Platelet-like catalyst design for high yield production of multi-walled carbon nanotubes by catalytic chemical vapour deposition

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Abstract

We investigated the effect of catalyst design on the synthesis of multi-walled carbon nanotubes (MWCNTs) by chemical vapour deposition (CVD). A set of highly active supported sol-gel Co-Mo/MgO and Ni-Mo/MgO catalysts was prepared systematically modifying the calcination temperature. First, the evolution of catalysts’ crystallographic phases and their morphology were studied by X-ray diffraction (XRD), Raman spectroscopy, scanning electron (SEM) and transmission electron (TEM) microscopy. Second, the catalysts were used for the CVD growth of MWCNTs. The resulting materials were analysed by SEM and TEM, Raman and XRD to establish a relation between catalyst design and MWCNT yield. We show that our catalyst synthesis route leads to the formation of laminar non-porous catalyst systems, which at a calcination temperature of 800 °C stabilise in a crystallographic phase of Me2Mg1-xMoO4 (Me = Co or Ni). We give evidence that increased MWCNT yields of more than 3000 wt% with

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respect to the catalysts are directly related to the aforementioned crystallographic phase. Finally, we propose a growth model based on the continuous exfoliation of platelet-like catalyst systems. This consistently explains the high catalytic activity towards MWCNT production using a non-porous catalyst. Our findings provide important insights for catalyst design strategies towards large-scale MWCNT production.

1. Introduction

The exceptional mechanical, thermal, electronic and chemical properties of carbon nanotubes (CNTs) [1] are the origin of enormous research activities in various fields of science and technology such as electronics, biology, materials science and energy conversion storage/conversion [2]. Progress for large-scale applications strongly depends on the availability of CNTs at competitive prices. Many efforts have been made in the last decade to develop low cost large-scale synthesis processes. Most promising are chemical vapour deposition (CVD) based methods, including fluidized bed techniques [3,4]. On one hand, they offer opportunities for a continuous and automatised production line. On the other hand, a broad set of process variables (temperature, carbon-precursor, catalyst, promoters, fluid-dynamics, etc.) ensures a high versatility to tune both CNTs characteristics and yield. However, understanding the role of the various process parameters is still a major issue when it comes to gain full control on CNT growth and to develop an effective large-scale production, which is of uppermost interest for further progress towards CNT-based products.

Among all the process parameters, catalysts are of special relevance for the formation of CNTs. Catalyst type (typically 3d metals), size (optimally in the range of a few nm) and activity largely define CNTs structural characteristics and the production yields [5-7]. The use of porous supporting matrices such as silica, alumina, zeolites, etc. is a highly
valuable way to achieve a homogeneous distribution of nanometer sized catalytic metal particles and ensure a high catalytic activity during the CVD process [8-10]. Synthesis strategies for these types of supported catalyst systems include impregnation, co-precipitation, combustion, and sol-gel methods [11-13]. The latter is based on low cost up-scalable chemical synthesis routes which can be adapted, e.g. by the addition of promoters, to design highly effective catalysts applicable as powders or even as coated films for tailored CNT growth [14-16]. However, very little information is available in what concerns the direct influence of catalyst preparation conditions on the growth of CNTs in high yields.

In this work we designed highly active Co-Mo/MgO and Ni-Mo/MgO sol-gel catalyst systems for high yield CVD production of MWCNTs. The effect of calcination temperatures on the structure and morphology of the catalyst was studied in detail and its direct impact on the influence of MWCNT yield was evaluated. We show that our synthesis strategy leads to layered non-porous catalyst systems in which a crystallographic phase of Me₅Mg₁₋₅MoO₄ (Me = Co, Ni), stabilized at a calcination temperature of 800 ºC, is responsible for achieving MWCNT yields of more than 3000 wt% with respect to the initial catalyst. Finally we present a growth model, which is based on the continuous exfoliation of the layered catalyst systems. This coherently explains the high activity of the developed non-porous catalysts and the high-yield production of MWCNTs organised in bundles grown from catalyst fragments whose sizes are related to the bundle diameters.

2. Experimental section

2.1 Catalyst synthesis and CVD reaction conditions
Catalysts were synthesized by the sol-gel method using MgO as catalyst support according to a procedure reported elsewhere [17]. In this way, two types of catalyst systems were prepared Co-Mo, and Ni-Mo with identical chemical compositions except for the active metal used which was Co, or Ni respectively.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Mo</th>
<th>Mg</th>
<th>Me(^{a)})</th>
<th>Mo/Me(^{a)})</th>
<th>Mo/Mg</th>
<th>Me(^{a)})/Mg</th>
<th>(Me(^{a)})+Mo)/Mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co-Mo</td>
<td>71.1</td>
<td>26.6</td>
<td>2.3</td>
<td>30.32</td>
<td>2.67</td>
<td>0.09</td>
<td>2.760</td>
</tr>
<tr>
<td>Ni-Mo(^{b)})</td>
<td>70.6</td>
<td>26.4</td>
<td>2.3</td>
<td>30.30</td>
<td>2.67</td>
<td>0.09</td>
<td>2.782</td>
</tr>
</tbody>
</table>

\(^{a)}\) Me= Co, Ni, for Co-Mo, and Ni-Mo catalysts respectively.

\(^{b)}\) Y was added in Y:Ni=1:4 molar fraction

Basically, metal nitrates precursors Me(NO\(_3\))\(_2\) (Me=Co, Ni), and Mg(NO\(_3\))\(_2\) are mixed in a citric acid solution, and stirred until complete dilution is achieved. The solution is dried at 120 °C into a foamy paste, which is mixed with molybdenum before performing the subsequent calcinations step. Catalyst metal loadings are shown in table 1. In the case of Ni-catalyst systems, yttrium (in the form of nitrate) was added in a Ni:Y molar ratio of 4:1 to enhance MWCNT yield, according to our former observations [18].

To study the influence of the catalyst preparation temperature on the synthesis of carbon nanotubes, the calcination process was performed at three selected temperatures, i.e. at 700°C, 750°C, and 800°C. In this way, 6 different catalyst samples were obtained and characterized. Subsequently, they were used under identical conditions in a CVD process designed for MWCNT production as described previously [17]. In brief, a flow composed by methane (1000 cm\(^3\)/min) and hydrogen (100:3 v/v) passes through a
quartz reactor (30 mm of inner diameter) placed inside a horizontal furnace. At a reaction temperature of 1000 °C, methane decomposes in the presence of the catalyst (50 mg) deposited as fine powder on a ceramic boat. After a reaction time of 30 minutes and the subsequent cooling period, MWCNT were collected from the ceramic boat, weighted and characterised.

Both catalysts and produced MWCNT materials were characterised in detail by different techniques. Structural and morphological characteristics were studied by transmission electron microscopy (TEM) on a JEOL JEM-2000FX II as well as by scanning electron microscopy (SEM) on a HITACHI S3400N. SEM-EDX mapping was applied to evaluate the homogeneity of metal deposition on the supported catalysts. Physical-chemical changes of the catalyst systems taking place during the calcinations step were studied using a combined differential scanning calorimetry-thermogravimetric analysis (DSC-TGA) in a SDT TA Instruments Q600. Samples were heated with a constant ramp of 20°C/min until 900 °C under a flow of 40 ml/min of argon and 50 ml/min of oxygen. X-ray powder diffraction measurements (XRD) on a Bruker D8 Advance Diffractometer were performed to identify the main crystallographic phases of the catalysts as well as to probe the crystalline structure of the produced MWCNT samples. Diffractograms were obtained using a Cu Kα radiation (λ=0.1542 nm) in a 2θ range from 3° to 80°, with 0.05° step size and 3 s of acquisition time. The qualitative identification of crystalline phases of the catalysts was carried out using the EVAV 8.0 program of the DIFFRAC plus package of Bruker AXS. Raman spectroscopy of the catalysts was employed to elucidate the chemical coordination of the different species in the catalyst systems thus completing structural information obtained by XRD. Raman spectroscopy also was used to elucidate the characteristics of the produced MWCNT materials. Corresponding measurements were carried out using a Horiba Jobin Yvon
HR800 UV spectrometer at an excitation wavelength of 532 nm. Spectra for catalysts and MWCNT materials were accumulated in 5 runs using an acquisition time of 45 s per run. The BET specific surface area of the catalysts was calculated from the nitrogen adsorption data acquired from nitrogen adsorption isotherms at liquid nitrogen temperature on a Micromeritics ASAP 2010 adsorption apparatus.

3. Results and Discussion

3.1. Catalysts’ physical-chemical evolution upon calcination

We first carried out a systematic DSC-TGA analysis in order to monitor the catalysts’ physical-chemical evolution at simulated calcination conditions. At the early stages of calcination, metal nitrate catalytic precursors Me(NO$_3$)$_2$ (Me=Co, Ni, Mg) decompose to the corresponding oxides which further react with molybdenum to form molybdate species. The evolution of the different precursors with temperature up to 700 ºC was followed by DSC-TGA experiments (not shown here). The transitions taking place are well described in the literature [19-21] and can be summarized as follows: below 200ºC citric acid decomposes to give CO$_x$ and water, leaving a carbonaceous residue that combust at about 400ºC. On the other hand, metal nitrates Me(NO$_3$)$_2$ decompose into nitrogen oxides (NO$_x$) and the corresponding oxides Me$_x$O$_y$ (Me= Co, Ni, Mg) at about 400ºC. In the range of calcination temperature between 500 and 750ºC, metal oxides react with molybdenum to form metal molybdates. Excess of molybdenum in these experiments oxidises beyond 500 ºC and forms non-stoichiometric MoO$_{3-x}$ oxides, coexisting with the molybdates. Molybdates’ phase transitions relevant for MWCNT growth should occur beyond 700 ºC. Therefore, the physical-chemical changes taking place in this range of temperatures comprise the focus of the following discussion.
DSC-TGA thermograms in Figure 1 show sharp endothermic peaks for both types of catalysts: at 758 °C and 849 °C for Co-Mo, and at 750 °C, 813°C and 840 °C for Ni-Mo. Furthermore, a mass loss of 50 % is observed in the temperature range between 750 °C and 900 °C, which together with the one observed at temperatures below 750 °C results in a total mass loss of around 60 % for both catalysts.

![DSC-TGA thermograms of calcination process from 650°C to 950°C for Co-Mo and Ni-Mo catalysts. DSC (--), TGA (---).](image)

The catalyst weight drop starting at 750 °C is attributed to the sublimation of MoO$_{3-x}$, formed by oxidation of the non-reacting molybdenum, as mentioned above. Consequently, similar mass losses are observed for both catalysts. Endothermic peaks could be assigned to the melt of a possible eutectic formed by the solid solution of the molybdate phases. Thus, molybdate phase transformations should take place in the vicinity of the endothermic peaks. We therefore selected three calcination temperatures around the first endothermic peak, i.e. 700 °C, 750 °C, and 800 °C, to prepare the
catalysts, and studied their crystalline composition by XRD and Raman spectroscopy. Further endothermic peaks observed around 850°C suggest additional phase transitions that will be subject of a forthcoming work.

3.2. Catalysts’ crystalline phase transitions

Figure 2 shows XRD diffractograms of Co-Mo and Ni-Mo catalysts at calcination temperatures of 700°C, 750°C, and 800°C. In general, both types of catalysts show similar XRD features for all the applied calcination conditions. At 700 °C, the MoO₃ phase is clearly identified for both Co-Mo and Ni-Mo catalysts. The corresponding diffraction peaks exhibit high intensity in 0k0 planes typical of a preferential crystal growth orientation in a laminar structure related to shear effects in MoO₃ at calcination temperatures higher than 600°C [22].

![Figure 2. XRD patterns of the Co-Mo and Ni-Mo catalysts obtained at calcination temperatures of 700°C, 750°C, and 800°C. Symbols: MoO₃ (*), Mo₄O₁₁ (^), MgMo₂O₇ (o), MgMoO₄ (<>), CoMoO₄ ([]), NiMoO₄ (x).](image)
Molybdate phases $\text{Mo}_4\text{O}_{11}$ and $\text{MgMo}_2\text{O}_7$ are also identified although with minor intensities. The molybdenum oxide phases ($\text{MoO}_3$, $\text{Mo}_4\text{O}_{11}$) are consequence of the molybdenum oxidation to form non-stoichiometric oxides at temperatures above 500 °C as mentioned above. The formation of the $\text{MgMo}_2\text{O}_7$ molybdate phase can be explained as reaction product of $\text{MgO}$ with $\text{MoO}_3$ occurring in the same temperature range.

XRD patterns for the catalysts prepared at 750°C indicate that $\text{MoO}_{3-x}$ compounds are not longer identified while $\text{MgMo}_2\text{O}_7$ is the only recognized phase. The loss of $\text{MoO}_{3-x}$ is consistent with the molybdenum oxide sublimation observed in the DSC-TGA experiments in this temperature range. Thus, catalysts obtained at a calcination temperature of 750 °C only comprise a $\text{MgMo}_2\text{O}_7$ phase whose formation already begins at temperatures below 700°C.

At calcination temperature of 800 °C, the XRD diffractograms clearly revealed the presence of $\text{MgMoO}_4$ phase. However, some differences are noticed between both Ni-Mo, and Co-Mo catalysts. XRD pattern for Ni-Mo catalysts clearly show the preservation of the $\text{MgMo}_2\text{O}_7$ phase. However this phase is not detected in the Co-Mo catalyst diffractogram. The appearance of the $\text{MgMoO}_4$ phase indicates that certain crystal reorganization processes take place after the endothermic melting of the eutectic observed at around 750°C by DSC-TGA (see Fig 2). During this reorganization, the $\text{MgMo}_2\text{O}_7$ phase completely transforms into the $\text{MgMoO}_4$ phase for the Co-Mo catalyst.

Peaks corresponding to $\text{NiMoO}_4$ and $\text{CoMoO}_4$ phases are also identified for Ni-Mo and Co-Mo catalysts respectively. Formation of the metal molybdate phases implies atomic substitution of magnesium by the catalytic metal (cobalt or nickel) within $\text{MgMoO}_4$ crystalline domains. In the case of Co-Mo catalyst, the $\text{CoMoO}_4$ phase exhibits a XRD pattern with very low intensity. This could be due to both, the low amount of cobalt in
the overall catalyst composition (see Table 1) and/or the very small cobalt particle size obtained by this method. On the contrary, the NiMoO₄ phase in the Ni-Mo catalyst shows much higher XRD intensities.

It finally remains to mention that cobalt and nickel species are not detected at lower calcination temperature possibly because the eutectic point has not been reached and the reorganization of the MeMoO₄ phase (Me=Co, Ni) has not taken place yet.

Further valuable information about the catalyst evolution during the calcinations process is obtained by probing the chemical coordination of the different molybdenum species by Raman spectroscopy. It is well known that MoO₃ has a Mo distorted octahedral coordination, α-MgMoO₄ has a distorted tetrahedral coordination and MgMo₂O₇ is considered as a polymeric solid with a mixture of both octahedral and tetrahedral coordination geometries [23]. In addition high temperature stable phase β-MeMoO₄ (Me= Ni, Co) is isotopic to α-MgMoO₄, whereas α-MeMoO₄ has a distorted octahedral coordination [20,22,24]. The different coordination geometries and chemical bond lengths of the catalyst phases alter the molecule polarizability, and thus the different phases clearly can be identified by distinctive Raman vibrational modes.

Figure 3 shows Raman spectra of the Co-Mo and Ni-Mo catalysts obtained at calcination temperatures of 700°C, 750°C, and 800°C. Raman spectra of Co-Mo and Ni-Mo catalysts obtained at 700 ºC exhibit sharp peaks at 993, 816, 663, 334, 280 cm⁻¹, which correspond to the characteristic Raman vibrations of MoO₃ in octahedral coordination geometry [25]. For catalysts prepared at 750°C, quite identical Raman spectra were obtained for both Co-Mo and Ni-Mo catalysts underlining the existence of one single phase, namely MgMo₂O₇ as identified by XRD. Additionally, the previously observed characteristic MoO₃ Raman peaks disappear in agreement with the XRD
results described before. At a calcination temperature of 800 °C, Raman spectra for Co-Mo and Ni-Mo catalyst exhibit different features indicating a change in coordination, representative for a phase transition.

![Raman spectra of Co-Mo and Ni-Mo catalysts](image)

Figure 3. Raman spectra of Co-Mo and Ni-Mo catalysts obtained at calcination temperatures of 700, 750, and 800°C. Symbols: MoO₃ (*), MgMo₂O₇ (○), MgMoO₄ (<>), CoMoO₄ ([]), NiMoO₄ (x).

In general, Raman spectra show increasing intensity with temperature for the peaks at 963 and 909 cm⁻¹, typical for Mo in tetrahedral coordination geometry. Therefore, Mo tetrahedral coordination associated to α-MgMoO₄ is becoming predominant upon increase of the calcination temperature. In particular, for the Ni-Mo catalyst, a superior enhancement of these peaks is clearly observed, which can be explained by the additional contribution of MgMo₂O₇ and β-NiMoO₄ phases, both having tetrahedral
coordination geometry. However, the Raman spectrum of the Co-Mo catalyst shows that tetrahedral related peaks (963 and 909 cm\(^{-1}\)) typical for the \(\alpha\)-MgMoO\(_4\) phase coexist with octahedral related features (993, 816, 663, and 280 cm\(^{-1}\)) assigned to an \(\alpha\)-CoMoO\(_4\) phase. It is important to remark that at this calcination temperature the Raman features are broadened and characterized by additional shoulder contributions indicative of multi-phase interactions in distorted crystal geometry of both catalysts [19, 20, 24-31].

In order to probe if crystallographic phase transitions eventually could take place at segregated catalysts’ domains, EDX mappings were performed for both types of catalyst systems at all calcination temperatures.

Figure 4 depicts typical EDX elemental composition maps of both, Co-Mo and Ni-Y-Mo catalysts. It clearly can be seen that all metals are homogeneously dispersed in the continuous Mo phase thus confirming the formation of solid-solid solutions. No changes are observed for different calcination temperatures indicating furthermore that
phase transitions take place on the overall solid-solid solution without affecting the dispersion of the active metals.

3.3. *Catalysts’ platelet-like structure*

Catalyst crystal preferential growth in a laminar structure was inferred by XRD analysis, and subsequently confirmed by electron microscopy. Figure 5 shows SEM and TEM images of both Co-Mo and Ni-Mo catalysts. Here, smooth surfaces arranged in a platelet-like structure are observed by SEM.

![SEM and TEM images of catalysts Co-Mo and Ni-Mo](image)

Figure 5. SEM (a,b) and TEM (c,d) micrographs of crystal structure obtained from catalysts Co-Mo (a,c) and Ni-Mo (b,d) at a calcination temperature of 800 °C. Insets show Fast Fourier Transformation images of the selected region of catalyst platelets.

A highly crystalline laminar structure is also clearly recognizable from TEM images and Fast Fourier Transformation analyses. In-plane preferential crystal growth in the form
of thin parallel sheets can be seen, especially at the edges of the catalytic particles. Additionally, nitrogen adsorption isotherms reveal surface areas below 2 m$^2$/g indicative of non-porous systems. Therefore, the observations consistently evidence the formation of non-porous catalysts systems having a platelet-like morphology.

3.4. MWCNT production and characterization

The aforementioned results clearly demonstrate that the calcination process strongly influences the formation of the solid-solid molbydate solutions. More specifically, the calcination temperature is the key to develop a specific phase. In the following the influence of the catalyst crystallographic phase on the production of MWCNT by CVD is evaluated.

<table>
<thead>
<tr>
<th>Catalyst type</th>
<th>Calcination temperature (ºC)</th>
<th>Crystallographic phase</th>
<th>Yield$_{MWCNT}$ (wt.%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co-Mo</td>
<td>700</td>
<td>MoO$_3$, MgMo$_2$O$_7$, Mo$<em>4$O$</em>{11}$</td>
<td>1834</td>
</tr>
<tr>
<td>Ni-Mo</td>
<td>700</td>
<td>MoO$_3$, MgMo$_2$O$_7$, Mo$<em>4$O$</em>{11}$</td>
<td>1190</td>
</tr>
<tr>
<td>Co-Mo</td>
<td>750</td>
<td>MgMo$_2$O$_7$</td>
<td>2686</td>
</tr>
<tr>
<td>Ni-Mo</td>
<td>750</td>
<td>MgMo$_2$O$_7$</td>
<td>2106</td>
</tr>
<tr>
<td>Co-Mo</td>
<td>800</td>
<td>Co$<em>x$Mg$</em>{1-x}$Mo$_4$</td>
<td>3064</td>
</tr>
<tr>
<td>Ni-Mo</td>
<td>800</td>
<td>MgMo$_2$O$_7$, Ni$<em>x$Mg$</em>{1-x}$Mo$_4$</td>
<td>3002</td>
</tr>
</tbody>
</table>

* Yield$_{MWCNT}$ (wt%) = (mass$_{MWCNT}$ + mass$_{Catalyst}$)/mass$_{Catalyst}$ * 100

High yields of MWCNTs are obtained by decomposition of methane in the presence of all the prepared sol-gel catalysts (see Table 2). In general Co-Mo catalysts show higher
catalytic performance than Ni-Mo catalysts, except for calcination treatments at 800 °C where both catalysts produce similar yields. It is also important to highlight that the highest yields directly correlate with a higher calcination temperature, and thus with the presence of the MgMoO₄ crystallographic phase in the catalyst. This again underlines the importance of the catalyst preparation conditions for the stabilization of certain catalytic phases (CoₓMg₁₋ₓMoO₄ and NiₓMg₁₋ₓMoO₄) and its relevance for the enhancement of MWCNT production yields.

Figure 6. SEM micrographs of MWCNT materials obtained with Co-Mo (a), Ni-Mo (b) catalysts.

SEM images of the as-synthesized carbon products (Figure 6) gives evidence of a homogeneous material consisting of high density of large curved bundles of several micrometers in length and several hundreds of nanometers in diameter. No noteworthy differences with respect to the type of metal and the calcination temperature are observed. The bundles consist of thin (3-5 nm) MWCNTs, and negligible amount of amorphous carbon (Figure 7). In general, Ni-Mo catalysts produce somewhat thicker MWCNT bundles ranging from 400 nm to 1 µm, in contrast to Co-Mo catalysts, which lead to average MWCNT bundle diameters below 300 nm.
Catalytic particles were in general difficult to find due to the high MWCNT yields obtained (elemental analyses indicate overall catalysts amounts of about 1 wt% in the produced MWCNT materials. However when observed, catalyst particles commonly appear as larger fragments attached to one extreme of MWCNT bundles. Interestingly, their lengths scale correlates to the size of the MWCNT bundle diameters (see Fig. 7a as example), which could be associated with a common starting growing plane for MWCNT growth.

![TEM micrographs of MWCNTs obtained from catalysts Co-Mo (a-c) and Ni-Mo (d-f).](image)

For all the produced materials, no RBM modes are detected in the corresponding Raman spectra (Figure 8), thus indicating the absence of SWNTs. However, typical D and G bands for MWCNTs are observed at 1400 cm\(^{-1}\) and 1600 cm\(^{-1}\), respectively. Looking at the intensity ratios of the D to G bands, no significant tendencies as a function of calcination temperature are found. Nevertheless, at all temperatures, MWCNTs produced by Co-Mo catalysts are characterized by higher I\(_G\)/I\(_D\) ratios,
indicative of higher degree of crystallinity compared to Ni-Mo catalyst systems. Even more important, in the region from 100 to 1000 cm$^{-1}$ none of the sharp peaks typical for the catalyst (see Figure 3) can be seen anymore. This once more shows, that catalyst materials are negligible in the produced materials, explains the difficulties to find any catalyst material in the final sample, and underlines the high production yields obtained.

Figure 8. Raman spectra of MWCNT materials produced by Co-Mo and Ni-Mo catalysts calcinated at 700ºC, 750ºC and 800ºC.

Figure 9. XRD patterns of MWCNT materials produced by Co-Mo (top) and Ni-Mo (down) catalysts at a calcination temperature of 700ºC. Symbols: Mo$_2$C (*), MgO (^), MWCNT (o).
X-Ray diffractograms of MWCNT produced by Ni-Mo and Co-Mo catalysts (Figure 9) show the typical broad peaks of MWCNTs as well as some minor peaks of molybdenum carbide and magnesium oxide. None of the sharp features typical for Ni or Co phases (see Figure 2) can be detected. Again, this is consistent with the Raman and TEM results confirming the negligible amount of metal particles in the final sample materials.

3.5. *MWCNT growth mechanism*

In previous sections, we have shown how an increase of the calcination temperature leads to crystallographic changes related to the formation and stabilization of crystallographic phases $\text{Me}_x\text{Mg}_{1-x}\text{MoO}_4$ (Me=Co, or Ni), relevant for the improvement of MWCNT production. Furthermore, the sol-gel method used here produces non-porous catalysts with laminar morphology, which are highly favourable for high yield MWCNT production. This is quite intriguing if one takes into account the general consideration that porous catalysts systems favour the dispersion state of the active catalyst nanoparticles, avoid massive sintering during the CVD process, and thus provide high surface areas and catalytic performance towards enhanced nanotube growth [32-36].

Having a non-porous catalyst system leading to high MWCNT yield production thus brings into focus its layered morphology, which apparently is an additional important factor to be taken into account. This will be discussed in the following by presenting a
growth mechanism for MWNCTs, which consistently explains the experimental findings of our work (Figure 10).

Four essential steps are considered. Step 1: Before the CVD reaction, Me$_x$Mg$_{1-x}$MoO$_4$ molybdates form a sol-sol solution. Once the reaction temperature is reached (1000ºC), a hydrogen flow is introduced into the reactor to reduce metal atoms (Co, Ni), which immediately arrange into catalytic active metal nanoparticles. Step 2: Reduced molybdenum oxide, MoO$_2$, is formed simultaneously and stabilises active metal nanoparticles, thus avoiding sintering into large metal cluster during the reduction process. Subsequently, methane introduced in the system starts to catalytically decompose, allowing carburization of the active metals and the available molybdenum as unstable and stable carbides, respectively. The correspondingly formed carbides act as a carbon sink, and regulate the carbon diffusion during the nucleation process. Step 3: Once saturation of unstable metal carbides is reached, nucleation stops and carbon precipitates in the form of MWCNTs in a typical base growth mechanism whereas the catalytic nanoparticles are still strongly attached to the support. Step 4: The directional mass diffusion of carbon into the form of MWCNTs at each nucleating active catalytic nanoparticle now may induce simultaneous stress between the layer-layer interfaces of the laminar catalytic system. These continued interlayer tensions could overcome weak interlayer attractive Van-der-Waals interactions leading to the exfoliation of individual layer-based catalyst domains.
Figure 10. Proposed growth mechanism in 4 steps: 1) Reduction to form catalytic active metal clusters. 2) Stabilization and nucleation of metal carbide nanoparticles. 3) MWCNT growth. 4) Exfoliation of platelet-like catalyst structure.

The exfoliation process then exposes fresh catalyst, and the cycle is repeated over and over again. In this way, all the potential nucleation sites of the layered catalyst system can effectively contribute to the formation of MWCNT in high yields. This growth mechanism is in agreement with the observation of wide MWCNT bundles attached at one extreme to catalyst fragments whose sizes are related to the bundle diameters. Large bundle diameters would be formed by Van-der-Waals interactions between thin MWCNTs, which are growing very densely due to the proximity of the nucleation sites within the catalyst fragments. Furthermore, the high aspect ratio of the produced MWCNTs underlines the high activity of the exfoliated catalyst domains allowing MWCNTs to easily reach micrometer lengths. In addition, XRD and Raman characterization of the produced materials confirm that the samples are composed (almost) exclusively of MWCNTs. The proposed model thus successfully explains MWCNT high yield production for these non-porous platelet-like catalyst systems.

Finally, the proper catalyst design also elucidates the effect of calcinations temperature on the MWCNT yields. At calcinations temperatures of 700 °C and 750 °C the important MeₓMg₁₋ₓMoO₄ (Me = Ni, Co) phase has not formed and massive sintering during nucleation becomes dominant due to a sub-optimal surface stabilization of the
catalytic active metals Ni and Co. Thus, this process limits the availability of active catalytic sites suitable for MWCNT growth, as well as the efficiency of the exfoliation cycle. On the contrary, deactivation is considerably diminished by the good dispersion and stability of active metal nanoparticles achieved once the Me$_x$Mg$_{1-x}$MoO$_4$ (Me = Ni, Co) phases are formed at catalyst calcinations temperature of 800 ºC.

4. Conclusions

Using sol-gel techniques, we have designed a set of non-porous platelet-like catalyst systems based on nickel and cobalt as active catalysts embedded in a magnesium oxide matrix. These have been used for the CVD growth of MWCNTs. By carefully controlling the calcination temperature during the catalyst preparation, we have established a relation between catalyst design and high yield MWCNT production. We have demonstrated that at a calcination temperature of 800 ºC catalysts stabilise in a crystallographic phase of Me$_x$Mg$_{1-x}$MoO$_4$ (Me = Ni, Co). We show that this phase is responsible for a high stabilization degree of the homogeneously dispersed active catalyst species, i.e. Ni and Co, and results in the formation of large bundles of MWCNTs in yields of more than 3000 wt% with respect to the initial catalyst. Finally, we propose a growth model based on the continuous exfoliation of platelet-like catalyst systems explaining the high yield MWCNT production achieved by non-porous catalysts. Our findings provide important insights for the rational design of highly active catalyst systems of great interest towards an effective and low-cost large-scale MWCNT production.
Acknowledgements

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References


Table Captions

Table 1: Catalyst metal loadings (at %).

Table 2: MWCNT production yields and main catalysts crystallographic phases at each catalyst calcination temperature.

Figure Captions

Figure 1. DSC-TGA thermograms of calcination process from 650°C to 950°C for Co-Mo and Ni-Mo catalysts. DSC (-), TGA (--).

Figure 2: XRD patterns of the Co-Mo, and Ni-Mo catalysts obtained at calcination temperatures of 700°C, 750°C, and 800°C. Symbols: MoO$_3$ (*), Mo$_4$O$_{11}$ (^), MgMo$_2$O$_7$ (o), MgMoO$_4$ (<>), CoMoO$_4$ (]), NiMoO$_4$ (x).

Figure 3: Raman spectra of Co-Mo and Ni-Mo catalysts obtained at calcination temperatures of 700, 750, and 800°C. Symbols: MoO$_3$ (*), MgMo$_2$O$_7$ (o), MgMoO$_4$ (<>), CoMoO$_4$ (]), NiMoO$_4$ (x).

Figure 4. SEM-EDX maps representing the dispersion of molybdenum (blue), magnesium (red), cobalt and nickel (green) of Co-Mo (a-d) and Ni-Mo (e-h) catalysts’ selected regions (indicated in a and c) at a calcination temperature of 700°C.

Figure 5: SEM (a,b) and TEM (c,d) micrographs of crystal structure obtained from catalysts Co-Mo (a,c), and Ni-Mo (b,d) at a calcination temperature of 800 °C. Insets show Fast Fourier Transforms of the selected region of catalyst platelets.

Figure 6: SEM micrographs of MWCNT materials obtained with Co-Mo (a), Ni-Mo (b) catalysts.
Figure 7: TEM micrographs of MWCNTs obtained from catalysts Co-Mo (a-c) and Ni-Mo (d-f).

Figure 8. Raman spectra of MWCNT materials produced by Co-Mo and Ni-Mo catalysts calcinated at 700ºC, 750ºC and 800ºC.

Figure 9. XRD patterns of MWCNT materials produced by Co-Mo and Ni-Mo catalysts at a calcination temperature of 700 ºC. Symbols: Mo$_2$C (*), MgO (^), MWCNT (o).

Figure 10: Proposed growth mechanism in 4 steps: 1) Reduction to form catalytic active metal clusters. 2) Stabilization and nucleation of metal carbides nanoparticles. 3) MWCNT growth. 4) Exfoliation of platelet-like catalyst structure.