Heterogeneous Catalyst Based on Phosphine-Containing Organic Polymer for Hydroformylation of Octene-1

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Abstract—A phosphine-containing porous polymer (TPDB), and a rhodium catalyst based on it, were synthesized. Proceeding from the relevant TEM, NMR, IR spectroscopy, and XPS data obtained, it was assumed that both the Rh(I) complexes and the Rh(0) nanoparticles that are stabilized in the polymer pores act as the catalyst’s active phase. The size of the nanoparticles (2–6 nm) is consistent with that of the polymer pores. The catalyst was tested in a model reaction of octene-1 hydroformylation at 2.0 MPa (CO : H2 = 1 : 1) in the temperature range of 60–120°C. It was demonstrated that the catalyst can be used repeatedly, and its activity remains constant from the third cycle of its use (aldehyde yield about 30% for 5 h, n/iso 2.5).

Keywords: hydroformylation, porous organic polymer, rhodium nanoparticles, rhodium complexes, catalysis, heterogeneous catalyst

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Hydroformylation of unsaturated compounds (i.e., catalytic interaction of olefins with syngas at elevated pressures and temperatures) is one of the most important organic synthesis reactions to obtain aldehydes with various structures [1, 2]. Industrial-scale olefin hydroformylation processes utilize homogeneous cobalt-based (140–200°C, 5.0–30.0 MPa) and rhodium-based (85–130°C, 1.5–5.0 MPa) catalysts, either in the form of hydridocarbonyls or modified with donor ligands, most often alkyl- or aryl substituted phosphines. However, even if the required levels of substrate conversion rate and reaction selectivity are achieved, the efficiency of the catalyst separation/recirculation step remains to be improved [3, 4]. The conventional extraction and distillation methods are very complex in terms of engineering design. Moreover, their implementation often leads to the deactivation and loss of the expensive catalyst as a result of the decomposition of the metal complex and the oxidation or destruction of the ligand; as a consequence, a regeneration step is required [5–7]. An alternative method was developed in Germany in 1984 [8, 9]. This method involves propylene hydroformylation under two-phase catalysis conditions using a water-soluble catalyst. However, this approach has proved to be applicable only to the hydroformylation of lower olefins because olefins with a chain length above C6 significantly reduce the reaction rate due to their low water solubility [10–13].

The challenge of catalyst separation from the reaction products, regardless of the type of substrate being processed, can be overcome by using heterogeneous catalysts. Considerable experience has been gained in this area. Various approaches to the fixation of the active metal on the surface, in the pores, or inside the structure of the support have been systematized in a number of review papers [14–17]. The main disadvantage of heterogeneous catalysts for liquid-phase hydroformylation is the leaching of active metal particles into the reaction medium, which causes their rapid deactivation and inability to operate continuously. The leaching is associated with a number of causes, in particular the formation of organic-soluble metal hydridocarbonyls, namely HCo(CO)4 or
HRh(CO)$_4$, under the effects of syngas and elevated temperatures [18, 19].

A very promising approach to the creation of heterogeneous hydroformylation catalysts involves the preparation of rhodium catalysts from porous organic polymers. These supports combine the high porosity and high density of ligand fragments with a good resistance to organic media and elevated temperatures. The provision of specific coordination sites in the polymer’s structure offers good opportunities for their use in selective synthesis of linear aldehydes or in asymmetric synthesis applications. The available information on the utilization of catalysts based on porous organic polymers in hydroformylation has been systematized in a review paper [20]. The general synthesis method for these polymers comprises the preparation of monomers (specifically, arylphosphine or phosphite derivatives that contain vinyl fragments) followed by their polymerization. The resultant porous materials were treated with rhodium precursors, which made it possible to prepare heterogeneous catalyst samples that exhibited high levels of stability, activity, chemoselectivity, and regioselectivity in the hydroformylation of a number of model substrates [21].

The purpose of this study was to prepare a heterogeneous rhodium catalyst based on a phosphine-containing porous organic polymer (TPDB) and to examine its activity in the liquid-phase hydroformylation of octene-1 as a model substrate.

**EXPERIMENTAL**

We used FeCl$_3$ (Sigma-Aldrich, 99.9%), $\alpha,\alpha'$-dibromo-$p$-xylene (Sigma-Aldrich, 98%), triphenylphosphine (Sigma-Aldrich, 99%), phenylsilane (Sigma-Aldrich, 96.5%), and octene-1 (Sigma-Aldrich). The following solvents were used: 1,2-dichloroethane (Acros Organics, 99.8%), methanol (Acros Organics, 99%), and toluene (Komponent-Reaktiv, CP grade). The solvents were purified according to standard procedures. Rhodium acetylacetonate dicarbonyl, or Rh(acac)(CO)$_2$, was analyzed by the method described in [22].

The starting materials, the polymeric supports and catalysts based on them, and the reaction products were analyzed by gas-liquid chromatography (GLC), spectrophotometry, IR spectroscopy, solid-state NMR spectroscopy, X-ray photoelectron spectroscopy (XPS), atomic absorption spectroscopy, transmission electron microscopy (TEM), and low-temperature nitrogen adsorption/desorption.

The GLC was performed on a Khromos chromatograph equipped with a flame ionization detector and a 30 m capillary column (DB-5 phase, programmed heating from 60 to 235°C at a rate of 10°C/min, helium as a carrier gas). The phosphorus and silicon contents were determined on a Micromeritics Gemini VII 2390 (V1.02t) analyzer according to a standard procedure. Prior to testing, the samples were degassed at 120°C and 3×10$^{-4}$ MPa for 12 h. The adsorption and desorption isotherms were recorded at 77 K. The porosity was calculated using a standard software package. The specific surface area was determined using a Brunauer–Emmett–Teller (BET) model at a relative partial pressure of $p/p_0 = 0.2$. The total pore volume and pore size distribution were evaluated using a Barrett–Joyner–Halenda (BJH) model at a relative partial pressure of $p/p_0 = 0.95$.

The phosphine-containing porous polymer (TPDB) was synthesized by the alkylation of triphenylphosphine with $\alpha,\alpha'$-dibromo-$p$-xylene followed by reduction with phenylsilane (Scheme 1).

A TP(O)DB polymer was prepared (the initial step) similarly to the method described in [23]. However, we used a different reagent mixing sequence to avoid triphenylphosphine quaternization, which took place under the original procedure and made it unnecessarily difficult to implement. Our synthesis procedure was as follows. First, 2.66 g of $\alpha,\alpha'$-dibromo-$p$-xylene was dissolved in 30 mL of 1,2-dichloroethane in a 250 mL three-neck flask equipped with a magnetic stirrer and a reflux condenser with a chloro-calcium tube. Then 6.48 g of anhydrous FeCl$_3$ was added. Next, the contents of the flask were cooled in an ice bath, and a solution of 2.64 g of triphenylphosphine in
20 mL of 1,2-dichloroethane was added. After 10 min, the flask was placed into a silicone oil bath, heated to 45°C, and stirred for 5 h. Then the flask was further heated to 80°C, and the reaction was conducted for 48 h. All these steps were carried out in an argon stream.

The product was washed with 1,2-dichloroethane and methanol, and vacuum-dried.

To prepare a phosphine-containing TPDB support, the TP(O)DB polymer was reduced with phenylsilane. For this purpose, 1.5 g of TP(O)DB was placed into
a 100 mL argon-filled round-bottom flask equipped with a magnetic stirrer, and 4.5 mL of phenylsilane and 10 mL of toluene were added. The flask, sealed tightly with a plug fitted with a Teflon gasket, was placed into a silicone oil bath. The reaction was carried out at 98°C under vigorous stirring for 48 h. The resultant precipitate was dried on a rotary evaporator at 60°C for 40 min. The product weighed 2.5 g.

The Rh/TPDB catalyst was synthesized in an inert atmosphere. 300 mg of the polymeric support and 18 mg of Rh(acac)(CO)$_2$ were placed into a 25 mL argon-filled flask equipped with a magnetic stirrer; then 4 mL of toluene was poured. The flask was tightly closed with a glass stopper. The mixture was stirred for 5 h, and the resultant precipitate was centrifuged, washed with toluene (3 times, 5 mL each), and vacuum-dried for 1 h. 243 mg of a light brown Rh/TPDB powder was ultimately obtained.

The hydroformylation reaction in the presence of the heterogeneous Rh/TPDB catalyst was carried out in a 25 mL steel autoclave equipped with a thermostat and a magnetic stirrer. The calculated amounts of catalyst, octene-1, and toluene were placed into the argon-purged autoclave. The reaction occurred under continuous stirring, at pressure of 2.0 MPa (CO : H$_2$ = 1 : 1) and a specific temperature (60–120°C), for a time period individually determined for each specific experiment. Then the autoclave was cooled to room temperature and depressurized. The catalyst was centrifuged from the reaction mixture and washed with toluene, before being reused in the reaction.

**RESULTS AND DISCUSSION**

The low-temperature nitrogen adsorption/desorption of the TP(O)DB sample demonstrates a high specific surface area of the polymer synthesized (590.5 m$^2$/g, at a pore volume of 0.9 cm$^3$/g and an average pore size of 6 nm). The adsorption/desorption isotherms (Fig. 1) display a hysteresis loop, which is indicative of a mesoporous structure. On the other hand, the high nitrogen absorption at low pressures, the further gradual increase in this parameter with an increase in $p/p_0$, and the abrupt rise in the high pressure region ($p/p_0 = 0.8–1.0$), prove the presence of both mesopores and micropores in the polymer.

The TP(O)DB polymer obtained was measured by solid-state $^1$H, $^{13}$C, and $^{31}$P NMR. The $^1$H NMR spectrum (Fig. 2a) contains overlapping signals at 3.0 and 6.5 ppm, which correspond to the hydrogen atoms of the polymer’s aromatic and aliphatic fragments, respectively. The carbon spectrum (Fig. 2c) displays a broad signal at 131 ppm, corresponding to the carbon atoms of the aromatic fragments, as well as signals between 10–60 ppm, typical of the aliphatic carbon atoms connected to the aromatic system. In the $^{31}$P NMR spectrum (Fig. 2b) of the TP(O)DB sample, the intense signal at 23 ppm indicates an almost complete oxidation of the triphenylphosphine groups during the synthesis. The interaction with phenylsilane is followed by a partial reduction of phosphine oxide fragments, as indicated by a signal at –5 ppm that appears in the TPDB spectrum (Fig. 2d). However, the phenylsilane treatment contaminates the sample with silicon compounds, as evidenced both by the elemental analysis data (2.48% Si) and by a significant reduction in the surface area (up to 0.03 m$^2$/g). The TEM data show the random porous structure of the support prepared (Figs. 3a, 3b).

Rhodium was applied by interaction between the TPDB polymer and the rhodium acetylacetonate dicarbonyl in toluene. The rhodium content in the Rh/TPDB, quantified by flame atomic absorption spectroscopy, is 1.2%.

Rh3d core level X-ray photoelectron spectrum of the Rh/TPDB sample (Fig. 3d) exhibits signals at 307.3 eV (5/2) and 312.1 eV (3/2), typical of rhodium in the zero oxidation degree [24] and, thus, indicative of Rh(0) nanoparticles in the sample. These nanoparticles likely result from rhodium reduction by phenylsilane during the catalyst synthesis, where the rhodium is retained in the support’s pores. At the same time, the 308.5 eV (5/2) and 313.2 eV (3/2) signals, as well as the 309.2 eV (5/2) and 314.0 eV (3/2) signals, are typical of Rh(+1) complexes [25]. The Rh/TPDB micrographs
The TP(O)DB’s IR spectrum contains absorption bands typical of the C–H (912, 870, 742, 721, and 692 cm\(^{-1}\)) and –C=C– (1437–1612 cm\(^{-1}\)) bonds of the aromatic ring, as well as P–C=C (1683 cm\(^{-1}\)) and Ph\(_3\)P=O (1186 cm\(^{-1}\)) [26]. The TPDB spectrum displays the appearance of absorption bands attributable to the valence vibrations of Si–H (2158 cm\(^{-1}\)) and Si–O–Si (1000–1100 and 814 cm\(^{-1}\)) [26, 27], which indicate the contamination of the sample with silicon compounds. The application of rhodium to the polymer results in the appearance of absorption bands near 2061 cm\(^{-1}\) and near 1993 cm\(^{-1}\) (corresponding to the Rh–CO vibrations), as well as signals in the range of 1635–1652 cm\(^{-1}\) (evidence of an acac\(^{-}\) anion in the coordination environment of rhodium) [28]. Rhodium nanoparticles of different sizes (2 to 6 nm, depending on the size of the pores in which they are located) are fixed in the polymer matrix. The polymer structure also contains rhodium complexes in which Rh(+1) is bonded to triphenylphosphine ligand, carbon monoxide, and an acac\(^{-}\) ion.

The catalytic activity of the resultant heterogeneous Rh/TPDB catalyst was examined in a model reaction of octene-1 hydroformylation. Although the catalyst demonstrated sufficiently high activity in the first catalytic experiment (Cycle 1), its activity noticeably declined during Cycle 2. This is most probably associated with the partial leaching, into the solution, of the rhodium acetylacetonate dicarbonyl that was sorbed but not bonded chemically to the polymer fragments. As a consequence, the reaction occurred not only on the active sites of the heterogeneous catalyst, but also on HRh(CO)\(_4\) (rhodium hydridocarbonyl) formed from Rh(acac)(CO)\(_2\) in the syngas atmosphere, thus leading to a higher aldehyde yield than in the subsequent cycles. It should be noted that the heterogeneous Rh/TPDB catalyst can be used repeatedly, and its activity remains nearly constant after Cycle 2 (Table 1).

The temperature effects on the hydroformylation in the presence of Rh/TPDB were evaluated based on the catalyst reuse data (Fig. 4a). At 60°C, the aldehyde conversion rate does not exceed 34%, the octene-1 conversion reaches about 50%, and the \(n/iso\) ratio is 1.8. However, raising the temperature to 80°C boosts the aldehyde yield to 57% (with \(n/iso = 1.4\) and 98% octene-1 conversion). At 100°C, despite a slight increase in the aldehyde yield (to 61%), the \(n/iso\) ratio declines to 0.6 as
a result of the accelerated iso-octene hydroformylation. The highest n/iso ratio (with yields being similar in all the cases compared) was achieved at 80°C. This temperature also corresponds to the lowest amount of by-products.

The evaluation of the substrate conversion rate and of the accumulation of the products of octene-1 hydroformylation in the presence of Rh/TPDB (Fig. 4b), revealed that the octene-1 conversion into aldehydes starts about 80 min after the experiment initiation. This is probably explained by the stabilization of diffusion processes in the system and the generation of the catalyst’s active form. The substrate undergoes an almost complete (98%) conversion within 300 min, where the initial reaction step mostly promotes the substrate isomerization. The iso-octene content in the reaction mixture reaches its maximum after about 200 min from the experiment initiation, followed by a decline as a result of hydroformylation into iso-aldehydes.

Thus, based on the data on the catalytic activity and the physicochemical properties of the Rh/TPDB catalyst,

Table 1. Hydroformylation of octene-1 in presence of Rh/TPDB

| Cycle no. | Conversion, % | Yield of iso-octenes, % | Yield of aldehydes, % | n/iso (yield ratio of n-aldehydes to iso-aldehydes) |
|-----------|---------------|-------------------------|-----------------------|-----------------------------------------------|
| 1         | 99            | 25                      | 74                    | 0.8                                           |
| 2         | 99            | 42                      | 57                    | 1.4                                           |
| 3         | 72            | 44                      | 28                    | 2.5                                           |
| 4         | 77            | 48                      | 29                    | 2.6                                           |
| 5         | 73            | 45                      | 28                    | 2.6                                           |
| 6         | 64            | 39                      | 25                    | 2.6                                           |
an assumption of its most likely structure was suggested (Fig. 5).

Rhodium nanoparticles can be stabilized by the $\pi$-system of aromatic polymer fragments and by heteroatoms and, thus, firmly fixed in the pores. In this case, the substrate may gain access to the “facets” of nanoparticles protruding onto the polymer surface, and these facets rather effectively catalyze isomerization of octene-1 (a side reaction). This assumption is further confirmed by the findings of the catalytic experiments. First, the sustaining catalytic activity in the reuse cycles proves that the active sites are firmly fixed on the support. Second, the substrate-related parameters (conversion rate, isomerization degree, and relative isomerization rate), the aldehyde yield, and the $n/\text{iso}$ ratio deviate, to some extent, from the values that would be expected if the active sites consisted only of rhodium-phosphine complexes. Although $n/\text{iso}$ reaches 2.5–2.6 during the reuse of Rh/TPDB, the final product mixture (5 h, aldehyde yield ca. 30%) contains a significant amount of $\text{iso}$-octenes. This leads us to expect the $n/\text{iso}$ ratio to be about 0.8–1.0 when unsaturated compounds are fully converted into aldehydes. In our case, $\text{iso}$-octenes mostly remain in the reaction mixture, and the low rate of their hydroformylation may stem from local steric hindrances near the active sites of the catalyst. The catalytic conversions mostly occur on nanoparticles protruding onto the catalyst surface. At 80°C, the turnover frequency (TOF) of octene-1 hydroformylation, on the basis of aldehydes produced, is about 100 h$^{-1}$ (at 50% substrate conversion). This is quite consistent with the values of the same parameter reported for other heterogeneous catalysts [18]. The relatively low TOF of the reaction in the presence of Rh/TPDB can be explained by the fact that the rhodium concentration on the surface is lower than its total concentration in the catalyst sample.

Thus, the catalyst under study exhibits sufficiently high activity in hydroformylation of octene-1, and the findings obtained from the examination of its structure suggest that its catalytic effect is associated both with the
Rh(+1) complexes fixed on the polymer surface and with the Rh(0) nanoparticles stabilized in the pores.

AUTHOR CONTRIBUTION

D.N. Gorbunov, Yu.S. Kardasheva, and E.A. Karakhanov developed the concept of the study, adapted a method for synthesis of polymer support to the research task, and proposed the method for heterogeneous catalyst synthesis and a set of methods for its physicochemical examination. M.V. Nenasheva and R.P. Matsukevich synthesized and isolated the polymer and polymer-based catalyst samples. M.V. Terenina and D.N. Gorbunov analyzed the data obtained by IR spectroscopy, solid-state NMR, XPS, atomic absorption spectroscopy, TEM, and low-temperature nitrogen adsorption/desorption. M.V. Nenasheva and R.P. Matsukevich carried out the catalytic experiments on hydroformylation of octene-1 in pressure reactors, interpreted the chromatography data on the product mixtures, and compiled the tables summarizing the main findings. D.N. Gorbunov and Yu.S. Kardasheva wrote the text part of the paper. All the authors participated in the discussion of the results. M.V. Nenasheva and D.N. Gorbunov prepared the graphic part for publication.

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CONFlict OF INTEREST

E.A. Karakhanov, a co-author, is an editorial board member for the Journal Nanogeterogeneny Kataliz (Nanoheterogeneous Catalysis). The other co-authors declare no conflict of interest requiring disclosure in this article.

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