Cobalt oxide-Carbon nanocatalysts with highly enhanced catalytic performance for the green synthesis of nitrogen heterocycles through Friedländer condensation

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Introduction

Development of new catalytic technologies by using carbon materials constitutes a currently hot topic. Nanocarbons as their different forms could be considered almost ideal supports for catalytic purposes due to their unique characteristics such as high surface areas, chemical inertness and thermal stability, among others, providing the appropriate chemical environment to produce fine chemicals.¹,² In this sense, in the last decades metal-supported carbon-based materials applied in catalysis have particularly attracted much attention.³ Activated carbons (ACs) have been used in catalysis for a long time.⁴ However, the application of nanostructured carbon forms such as carbon nanotubes (CNTs) and more recently carbon aerogels⁵ are issues of interest and continuously growing. For instance, Pd/Co nanoparticles supported on multi-walled CNTs have recently reported in the microwave-assisted efficient synthesis of substituted tetrazoles⁶, Cu loadings on Ni/carbon nanotubes catalysts in the hydrodeoxygenation reaction of guaiacol⁷, and RuCNT nanohybrid in the selective catalytic reduction of nitrochalcones to quinoline N-oxides⁸. AuCNTs have been also found to be interesting catalysts in diverse chemical transformations such as oxidation reaction of silanes⁹, alcohols¹⁰, and phenols¹¹, reductive amination¹² deoxygenation of amine N-oxides¹³, aldehydes N-formylation¹⁴ and in the synthesis of quinoxalines¹⁵. More recently, cobalt-carbon composites have been applied in the reduction of p-nitrophenol and p-dimethylaniline¹⁶ and as electrocatalysts for oxygen evolution reactions¹⁷.

As one of our research lines, we have developed different environmental-friendly and efficient methodologies for the synthesis of quinolines, through Friedländer condensation¹⁸ involving acid and basic activated carbons¹⁹,²⁰ and much more recently metal-free carbon aerogels²¹ and transition metal doped carbon aerogels.²²-²³ As far as we know, our studies represent the first examples of transition metal doped carbon aerogels implied in the aldol-based reactions, demonstrating that Co⁰ or Cu⁰ nanoparticles over the surface on carbon aerogels are the main active catalytic species producing quinolines in high yields.

With these items in mind, the goal of this paper is aimed onto the comparative study of using three types of carbon forms in the synthesis of quinolines and related compounds, via Friedländer reaction, in which Co nanoparticles are supported. The carbon-based materials of our choice are a carbon aerogel B-500, commercial AC, Norit RX 3, and multiwall carbon nanotubes (MWCNTs) showing different morphologies and textures. While Norit RX 3 AC is mainly a microporous sample, the B-500 aerogel shows a mesoporous character. It is noticeable that the polymerization resorcinol-formaldehyde (R-F) leading to nanostructured carbon gels allows the control of the porosity and morphology of these materials as well as of their tuned surface chemistry. In fact, the relationship of the surface properties and the catalytic behaviour of these nanocarbons are continuously revised hot topics in this field.⁵ In this work we seek to develop a novel sustainable series of carbon-based Co catalysts with enhanced catalytic performance. At this regard, the relationship between the morphology, textural properties of the different carbon samples and the effect of the concentration and dispersion of active phases onto the catalysts and the catalytic performance are envisioned.
Materials and Methods

Synthesis and characterization of the nanocatalysts

Three series of Co-catalysts were prepared by impregnation of different types of carbon materials with cobalt salts. Commercial AC Norit RX3 and DWCNTs have been purchased from NORIT Nederland B.V. and Sigma-Aldrich, respectively. Norit RX3 was milled and sieved to a particle size below 0.25 mm, without any chemical treatment, but DWCNTs were firstly treated with concentrated HNO₃, under refluxing conditions, according the experimental protocol reported by Trépanier et al. B500 carbon aerogel was synthesized in our laboratories from resorcinol (R) and formaldehyde (F) polymerization. The experimental protocol for the synthesis was reported elsewhere. In this case, resorcinol (R), formaldehyde (F) (molar ratio R/F = 1:2) and Na₂CO₃ (C) (R/C = 300) were dissolved in water (W) and cast into glass moulds (25 cm length x 0.5 cm internal diameter). After the curing period (1 day at room temperature, 2 days at 303 K and 5 days at 353 K), the gel rods were cut in 5 mm pellets, exchanged with acetone and supercritically dried with carbon dioxide. B500 carbon aerogel was obtained by carbonization of this organic aerogel at 773 K in N₂ flow in a tubular furnace using a temperature ramp of 1.5 K min⁻¹ and a soak time of 2h. No activating agents were applied. The final carbonized is milled and sieved also below 0.25 mm.

The cobalt-supported catalysts were prepared by treatment of the different carbon materials with the appropriate aqueous solution of cobalt nitrate (Co(NO₃)₂), cobalt acetate, ((CH₃CO₂)₂Co), to obtain the corresponding Co-loading. Briefly, carbon supports (5 g) were impregnated with differently concentrated Co(NO₃)₂ water solutions (25 mL). The used Co(NO₃)₂·6H₂O amounts were 0.247, 0.741, and 1.235 g corresponding to 1, 3 and 5% wt. After drying overnight at 383 K, the samples were pre-treated in He-flow at 573 K to obtain the Co active phase. Similar protocol was performed for the (CH₃CO₂)₂Co impregnated samples by using the appropriated cobalt salt amounts. Three catalyst series: CNTₙ, nCo, B500nCo and NoritinCo were obtained, where n is the theoretic Co loading expressed as % wt. Samples in which the cobalt salt for impregnation was (CH₃CO₂)₂Co were marked using Ac notation.

The morphology of carbons was analysed using a scanning electron microscope (Leo, Carl Zeiss, Geminxy-153). Images were obtained at excitation energy of 20 kV. The textural characteristics of the samples were obtained by mercury porosimetry and gases adsorption. The macro-mesoporosity of the samples was analysed by mercury porosimetry (Autoscan60 from Quantachrome). The adsorption isotherms of N₂ and CO₂ at 77 and 273 K were determined using an Autosorb 1 from Quantachrome after outgassing samples overnight at 383 K under high vacuum (10⁻⁶ mbar). Parameters as BET surface area and micropore volume (Wₜ) were calculated by applying respectively the BET or Dubinin-Radushkevich equations to the isotherms. The pore size distributions (PSD) were determined by applying Quenched Solid Density Functional Theory (QSDFT), assuming slit-shaped pores, to the N₂ adsorption isotherms. The total pore volume (Vₜ) is considered as the volume of adsorbed nitrogen at a relative pressure P/P₀ = 0.95.

Thermal treatments of samples were simulated by thermogravimetric analysis using a SHIMADZU mod. TGA-50H for TG-DTG experiments. The metal content of the supported catalysts was obtained by burning a fraction of it in the thermobalance at 1123 K under air flow up to constant weight. Thermal Programmed Desorption (DTP) curves of carbon monoxide, carbon dioxide, water and hydrogen were recorded. Experiments were carried out by heating 150 mg of sample at 303 K min⁻¹ up to 1273 K using He as the carrier gas (60 cm³ min⁻¹). The analysis of desorbed gases was performed with a Mass Spectrometer model Prisma (Pfeiffer).

The chemical nature and dispersion of the metal were studied by X-ray diffraction (XRD, Phillips PW1710 diffractometer), high-resolution transmission electron microscopy (HRTEM, Phillips CM-20 or HAADF FEI TITAN G2 microscope) equipped with FEI system of EDAX analysis. More than 100 metal nanoparticles were analysed in each case from the corresponding microphotographs using the appropriate software (ImageJ) where a mean diameter is calculated. X-ray photo emission spectra (XPS) were recorded using a Kratos Axis Ultra-DLD spectrometer. This instrument utilizes as the radiation source Mg Kα (1253.6 eV) and a hemispheric electron analyser operating at 12 kV and 10 mA.

Catalytic performance

In a typical experiment, a mixture of the corresponding 2-aminoaldehyde 1 (0.5 or 2 mmol) and ethyl acetoacetate 2 (5 mmol) in a three-necked vessel, equipped with thermometer, was placed on a multieperiment work station StarFish (Radley’s Discovery Technologies IUK). When the temperature reaches 303 or 323 K, the catalyst was added (25 mg) and the reaction mixture was stirred during 240 min. The samples were periodically taken at 15, 30, 60, 120, 180 and 240 min, diluted with acetone (0.5 mL), and the catalyst was filtered off and the solvent evaporated in vacuo. The reactions were followed by TLC chromatography performed on DC-Aulofolien/Kieselgel 60 F245 (Merk) using mixtures of CH₂Cl₂/EtOH 98:2 as eluent. Product was characterized by ¹H NMR spectroscopy. NMR spectra were recorded by using a Bruker AVANCE DPX-300 spectrometer (300 MHz for ¹H). ¹H chemical shifts in [d₆]DMSO are referenced to internal tetramethylsilane.
Results and Discussion

Physico-chemical characterization of the carbon catalysts

The physico-chemical properties of the final catalysts will be related with the characteristics of supports. Considering that the samples were prepared by impregnation, the metal content obtained by TG for the different catalysts fits according the programmed loading. The elemental analysis of CNT$_{ox}$, B500 and Norit supports and the oxygen content calculated from the DTP experiments are collected in Table 1. Heating in inert atmosphere the weight loss (WL) undergone by the samples corresponds to the amount of gases evolved as CO$_2$ from the decomposition of the oxygenated surface groups (OSG), which can be also characterized as a function of their thermal stability. The higher oxygen content determined by elemental analysis for the Norit support is influenced by ash content of this sample (4.8 %). In fact, the oxygen content determined by DTP is only around 2%. Lower oxygen content is determined for B500 aerogel. In this case, the only oxygen source is the resorcinol-formaldehyde used for the synthesis that is forming methylene-ether links between aromatic rings in the organic aerogels, but these links are mainly evolved during the carbonization process.$^{30}$ This sample is obviously ash free, because it is synthesized from pure organic monomers. The corresponding CO and CO$_2$ DTP profiles are showed in Fig. 1A and 1B. The presence of different maxima in the profile shows the presence of different types of OSG with different stability. The first peak at low temperature (at around 543 K) corresponds to the evolution of carboxylic acid groups. When increasing temperature, the decomposition of OSG such as anhydrides, formed from dehydration of adjacent carboxylic groups occurs. Mainly in the case of Norit sample there are present other CO$_2$-evolving groups with higher thermal stability. The peak at around 973 K is associated to the decomposition of lactone groups. In the case of the CO-profile, CO is also formed from anhydride groups (CO + CO$_2$), and with increasing temperatures from phenol, carbonyl or ether OSG, the latter decomposing at around 1173 K.$^{31,32}$ Similar conclusions that those previously described from TPD was pointed out when analysing the profiles obtained by TG-DTG in He flow (Fig. 1C).
The oxygen content for CNT\textsubscript{ox} support evidently increases after HNO\textsubscript{3} treatment regarding raw CNT (Table 1), changing also the nature of the oxygen surface groups. Thus, it is observed that effectively the WL of CNT\textsubscript{ox} > CNT due to the formation of OSG. The clear maximum at around 550 K observed for CNT\textsubscript{ox} sample corresponding to the evolution of carboxylic acid groups while the main desorption occurring at around 973 K associated to the CO evolved from ether or carbonyl groups.

The morphology of the samples was investigated by SEM/TEM (Fig. 2). The structure of aerogel B500 corresponds to the well-known structure built by aggregation of nearly spherical primary particles. The support CNT\textsubscript{ox} also maintains evidently the typical morphology of MWCNT while Norit AC is a particulate material with more or less rough surface (not shown). This morphology is also related with the porosity of the samples.

![Fig. 2 Morphology of A) B500 and B) CNT\textsubscript{ox} supports.](image)

The morphology of the samples was investigated by SEM/TEM (Fig. 2). The structure of aerogel B500 corresponds to the well-known structure built by aggregation of nearly spherical primary particles. The support CNT\textsubscript{ox} also maintains evidently the typical morphology of MWCNT while Norit AC is a particulate material with more or less rough surface (not shown). This morphology is also related with the porosity of the samples.

The textural characteristics of supports and catalyst derivatives showed, as an example, the influence of the precursor on the textural properties of the catalysts. Although as commented, the microporosity decreases in both cases regarding the activated carbon support, this blockage is stronger, for the same metal loading and pre-treatment, when using nitrate instead cobalt acetate. Nevertheless, both catalysts showed a similar hysteresis cycle at the same range of P/P\textsubscript{0} than those observed in the support, indicating that the mesoporosity is preserved. The textural characteristics of supports and catalyst derivatives are numerically summarized in Table 2. The study of the microporosity is also complemented by CO\textsubscript{2} adsorption, considering that this adsorbate can be adsorbed into the narrowest micropores, i.e. in pores with diameter < 0.7 nm. The higher surface values correspond to the Norit support as consequence of the high microporosity of this sample as above commented, while CNT\textsubscript{ox} presents the lowest surface area value in spite to present the highest total pore volume, V\textsubscript{t}. In all cases, the porosity of the catalysts changes regarding their supports, this feature being influenced by the location and distribution of the metallic particles, as detected in previous works. Thus, when supported on CNT\textsubscript{ox}, S\textsubscript{BET} surface area values do not change with the Co-loading, even the micropore volume determined by N\textsubscript{2}-adsorption increasing. The pore blockage in this case occurs inside the narrowest micropores, determined by CO\textsubscript{2}-adsorption (W\textsubscript{o} (CO\textsubscript{2}) which progressively decreased (Table 2). On the contrary, using Norit as support the stronger micropore blockage occurs in the larger microporosity, W\textsubscript{o} (N\textsubscript{2}), and is stronger when using nitrate instead acetate, suggesting a different metal nanoparticle size. In this case W\textsubscript{o} (CO\textsubscript{2}) and consequently the S\textsubscript{BET} surface area, increases regarding the corresponding supports because the narrowest microporosity is enhanced due to the formation of metal nanoparticles inside the larger microporosity.

\begin{table}[h]
\centering
\caption{Elemental analysis of the supports.}
\begin{tabular}{lcccccc}
\hline
Sample & C & H & N & S & O\textsuperscript{[a]} & O\textsubscript{DTP} \\
\hline
Norit & 89.0 & 0.3 & 0.7 & 0.0 & 11.0 & 2.8 \\
B500 & 96.3 & 0.2 & 0.0 & 0.0 & 3.5 & 1.2 \\
CNT\textsubscript{ox} & 92.0 & 0.4 & 0.2 & 0.0 & 7.4 & 5.1 \textsuperscript{[b]} \\
CNT & 94.2 & 0.5 & 0.6 & 0.1 & 4.7 & 2.5 \textsuperscript{[b]} \\
\hline
\end{tabular}
\textsuperscript{[a]} Oxygen content obtained by difference. \textsuperscript{[b]} From TG experiments.
\end{table}
Table 2. Porosity parameters of the samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>( S_{\text{BET}} ) (m(^2)g(^{-1}))</th>
<th>( S_{\text{CO}_2} ) (m(^2)g(^{-1}))</th>
<th>( W_{0(N_2)} ) (cm(^3)g(^{-1}))</th>
<th>( W_{0(CO_2)} ) (cm(^3)g(^{-1}))</th>
<th>( V_T ) (cm(^3)g(^{-1}))</th>
</tr>
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<tbody>
<tr>
<td>CNT(_{\text{ox}})</td>
<td>339</td>
<td>121</td>
<td>0.14</td>
<td>0.10</td>
<td>2.08</td>
</tr>
<tr>
<td>CNT(_{\text{ox}})1Co</td>
<td>219</td>
<td>65</td>
<td>0.08</td>
<td>0.05</td>
<td>1.49</td>
</tr>
<tr>
<td>CNT(_{\text{ox}})3Co</td>
<td>228</td>
<td>41</td>
<td>0.09</td>
<td>0.04</td>
<td>1.45</td>
</tr>
<tr>
<td>CNT(_{\text{ox}})5Co</td>
<td>222</td>
<td>32</td>
<td>0.12</td>
<td>0.03</td>
<td>1.64</td>
</tr>
<tr>
<td>Norit</td>
<td>1233</td>
<td>621</td>
<td>0.54</td>
<td>0.27</td>
<td>0.61</td>
</tr>
<tr>
<td>Norit3CoAc</td>
<td>1147</td>
<td>765</td>
<td>0.31</td>
<td>0.48</td>
<td>0.60</td>
</tr>
<tr>
<td>Norit3Co</td>
<td>1050</td>
<td>696</td>
<td>0.27</td>
<td>0.44</td>
<td>0.54</td>
</tr>
<tr>
<td>B500</td>
<td>629</td>
<td>550</td>
<td>0.22</td>
<td>0.16</td>
<td>1.25</td>
</tr>
<tr>
<td>B5003Co</td>
<td>619</td>
<td>470</td>
<td>0.22</td>
<td>0.17</td>
<td>1.20</td>
</tr>
</tbody>
</table>

\( W_{0(N_2)} \) = micropore volume and \( V_T \) = total volume both determined by \( N_2 \) adsorption. \( W_{0(CO_2)} \) = micropore volume determined by \( CO_2 \) adsorption.

Fig. 3 Porous characteristics of supports. A) PSD obtained by mercury porosimetry. B) \( N_2 \)-adsorption isotherms (close symbol: adsorption open symbol: desorption branch). C) PSD obtained from \( N_2 \)-adsorption isotherms by applying the DFT method. D) Influence of the metal precursor on the porosity of catalysts.

The behaviour of B500 as support is clearly intermediate, because this material presents both a well-developed mesoporosity but also significant microporosity, allowing therefore the deposition of the metal particles on the large external surface of the mesopores and leading to a smaller reduction of the porosity.

The location and distribution of metallic nanoparticles (dispersion) on the different supports were analysed by HRTEM (Fig. 4). In the case of CNT\(_{\text{ox}}\), the internal hollow is around 7 nm in diameter. Inside this porosity sometimes small Co-nanoparticles are placed (marked in the Fig. 4 with dense arrows) the most of them being located on the external surface of the CNTs (marked with hollow arrow). The particle size distribution obtained for these samples are showed in Figure 5. Although in general external particles are bigger than those located inside the CNTs, many of them are also quite small but their particle sizes tend to increase with increasing the Co-
loading thus the distribution shifted from particles with a typical diameter around 7 nm to particles of around 13 nm, and also the formation of larger particles, exceeding the 20 nm were observed in this sense, which also favor the CNTs grouping, this fact being related with the detected increase of W0 (N2).

**Fig. 4** A) CNTox3Co, B) CNTox5Co, C) B500-1Co, D) B500-5Co, E) Norit3Co and F) Norit3CoAc samples. Dense arrows point out small CoO-nanoparticles inside the pores whereas hollow arrows indicate those located at external surface.

**Figure 5.** Distribution of the nanoparticle size determined by HTREM for CNTox nCo

As previously commented, the support B500 is composed by nearly spherical carbon particles that coalesce between them, forming a solid with high mesopore volume between these primary particles and giving a granulated aspect to the support. The high external surface associated to these mesopores provides an adequate support to develop highly dispersed catalysts, in such a manner that Co-particles are quite difficult to be distinguished at low Co-loading (Fig. 4C) and causing a poor variation of the porous texture. Nevertheless, sintering increases at high Co-loading (Fig. 4D). The Norit AC is a denser but shows large micropores allowing a very high dispersion of Co-nanoparticles. It is important to stress that the Co-particle size increases when use acetate instead nitrate salt (Fig. 4E and 4F). However, even in the worst case, well distributed nanoparticles with diameter of around 13 nm were obtained. Similar information is also obtained by mapping of different areas of the catalysts (Fig. 6), the images showing the good and homogeneous dispersion of Co on the CNTs surface.

**Fig. 6** Mapping of Co-species on the CNTs (CNTox3Co).

Although the Co-nanoparticles reach a significant size in some samples, their XRD patterns do not present any diffraction peak associated to the Co-phase. At this regard, we try to identify the chemical nature of the metallic particles and the interactions with the supports by XPS analysis of the samples. Thus, the oxygen, carbon and cobalt spectral regions were analysed (Table 3). The peak C1s at the carbon region was fitted using four components. The first one at 284.9 eV corresponding to aromatic and aliphatic carbon, while the other ones are associated to the presence of heteroatoms; the component at 286.5 eV corresponding to single C-O bonds, the component at 288 eV to C=O and finally the carbonate species at around 290 eV. The oxygen region O1s is fitted using two components, the first one at around 532 eV associated to C=O bonds and the second one at 533 eV corresponding to C-O bonds. In the catalyst, although this region is fitted also using only two components, clearly the component at low B.E. is favoured after impregnation, due to the contribution of oxygen linked to the metal nanoparticles. Consequently, the oxygen content also significantly increases. The atomic Co-content on surface is 5.8%, corresponding to a 18.4% wt. This value is clearly higher than the nominal loading of 3% indicating that the metal is well dispersed only on the high external surface of the support. The analysis of the Co-spectral region is notably more difficult due to the presence of shake-up satellites together with the Co(2p3/2) and Co(2p1/2) photolines (Fig. 7). The components at 781.9 Co(2p3/2) and 797.7 eV Co(2p1/2) with the satellite at 787.1 eV suggest the presence of Co3+, probably as oxide/hydroxide species.
The existence of oxidized Co-species on the catalysts surface is also pointed out by TG-DTG experiments carried out in He flow (Fig. 8). Comparing this result with those previously presented for the supports in Fig. 1C, it is noteworthy the new significant peak at around 640 K, that progressively increases with increasing the metal loading. This peak is associated to the WL produced by the reduction of the Co oxide/hydroxide by the carbon matrix, evolving COx.

Interestingly, it is important to remark that these catalysts significantly differ from those previously developed and used in the reaction under study. Previous Co-catalysts were prepared by metal-doping during the sol-gel synthesis of the carbon aerogel, and thereafter carbonized at high temperature (1273 K), therefore metal particles were reduced to Co0 during the thermal treatment. In this case, carbon catalysts are prepared by impregnation, metal particles formed exclusively on the support surface during a soft thermal treatment at 573 K. In such a way, amorphous nanoparticles remaining as cobalt oxides/hydroxides are obtained, which should present therefore also a different catalytic performance as showed below.

**Figure 7** Fitting the XPS spectral region of Co in the B5003Co catalysts.

Fig. 9A depicts the comparison of yield vs time for the carbon supports Norit, B500 and CNTox when operating at 323 K exhibiting notable differences at the shortest reaction times. In these cases, the porosity of the samples seems directly affect to the reactivity, the presence of the mesopores in B500 leading to increased yields of compound 3a. Similar trend was observed for the CNTox regarding the B500 sample. In this case, the smaller surface can be compensated by large interparticle volume and the presence of OSG generated during the acid treatment, which could be also responsible of the selective formation of quinoline 3 together π,σ-stacking interactions between catalysts and reactant as accounted for the previous reported studies concerning the oxygenated carbon materials.

The reaction under the same experimental conditions used above and in the presence of the Co-supported nanocarbons afforded the quinoline 3a in almost quantitative yields after 15 min of reaction time. Then, CoO nanoparticles supported over the carbon surface notably increased the yields of 3a. Remarkably CoO is now the catalytic specie, much more active than the Co2 nanoparticles.

Having these results in mind, we carried out the reaction notably increasing the reagent molar ratio (2:5 vs 0.5:5) but also decreasing the reaction temperature, operating at temperature close to room temperature, 303 K. The catalytic behaviour of the samples based on mesoporous nanocarbons by varying the concentration of the active phase is summarized in Fig. 10. For the carbon aerogel-based catalysts, it was observed an increment of yield of 3a as function of the concentration of Co active phase in such a manner that the B5005Co sample led to compound 3a with increased yields.

**Catalytic performance**

First, the catalytic performance of the investigated nanocarbons used as supports was checked on the condensation of 2-amino-5-chlorobenzaldehyde (1a) and ethyl acetoacetate (2) for the synthesis quinolines 3a (Scheme 1) as an analogue of biologically relevant compound. As demonstrated in our previous studies, the carbon support is active in the reaction when operating at 323 K, producing the quinoline 3a in low yields probably involving π,π-stacking interactions.

![Scheme 1 Friedländer condensation from 2-aminoaryl aldehydes (1) and ethyl acetoacetate (2), under solvent-free conditions.](image-url)
Table 3. Atomic Surface composition (by XPS) and Binding Energy of B500 support and B5003Co catalyst.

<table>
<thead>
<tr>
<th>Sample</th>
<th>C (at%)</th>
<th>O (at%)</th>
<th>Co(at%)</th>
<th>B.E. C (eV)</th>
<th>B.E. O (eV)</th>
<th>B.E. Co (eV)</th>
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<tr>
<td>B500</td>
<td>92.8</td>
<td>7.2</td>
<td>-</td>
<td>284.9 (74%)</td>
<td>531.9 (24%)</td>
<td>--</td>
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<td></td>
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<td>286.5 (18%)</td>
<td>533.8 (76%)</td>
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<td>288.8 (6%)</td>
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<td></td>
<td>290.8 (2%)</td>
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<tr>
<td>B5003Co</td>
<td>72.6</td>
<td>21.6</td>
<td>5.8</td>
<td>284.8 (73%)</td>
<td>531.5 (41%)</td>
<td>781.9 (100%)</td>
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<td></td>
<td>286.3 (16%)</td>
<td>533.6 (59%)</td>
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<td>289.5 (5%)</td>
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However, at short reaction times this catalyst is slower than B5003Co, probably associated to the worse dispersion of the active phase previously showed. This behaviour is even more significant in the CNT$_{ox}$nCo serie in which it seems that the CNT$_{ox}$3Co nanocarbon shows the optimal amount of CoO exhibiting the highest yields to product.

Regarding to that, the CNT$_{ox}$1Co and NTC$_{ox}$5Co samples showed a similar catalytic behaviour affording the quinoline 3a with almost identical yields. In these cases, the highest metal content distributed over the lowest surface area of CNT$_{ox}$ also produce the formation of CoO aggregates as demonstrated by the HRTEM images. The effect of the porosity and the nanoparticles location is even more noticeable at intermediate metal loadings (Fig. 10). CoO supported on B500 are faster than when supported on CNTs, in this case favoured by a high dispersion degree (Fig. 4C). Thus, although B5003Co and CNT$_{ox}$3Co afforded the quinoline 3a in high yields after 1h of reaction time (up to 60%), at short reaction times (15 min) B5003Co sample yielded the quinoline 3a in 72% vs 37 and 16% for the CNT$_{ox}$3Co and Norit3Co samples, respectively (Fig. 10), this feature demonstrating the advantage of using carbon aerogels.
present in the Norit support. Nevertheless, this sample became finally the most active due to the higher catalytic surface and high Co-dispersion.

Figure 11 Friedländer condensation from 2-amino-5-chlorobenzaldehyde (1a) and ethyl acetoacetate (2) catalysed by CoO-supported nanocarbons − B5003Co, CNT_o3Co and Norit3Co − at 303 K, under solvent-free conditions.

Figure 12 Friedländer condensation from 2-amino-5-chlorobenzaldehyde (1a) and ethyl acetoacetate (2) catalysed by CoO-supported nanocarbons − CNT_o3CoAc and Norit3CoAc − at 303 K, under solvent-free conditions.

Concerning the reaction mechanism for the investigated condensation, when using CoO doped carbon aerogels we previously proposed the in situ generation of Co\(^{0}\)-Co\(^{2+}\) catalytic system, which would initiate the condensation reaction between reactants.\(^{15}\) The enhanced catalytic performance observed for the nanocarbons herein reported is then attributed to the presence of CoO. Based on our experience, CoO could act as dual catalyst in which O atom would initiate the condensation reaction between reactants, whereas Co\(^{2+}\) would activate the ethyl acetoacetate, through its enol form, thereby strongly favouring the reaction. Subsequent heterocyclization followed by double dehydration steps would yield the corresponding quinoline.

Finally, Table 4 summarizes the catalysts described for the synthesis of quinolines 3, demonstrating the high potential of CoO supported carbon catalyst under study. These catalytic materials were found to be easily prepared as a sustainable alternative to metal organic frameworks (MOFs), particularly Cu-BTC, bifunctional mesoporous silicas (AP/AI-MCF) and even other basic carbocatalysts with high metal oxide loadings (PET/MAG), showing both superior thermal stability and enhanced catalytic performance.

### Table 4: Catalysts reported for the Friedländer reaction from 2-aminoaryl aldehydes (1) and ethyl acetoacetate (2).

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Reaction product</th>
<th>Reaction Conditions</th>
<th>Time (min)</th>
<th>Yield (%)</th>
<th>Ref.</th>
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</thead>
<tbody>
<tr>
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<td>1a (0.5 mmol), 2 (5 mmol), catalyst (25 mg), 293 K</td>
<td>60</td>
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<td>Cu-BTC</td>
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<td>58</td>
<td>22</td>
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<tr>
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<td>3a</td>
<td>1a (2 mmol), 2 (5 mmol), catalyst (25 mg), 303 K</td>
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<td>This work</td>
</tr>
<tr>
<td>CNT_o3CoAc</td>
<td>3a</td>
<td>1a (2 mmol), 2 (5 mmol), catalyst (25 mg), 303 K</td>
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<td>73</td>
<td>This work</td>
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<td>AP/AI-MCF</td>
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</table>
Conclusions

We report herein the first examples of CoO supported on carbon nanomaterials highly efficient in the synthesis of nitrogen heterocyclic compounds through the Friedländer condensation, under solvent-free and mild conditions. Three series of eco-sustainable catalysts, exhibiting enhanced catalytic performance are prepared and characterized and these properties related with their catalytic performance. The role of the support (AC, CNT or carbon aerogels) porosity and surface chemistry influences the formation of CoO nanoparticles on different porosity ranges. The size and distribution of these nanoparticles are also influenced by the Co-precursor salt. Sintering increased with increasing the metal loading. In the case of mesoporous aerogels the nanoparticles can be located in the mesoporosity and large micropore range, favouring a homogeneous distribution and the accessibility of reactants (fast reactions). On the contrary, microporous ACs with high specific surface area induce slowly reaction, but yields increased with the reaction time obtaining yields even greater than in the previous case. Using CNTox in spite that present the highest pore volume, some of the metal nanoparticles are formed inside the nanotubes, showing an intermediate behaviour between mesoporous carbon aerogels and microporous ACs. Among the explored samples, BS005Co was found to be the most efficient catalyst providing highest conversions to product, at the shortest reaction times, demonstrating the advantage of using mesoporous carbon aerogels as pure carbon support. The CoO-Carbon catalysts herein reported are found to be much more efficient than MOFs, bifunctional mesoporous silicas and even basic carbon catalysts, showing also a notable stability along different reaction cycles. Then, these interesting results open the doors to the development of functional carbon aerogels able to efficiently catalyse the synthesis of relevant heterocyclic systems through cascade reactions.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Notes and references