Preparation and characterization of a supported system of Ni2P/Ni12P5 nanoparticles and their use as active phase in chemoselective hydrogenation of acetophenone

To cite this article before publication: Dolly C Costa et al 2018 Nanotechnology in press https://doi.org/10.1088/1361-6528/aab3a8

Manuscript version: Accepted Manuscript

Accepted Manuscript is “the version of the article accepted for publication including all changes made as a result of the peer review process, and which may also include the addition to the article by IOP Publishing of a header, an article ID, a cover sheet and/or an ‘Accepted Manuscript’ watermark, but excluding any other editing, typesetting or other changes made by IOP Publishing and/or its licensors”

This Accepted Manuscript is © 2018 IOP Publishing Ltd.

During the embargo period (the 12 month period from the publication of the Version of Record of this article), the Accepted Manuscript is fully protected by copyright and cannot be reused or reposted elsewhere.

As the Version of Record of this article is going to be / has been published on a subscription basis, this Accepted Manuscript is available for reuse under a CC BY-NC-ND 3.0 licence after the 12 month embargo period.

After the embargo period, everyone is permitted to use copy and redistribute this article for non-commercial purposes only, provided that they adhere to all the terms of the licence https://creativecommons.org/licenses/by-nc-nd/3.0

Although reasonable endeavours have been taken to obtain all necessary permissions from third parties to include their copyrighted content within this article, their full citation and copyright line may not be present in this Accepted Manuscript version. Before using any content from this article, please refer to the Version of Record on IOPscience once published for full citation and copyright details, as permissions will likely be required. All third party content is fully copyright protected, unless specifically stated otherwise in the figure caption in the Version of Record.

View the article online for updates and enhancements.
Preparation and characterization of a supported system of Ni$_2$P/Ni$_{12}$P$_5$ nanoparticles and their use as active phase in chemoselective hydrogenation of acetophenone.

Dolly C Costa$^1$, Analía L. Soldati$^2$, Gina Pecchi$^3$, José Fernando Bengoa$^1$, Sergio Gustavo Marchetti$^1$, Virginia Vetere$^1$

1- Centro de Investigación y Desarrollo en Ciencias Aplicadas “Dr. Jorge J. Ronco”-CINDECA (UNLP-CONICET-CICBA). Calle 47 Nº 257, (1900) La Plata, Bs. As. Argentina

2- Grupo de Caracterización de Materiales, Centro Atómico Bariloche, CONICET. Av. Bustillo 9500, SanCarlos de Bariloche, Río Negro, Argentina.

3- Departamento de Físico Química, Facultad de Ciencias Químicas, Universidad de Concepción. Casilla 160-C, Concepción, Chile

*Corresponding author: vetere@quimica.unlp.edu.ar

Abstract

Ni$_2$P/Ni$_{12}$P$_5$ nanoparticles were obtained by thermal decomposition of nickel organometallic salt at low temperature. The use of different characterization techniques allowed us to determine that this process produced a mixture of two nickel phosphide phases: Ni$_2$P and Ni$_{12}$P$_5$. These nickel phosphides nanoparticles, supported on mesoporous silica, showed activity and high selectivity to produce the hydrogenation of the acetophenone carbonyl group to obtain 1-phenylethanol. This is a first report that demonstrates the ability of supported Ni$_2$P/Ni$_{12}$P$_5$ nanoparticles to produce the chemoselective hydrogenation of the acetophenone. We attribute these special catalytic properties to the particular geometry of the Ni-P sites on the surface of the nanoparticles. This is an interesting result because the nickel phosphides have a wide composition range (from Ni$_3$P to NiP$_3$), with different crystallographic structures, therefore, we could think that different phases could be active and selective to hydrogenate many important molecules with more than one functional group.

Keywords: monodisperse nanoparticles of nickel phosphides, Ni$_2$P, Ni$_{12}$P$_5$, hydrogenation, chemoselectivity, acetophenone.
1. Introduction.

Over the last few decades a new family of hydrotreating catalysts, the transition metal phosphides, has emerged. They have been widely used in hydrosulfurization (HDS) and hydrogenation (HDN) processes [1,2]. The Oyama’s group have worked specially in this area [3-6]. These phosphides have physical properties, which are similar to those of ordinary metallic compounds: they are hard and strong, good heat and electricity conductors, and have high thermal and chemical stability.

Among the various transition metal phosphides, nickel phosphide is the most active catalyst for HDS and HDN reactions [7-9]. Since 1990 a temperature-programmed reduction method has been developed as a simple and economical way to prepare metal phosphides [10]. However, the main disadvantage of this method is the high reduction temperature required due to the great stability of the P-O bond. These conditions lead to a wide distribution of sizes of the nickel phosphide particles [11, 12].

More recently, nickel phosphides nanoparticles (NPs) have been obtained by thermal decomposition of nickel organometallic salts in the presence of a reducing agent and alkyl or phenyl phosphines as phosphorus source [12-15]. It is well known that up to 850 °C there are many stable phases of nickel phosphides with compositions between Ni₃P and NiP₃. However, Ni₂P is obtained in most of the cases in the NPs range. Other composition that frequently appears is Ni₁₂P₃. This versatility in the arrangements allows to produce active sites with very different geometric characteristics, changing the electronic density of the Ni atoms. Thus, the selectivity of the catalytic reaction could be drastically changed.

As it was mentioned, the excellent catalytic properties of nickel phosphides for HDS and HDN, would probably indicate that they have good activity in hydrogen transfer reactions, such as hydrogenation. Additionally, the particular geometric characteristics of the active sites could produce the chemoselective hydrogenation of double bonds in those molecules with more than one functional group. Previous researches about this behavior, with nickel phosphides catalysts, are scarce. Wang et al. have studied the chemoselective hydrogenation of cinnamaldehyde with nickel phosphides catalysts supported on SiO₂ [16]. They prepared these catalysts by temperature programmed reduction of nickelphosphates. These catalysts showed a high selectivity in the hydrogenation of this unsaturated aldehyde to the saturated one. Thus, they obtained a great production of hydrocinnamaldehyde from hydrogenation of cinnamaldehyde. However, it is well known that in the hydrogenation of α,β-unsaturated aldehydes thermodynamics favours the hydrogenation of aliphatic C=C bond over C=O group in around 35 kJ/mol and due to kinetic reasons the reactivity of the C=C bond is higher than that of C=O [17]. Therefore, the article of Wang et al. shows the hydrogenating capacity of Ni₃P and Ni₁₂P₃, but without chemoselectivity.

On the other hand, there is a higher challenge related to aromatic alcohols if these have to be obtained from a selective hydrogenation of aromatic ketones. These types of alcohols are widely used in the production of drugs and in fine chemical. The aromatic ketones molecules have two groups that can be hydrogenated. Therefore, great efforts to achieve these chemoselective hydrogenations have been performed. When Pt, Rh and
Ru catalysts are used, they exhibit comparable reactivity towards the hydrogenation of carbonyl group and the aromatic ring [18-24]. In order to enhance the selectivity of C=O hydrogenation, a suitable support or modifications by the use of additives on these metals, are needed. On the other hand, Pd supported catalysts are highly reactive for the C=O hydrogenation towards to C–OH, but then they catalyze the hydrogenolysis of C–OH to give ethylbenzene as a final product [25-28]. Considering these difficulties, we guess that the expected results could be obtained using a compound with metallic characteristics with moderate hydrogenation capacity and catalytic sites with particular geometries. Thus, the spatial configuration of the adsorbed molecules and the structural characteristics of the catalytic sites would allow to “tune” the proper arrangement to reach the appropriate hydrogenation. These characteristics could be present in nickel phosphides because they have metallic properties that are able to produce moderate hydrogenations due to the diluting effect of the phosphorus atoms. Besides, the wide range of stoichiometric compositions, with very different crystallographic structures, produces surface sites with very diverse geometries. Therefore, there could be catalytic sites, with particular geometries that could hydrogenate different aromatic ketones to the desired product. Following these ideas, the aim of the present work is to explore the possibility to obtain a chemoselective hydrogenation of acetophenone to produce 1-phenylethanol using nickel phosphide species as catalyst. To our knowledge, this is the first report on this application until now.

Besides, in order to avoid possible effects of the NPs sizes on the reaction selectivity, we pre-synthesized NPs of nickel phosphides monodisperse which were then supported on mesoporous silica and used as catalyst in the acetophenone hydrogenation in liquid phase.

2. Experimental.

2.1. Nanoparticles Synthesis.

In a one-pot synthesis, determined amounts of nickel(II) acetylacetonate (Ni(acac)$_2$, 1 mmol), oleylamine as solvent and reductant (OA, 10 mmol) and triphenylphosphine as ligand and phosphorus source (PPh$_3$, 0.8 mmol) were directly added into a three-neck round bottom flask fitted with a condenser and magnetic stirring. The two remaining necks were used to introduce a thermocouple with a glass sheath and a flow of Ar. The mixture was heated at 220 °C for two hours. Finally, the NPs were purified and isolated precipitating the suspension with acetone and re-dispersed in n-hexane. The NPs were characterized by X-ray diffraction (XRD), diffuse light scattering (DLS), transmission electron microscopy (TEM), selected area electron diffraction (SAED), Fourier transformer infrared spectroscopy (FT-IR) and magnetization vs. applied field (M vs. H). Besides, the Ni content in the suspension and in the catalyst was determined by atomic absorption spectroscopy (AA).

2.2. Support synthesis.
The mesoporous silica, used as support, was synthesized following the method proposed by Ishihara et al. [29]. An acid-base-catalyzed sol-gel process with a gel skeletal reinforcement was adopted. Briefly, tetraethyl orthosilicate (TEOS) was used as silica source. This compound was dissolved in 2-propanol and HCl at 25 °C. The pH of the mixture was adjusted to 5 by adding aqueous NH₃ solution and the gelation process was performed at 50 °C during 5 h. This gel was then introduced in deionized water at 50 °C for 24 h. After this treatment, the gel was thoroughly washed with 2-propanol to remove the water present into the pores. After this step, the gel was contacted with 80% TEOS/20% 2-propanol solution at 50 °C for 48 h. Finally, the gel was washed with 2-propanol to remove the remaining 80% TEOS/20% 2-propanol solution, dried and finally calcined at 600 °C for 3 h in air flow. The final solid was characterized by N₂ adsorption at -196 °C and TEM.

2.3.Catalyst preparation.

The catalyst was prepared by wetness impregnation of the mesoporous silica with the NPs suspension with the aim to obtain a nominal Ni loading of 5 % wt/wt. In order to eliminate the surfactant (PPh₃) from the surface of the NPs two different treatments were performed. The first one was a thermal treatment at 400 °C in Ar flow (60 cm³/min) during 24 h, and the second one consisted of several washing with CHCl₃. The catalyst was characterized by thermal gravimetric analysis and differential thermal analysis (TGA-DTA), TEM and scanning electron microscopy (SEM).

2.4.Characterizations.

XRD patterns were recorded using a standard automated powder X-ray diffraction system Philips PW1710 with diffracted-beam graphite monochromator, using Cu Kₐ radiation (λ= 1.5406 Å) in the range 2θ = 35–60° with steps of 0.05° and counting time of 6 s/step.

In the DLS measurements, the time correlation function G(q, t) of the light scattering intensity was measured at the scattering angle 0 = 90° with a goniometer ALV/CGS-5022F with multiple tau digital correlator ALV-5000/EPP covering a 10⁻⁶–10³ s time range. The light source was a helium/neon laser (λ = 632.8 nm) operating at 22 mW. All of the measurements were carried out at room temperature. In order to obtain appropriate estimations of the size of the particles, number and weight average hydrodynamic radii were calculated from the time correlation function with the package CONTIN [30].

For TEM measurements two different equipments were used. Samples for analysis were prepared by drying a dispersion of the NPs on amorphous carbon coated copper grids. TEM micrographs were obtained on a JEOL model JEM-1200 EX II microscope and on a Philips CM 200 UT TEM and HRTEM equipped with an ultra-twin objective lens. The electron source used was a LaB6 filament operated at 200 keV. The nominal resolution was 0.2 nm for high resolution mode. The microscope was equipped with a
CCD camera for digital acquisition; contrast and illumination were linearly adjusted after using commercially available image treatment programs. Point elemental analysis was done using an EDS (EDAX) system coupled to the TEM. The excited spot was 150 nm diameter on the surface of the sample and acquisition time was 100 s. Besides, the electron diffraction of a selected area (SAED) was obtained.

The TGA-DTA measurements were performed on a RIGAKU Thermo plus EVO equipment. The samples were located inside an alumina crucible and heated from room temperature up to 1000ºC at a heating rate of 10ºC/min under air flow (100 cm³/min). A FT/IR Jasco spectrometer model 4200 equipped with a PIKE diffuse reflectance IR cell with a resolution of 1 cm⁻¹ was used. From 200 to 400 scans were accumulated in each case.

The magnetic measurements were carried out using a multipurpose physical magnetic system (MPMS) superconducting quantum interference device (SQUID) from Quantum Design. The magnetization versus magnetic field (M vs. H) curve was recorded at 6 K up to a maximum magnetic field of 5 T.

The textural properties, specific surface area (S_g), specific pore volume (V_p) and pore diameter (D_p), were measured with a Micromeritics ASAP 2020 V1.02 E device.

The SEM measurements were performed with a scanning electron microscopy model FEI ESEM Quanta 200. The Ni/P ratio on the catalyst surface was determined by energy dispersive analysis of X-ray with an EDAX SDD Apollo 40 probe.

2.5. Catalytic tests.

Hydrogenation of acetophenone was carried out in a stirred autoclave reactor at a pressure of H_2 of 1 MPa and a temperature of 80 ºC, using 0.25 g catalyst and n-heptane as solvent. The experimental conditions for the catalytic tests were specifically chosen to avoid mass transfer control. The course of the reaction was followed by gas chromatography in a GC Varian 3400 chromatograph equipped with a capillary column of 30 m CP wax 52 CB and FID. The identification of the diverse reaction products was accomplished by GC/MS using a Shimadzu QP5050 equipment.

3. Results and Discussion.

In Figure 1, the XRD pattern of the as synthesized NPs is shown. The detected peaks at around 20 = 40.5°, 44.2°, 46.8° and 54.0° could be assigned to Ni₁₂P₅ (tetragonal, PDF 74-1381) or Ni₂P (hexagonal, PDF 74-1385) respectively. Besides, the peaks at about 38.4° and 48.8° appear exclusively in Ni₁₂P₅. The crystalline planes assignable to Ni₁₂P₅ are indicated in red and those corresponding to Ni₂P in blue. After analyzing some aspects of the diffractogram we can observe that:

- the signal at 48.8° appears exclusively in Ni₁₂P₅ and represents its most intense peak,
- both phosphides show a peak at around 40.5° which corresponds to the most intense signal of the Ni₂P. However, for Ni₁₂P₅ it represents approximately 16% of its most intense peak.
Considering that the peaks at 40.5° and 48.8° have similar intensities, we can conclude that a mixture of both phosphides was obtained. All peaks are considerably broadened, due to the small size of the NPs (this topic will be described below). Consequently, the precise assignment of the 2θ values is not possible in some of them. For the same reason, the peaks at 2θ = 44.6° and 41.6° assignable to the planes (0 2 1) of Ni₂P and (3 3 0) of the Ni₁₂P₃, respectively, cannot be distinguished. Finally, it is worth noting that the quality of the diffractogram is not optimal because to deposit a good quantity of the NPs on the XRD holder, in order to reach a homogeneous covering, is a hard task.

In order to obtain the size distribution of the NPs we have performed DLS measurements. It is worth mentioning that DLS measurements are easy and fast to perform and also provide significant statistical information. As a disadvantage, the values obtained are the hydrodynamic diameters, that is, the sum of the NPs diameter plus the thickness of the NPs coverage. Depending of the system, a significant or a negligible difference compared to the real diameter, can be found [31]. The result obtained by this technique show that the NPs suspension is monodisperse (polydispersity index < 0.1) and they have an average diameter between 12 and 14 nm.

In order to confirm this average diameter we obtained TEM micrographs. In Figure 2, can be seen two representative micrographs. Figure 3 shows the histogram obtained counting 622 NPs. The histogram was fitted using a log-normal distribution according to previous studies showing that very small particles (i.e. lower than 20 nm) present a log-normal size distribution [32]. The statistical parameters obtained from the fitting produced an average diameter of 9.6± 0.2 nm. Considering the DLS results the NPs are monodisperse and the average diameter value is very similar to that obtained by TEM, although slightly lower in coincidence with the previous explanation.
To obtain structural information of the NPs, the electron diffraction of a selected area (SAED) was acquired and analyzed. Table 1 shows the lattice spacing measured from the rings of the diffraction pattern (Figure 4) and compares them with the known lattice spacing for bulk Ni$_{12}$P$_5$ and Ni$_2$P. This technique would indicate the presence of a mixture of both phosphides, but in a less categorical way because, in the analyzed spot, the diffraction rings corresponding to the (1 1 2) and (3 1 2) crystallographic planes of Ni$_{12}$P$_5$ were not detected.

Figure 3: histogram of the sizes of nickel phosphide nanoparticles obtained from TEM. Blue line was obtained fitting the results by assuming a log-normal distribution.
Table 1. Comparison of lattice spacing between the nickel phosphide NPs, measured by SAED technique, with the powder diffraction files data.

<table>
<thead>
<tr>
<th>Experimental d-spacing (Å) obtained from SAED</th>
<th>d-spacing (Å) from data files</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ni$_2$P</td>
</tr>
<tr>
<td>2.27</td>
<td>2.21</td>
</tr>
<tr>
<td>2.00</td>
<td>2.03</td>
</tr>
<tr>
<td>1.76</td>
<td>1.69</td>
</tr>
</tbody>
</table>

On the other hand, the lattice spacing was obtained using the inverse Fourier transform of HRTEM images of three different and isolated NPs (Figures 5A, 5B and 5C). These values were: 2.03 Å, 2.21 Å, 2.50 Å and 2.34 Å. The first three values could be assigned to (0 2 1), (1 1 1) and (2 0 0) planes of Ni$_2$P or (3 3 0), (2 0 2) and (3 0 1) planes of Ni$_{12}$P$_5$ respectively, and the spacing of 2.34 Å corresponds exclusively to (1 1 2) plane of Ni$_{12}$P$_5$. It can be seen that only one class of crystalline planes is present in each of the NPs, there are no planes changing their direction and all planes are straight. Therefore, each NPs has only one nickel phosphide phase. In summary, the suspension would consist of a mixture of pure NPs of Ni$_{12}$P$_5$ and Ni$_2$P.

Figure 4: d-spacing of the nickel phosphide nanoparticles determined by electron diffraction of a selected area (SAED).
Figures 5A, 5B and 5C: HRTEM of three different isolated nanoparticles (right) and their inverse Fourier transforms (left).

The magnetic characterization was performed on the NPs suspension. M versus H loop obtained at 6 K is shown in Figure 6. This curve can be fitted with a Brillouin function denoting a paramagnetic behavior. This result is coherent with the presence of Ni$_2$P or Ni$_{12}$P$_5$ [33-36]. It is important to remark that a superparamagnetic behavior can be discarded taking into account that the temperature at which the measurement was done.
This conclusion can be obtained considering that metallic nickel nanoparticles smaller than 15 nm are typically superparamagnetic at room temperature, but they are magnetically blocked at 10 K [37]. As the magnetic anisotropy constant of nickel phosphides is about six times higher than the one of the metallic nickel, their magnetic blocking temperature must be considerably higher than 6 K [38].

![Graph](image)

**Figure 6:** hysteresis cycle of nickel phosphide nanoparticles at 6 K using a maximum applied magnetic field of \( H = 5 \) T.

On the other hand, the Ni/P ratio obtained from the average of several EDAX measures on the calcined catalyst was of 2.35. Taking into account that the stoichiometric ratios are 2.4 for \( \text{Ni}_{12}\text{P}_5 \) and 2 for \( \text{Ni}_2\text{P} \), a combination of the presence of both phases could explain this experimental result with \( \text{Ni}_{12}\text{P}_5 \) as a dominant phase. Using the expression: 

\[
(Ni/P)_{\text{exp}} = 2 \times X + 2.4 \times Y,
\]

where \( X \) = molar fraction of \( \text{Ni}_2\text{P} \) and \( Y \) = molar fraction of \( \text{Ni}_{12}\text{P}_5 \) and considering that \( X+Y = 1 \), the following composition of the mixture can be obtained: 87% molar of \( \text{Ni}_{12}\text{P}_5 \) and 13% molar of \( \text{Ni}_2\text{P} \). It is necessary to highlight that EDAX analysis attached to SEM is not fully reliable for obtaining quantitative values. However, this technique can be used to obtain a rough estimation of the molar percentage of the nickel phosphides species. Besides, this assay reinforces the conclusion about the presence of both phases of nickel phosphides. It is interesting to mention that, in the catalyst without calcination, the Ni/P ratio determined by this technique was 0.80. This result can be explained because the presence of Ph\(_3\)P on the NPs surface increases the P content.

There is a controversy about the synthesis temperature necessary to obtain nickel phosphide from organophosphorus compounds. In this way, the research groups led by Dr. Sanchez and Dr. Brock (probably the groups that have studied this subject more in depth) consider that temperatures higher than 330 °C are required [36, 39, 40]. Other authors have used the same temperature [12, 41, 42]. However, Wang et al. suggest that
this decomposition can start at 240 °C [42]. Subsequently, it was confirmed that at 200-230 °C, an amorphous Ni$_x$P$_y$ is produced. Besides, crystalline Ni$_2$P can be obtained maintaining this temperature during 24h [43, 44]. It is important to remark that, in these studies, the phosphorus compound used is trioctylphosphine instead of triphenyphosphine. On the other hand, it is commonly accepted that the small nickel nuclei act as a catalyst to break the P-organic ligand bond. Then, this free P diffuses inside the nickel NPs and produces nickel phosphide. We think that the different phosphorus compound used and the quantity of nickel nuclei produced in the initial step (as a consequence of the nickel(II) acetylacetonate : oleylamine ratio) would explain the obtained results.

To determine if the OA and PPh$_3$ remain on the NPs surface, we obtained the FT-IR spectrum (Figure 7). It was acquired by mechanical mixing of the NPs suspension with KBr and subsequent drying. We chose this procedure, instead of use the supported system, because SiO$_2$ support has bands that overlap with the ones belonging to OA and P$_3$H$_3$. The bands indicated by red arrows can be assigned to OA. Thus, the bands at 2927 and 2854 cm$^{-1}$ correspond to stretching C-H of methyl and methylene groups respectively, at 1710 cm$^{-1}$ to C=O stretching (the carbonyl group is produced by reaction between OA and acetylacetonate groups) and at 1580 cm$^{-1}$ to aliphatic C=C [39, 45, 46]. On the other hand, the bands pointed out by green arrows indicate the presence of PPh$_3$. The weak signal at 3000 cm$^{-1}$ is assigned to stretching of C-H bound and at 1655 cm$^{-1}$ to aromatic C=C. Besides, the bands at 1454 and 1421 cm$^{-1}$ correspond to the stretching of the bound P-C$_6$H$_5$. Finally, the bands at 1065 and 960 cm$^{-1}$ are attributed to aromatic C-H in-plane vibrations and at 742 and 720 cm$^{-1}$ to out-of-the-plane vibrations [39, 46, 47]. The characteristic bands appear slightly shifted, indicating that the hydrocarbon chains in the monolayer surrounding the nanoparticles are in a closed-packed [48]. The presence of these surfactants on the NPs surface would explain the diameter differences between DLS and TEM results. Finally, the blue arrow at 1080 cm$^{-1}$ can be assigned to the stretching of metallic Ni atom bound to P=O. This species would be originated by the oxidation of the PPh$_3$ on the surface of the NPs when they are contacted with air during the FT-IR spectrum measurement. Considering that the spontaneous oxidation of PPh$_3$ by O$_2$, either in the solid state or in solution is very slow in air contact, the nickel phosphide surface would act as a catalyst in this process [39]. The coverage of the NPs surface by OA and PPh$_3$ would block the catalytic active sites. This has been confirmed based on the fact that the catalytic reaction, using the catalyst as it was prepared (without any post-synthesis treatment), was found inactive.

The silica support is made up of nanometric spheres of about 200 nm with interparticular channels (Figure 8). Their textural properties are: BET specific surface area: 508 m$^2$/g, average pore diameter, obtained by BJH method: 27 nm and pore volume: 2.5 cm$^3$/g. The Ni loading of the catalyst, determined by AA, was of 5 % wt/wt.

After wetness impregnation of the silica support with the NPs nickel posphides suspension, TEM micrographs show that the NPs did not change their size and they are located preferentially on the surface of the SiO$_2$ spheres, but some of them are placed inside the interparticular mesoporous (Figure 8).
Figure 7: FTIR of nickel phosphide nanoparticles as prepared (black spectrum), nanoparticles with thermal treatment (red spectrum) and nanoparticles washed with \( \text{Cl}_3\text{CH} \) (blue spectrum).
With the purpose of studying the elimination of the NPs superficial covering, a TGA-DTA measurement was performed. Figure 9 shows that the highest velocity of mass loss (attributed to the surfactants elimination) occurs at about 330 °C. Taking into account that the nickel phosphides are stable up to 890 °C, the catalyst was heated at 400 °C in Ar flow during 24 h in order to eliminate the surfactant before performing the catalytic test [40]. Figure 10 shows a TEM micrograph that reveals that NPs sintering did not occur during this treatment. This result is coherent with the high Tamman temperature of the nickel phosphide (about 560 °C). Above this temperature the mobility and reactivity of the atoms become appreciable and sintering could occur.

Figure 7 displays the NPs FT-IR spectrum after this thermal treatment. Again, the spectrum was obtained by mechanical mixing of the NPs suspension with KBr and this solid was then heated following the thermal treatment previously described. Figure 7 shows that all peaks assigned to OA have disappeared (bands at 2927, 2854, 1710 and 1580 cm\(^{-1}\)). Another interesting point is that the PPh\(_3\) partially disappears. Thus, in the range of 720-740 cm\(^{-1}\), the bands attributed to aromatic C-H out-of-the-plane vibrations are clearly visible. On the other hand, the fine structure of the peaks at 1655, 1454 and 1421 cm\(^{-1}\) is lost and only a broad peak can be seen. According to the TGA-DTA results, the thermal treatment would be enough to eliminate the surfactants. A greater sensitivity of the FT-IR to detect surface species, in comparison with TGA-DTA technique, would explain this result. Therefore, we can infer that there are different types of surface sites able to adsorb the PPh\(_3\) with different strength. This would be the result of the complex surface geometric structure of the nickel phosphides, as it will be discussed below, and maybe these blocked sites would produce a decrease in the catalytic activity.

The main products of acetophenone hydrogenation are shown in Scheme 1. If the carbonyl group is hydrogenated, 1-phenylethanol (PE) is obtained. On the other hand, the hydrogenation of the aromatic ring leads to the cyclohexylmethylketone (CMK).
The hydrogenation of both products can continue and 1-cyclohexylethanol (CE) could be obtained. Besides, traces of ethylbenzene (EB) and ethylcyclohexane (EC) can be produced. These products can be obtained by hydrog enolysis of C-O bond of intermediates alcohols or initial hydrogenation of C=O followed by dehydration and subsequent addition of hydrogen to C=C bond formed. Our objective is to produce a chemoselective hydrogenation of the carbonyl group to obtain 1-phenylethanol. Figure 11 shows that after 7 h of reaction time a conversion of 30% was obtained and the selectivity to 1-phenylethanol was of 90%. Therefore, the Ni$_2$P-Ni$_{12}$P$_5$/SiO$_2$ system is active in the acetophenone hydrogenation, and it has a high chemoselectivity to the desired product.

![Figure 9: TGA-DTA diagrams of nickel phosphide nanoparticles.](image)

With the aim of finding an alternative method to eliminate the surface covering of the NPs, several washes of the supported catalysts with CHCl$_3$ were performed, following the method proposed by Senevirathne et al. for a similar system (bulk Ni$_2$P unsupported) [12]. The FT-IR spectrum (Figure 7) is very similar to the one after the thermal treatment of the sample. In agreement with this result, the activity and selectivity achieved after this treatment are very similar to those obtained with the thermal treatment (Table 2). The advantage of the washing method is that it significantly reduces the surfactants elimination time. In addition, the EDAX analysis did not detect the presence of chloride ions. This is an important result because these ions can poison the active sites.
Figure 10: TEM image of nickel phosphide nanoparticles supported on mesoporous silica after thermal treatment.

Table 2. Conversion and selectivity to PE after 7 h of reaction for different treatments.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Conversion (%)</th>
<th>S_1-phenylethanol (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As prepared</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>Calcination in Ar</td>
<td>28</td>
<td>90</td>
</tr>
<tr>
<td>Washing with CHCl₃</td>
<td>30</td>
<td>92</td>
</tr>
</tbody>
</table>

Figure 11: selectivity diagram of acetophenone hydrogenation (operation conditions: (353 K, 1MPa H₂, 0.25 g catalyst).
In order to analyze the chemoselective results of these nickel phosphide NPs, we will begin describing the explanation suggested by Chen et al. for acetophenone hydrogenation on Pt/SiO$_2$ catalysts [49]. These authors proposed that the acetophenone has two preferential adsorption modes: $\eta^1$(O) and $\eta^2$(C,O) on Pt surface. In the first mode, the molecule is coordinated to the catalyst surface by the oxygen atom of the carbonyl group and the aromatic ring remains parallel to the surface. Under these conditions, both the carbonyl group and the aromatic ring could be hydrogenated. In the second mode, the carbonyl group is coordinated through its $\pi$ electrons (bridge coordination) and the aromatic ring is tilted with respect to the surface. This configuration would produce mainly 1-phenylethanol and the aromatic ring would not be hydrogenated. When nickel phosphides are used as catalyst to hydrogenate acetophenone some important differences with pure metals must be considered. Thus, in these phases, P atoms have higher electronegativity than Ni atoms. As a consequence these atoms can be represented as P$^{\delta-}$ and Ni$^{\delta+}$, respectively. The Ni$^{\delta+}$ surface atoms behave as Lewis acid sites, attracting the atoms with negative charge density of the acetophenone, and at the same time, as metallic sites for hydrogenation [50]. However, this is not the only difference in comparison with pure transition metals because, Sawhill et al. found that C=O is adsorbed on Ni catalysts with two different geometries: linear and bridge-like. Instead, when Ni$_{12}$P$_3$ and Ni$_2$P were studied, the bridge adsorption disappears and only the linear geometry remains [51]. The different adsorption modes of CO between Ni and Ni$_{12}$P$_3$ and Ni$_2$P cannot be justified by differences between the interatomic distances of the metallic centers. Thus, interatomic distance Ni-Ni, in metallic Ni, is of 2.49 Å, while in Ni$_{12}$P$_3$ and in Ni$_2$P these distances are around 2.45 Å and 2.61 Å respectively [52]. Therefore, these different adsorption modes could be attributed to a particular spatial arrangement of the nickel phosphide surfaces. Considering that, the electronic similarities between the carbonyl group and carbon monoxide, at a first approximation we could think, that on Ni$_{12}$P$_3$ and Ni$_2$P, the acetophenone preferential adsorption mode would be $\eta^1$(O). However, a $\eta^1$(O) adsorption mode would favor the aromatic ring hydrogenation but our experimental results determine the opposite. We attributed this fact to the special configuration of the catalytic sites on the surface of the nickel phosphide. If the bond length between the aromatic ring and the carbon atom of the carbonyl group (about 1.5 Å) is compared with interatomic distances Ni-P in Ni$_{12}$P$_3$ and Ni$_2$P: 2.21 - 2.59 Å and 2.20 Å, respectively [52] we can concluded that, the position of P$^{\delta-}$ would match, approximately, with the center of the aromatic ring. This situation would produce a strong electrostatic repulsion between them, moving away the phenyl group from the catalyst surface. As a consequence, a chemoselective hydrogenation of the carbonyl group is produced. On the other hand, the phosphorus atoms that surround the nickel atoms would produce a diluting effect on the nickel assembly decreasing the hydrogenation capacity. This would explain why the conversion value is about 30%.

Conclusions
In this work, nickel phosphides NPs were obtained by thermal decomposition of a nickel organometallic salt at low temperature. These NPs are monodisperse and their average diameter is of 10 nm. Different characterization techniques allowed us to identify these phosphides as a mixture of Ni$_{12}$P$_5$ and Ni$_2$P, where each NPs is monophasic.

These NPs, supported on mesoporous SiO$_2$, were used in the catalytic hydrogenation of acetophenone. The catalyst showed hydrogenation capacity and high chemoselectivity to 1-phenylethanol. To our knowledge this is the first time that these types of results are reported.

We attributed the preferential hydrogenation of the carbonyl group to a combination of the following effects:
- the P and Ni surface atoms have negative and positive charge densities respectively, because P atoms have higher electronegativity than Ni atoms. As a consequence, Ni$^{2+}$ can attract the oxygen atom of the carbonyl group,
- the special geometry of the Ni-P sites on the surface of the NPs allows an adequate matching, to produce a strong electrostatic repulsion of the phenyl group from the surface catalyst.

It should be pointed out that the nickel phosphides have a very wide range of compositions from Ni$_3$P to NiP$_3$. Among them there are great structural differences, producing very diverse catalytic sites. Therefore, we could guess that there would be many different types of molecules, with more than one functional group, on which a chemoselectivity hydrogenation could take place. As a consequence of these results, we can infer that the great versatility of these phases would justify new studies with different compositions and substrates.

Acknowledgments.

This work was supported by Universidad Nacional de La Plata (Projects X633 and X710). The authors are grateful to María Cecilia Moreno, CICPBA translator, for checking the English version.

References.


