, which suggests incorporation of cobalt into the precursor’s structure (not just a copper oxide).

The characteristic fragments of PAA-Co and PAA-Cu ions present in positive mass spectra are shown in Tab. 2.

The presence of [COCo]+ and [COCu]+ fragment ions provides evidence for the chemical interaction between the catalyst precursor (PAA-Na) and both the cobalt and copper cations. It means that these metals are not simply supported as their oxides on PAA-Na but they are chemically incorporated into it.
2.4. The oxidation of hydrocarbons in presence of received catalysts

In order to check activity of the prepared catalysts several reactions were carried out. In a typical procedure catalyst (30 mg), acetonitrile (30 mL), and substrate (2 mmol) were placed in a reactor. The mixture was heated up to 60ºC, stirred with magnetic stirrer and bubbled with molecular oxygen. 2-methylpropanal was added (6 mmol) to the reaction mixture. The oxidation yields the following hydrocarbons: trans-stilbene, 1-decene and indene.

3. Results and discussion

The results of catalytic oxidation of organic compounds are collected in Tab. 3. Selectivity of the main products was given and the reaction time was shown in brackets.

<table>
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<tr>
<th>Catalysts</th>
<th>Products</th>
<th>Selectivity [%]</th>
<th>(time [h])</th>
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<tbody>
<tr>
<td>PAA-Co</td>
<td>epoxystilbene</td>
<td>85 (0.5)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>epoxydecene</td>
<td>62 (30)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>indene-2-one</td>
<td>98 (1)</td>
<td></td>
</tr>
<tr>
<td>PAA-Cu</td>
<td>epoxystilbene</td>
<td>91 (0.5)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>epoxydecene</td>
<td>56 (32)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>indene-2-one</td>
<td>97 (1)</td>
<td></td>
</tr>
</tbody>
</table>

Trans-stilbene and 1-decene give epoxides, but indene gives ketone (1,3-dihydro-2H-inden-2-on) with very high selectivity. The oxidation reaction of 1-decene needs longer time than the oxidation of trans-stilbene and indene. Under these conditions 1-decene gives epoxide with very high selectivity. No changes in yield were observed when the same reactions were carried out longer. The oxidation of double bonds proceeds easier when the double bond is activated by the presence of another group. In comparison, polymers-supported manganese porphyrin gave similar results as a catalyst in the oxidation reactions, but longer reaction time was necessary [14].

4. Conclusions

In conclusion, we have described the synthesis of poly(aspartic acid) – based catalysts, which are very effective in oxidizing of alkenes under mild conditions. The main advantages of these catalysts are their efficiency, selectivity and simplicity of separation of the heterogeneous catalysts from the reaction medium by filtration. The oxidation reactions occur in a relatively short time. Generally, poly(aspartic acid) can be considered as a new group of macromolecular oxidation catalysts, with advantageous properties.
References