Characterisation of commercially CVD grown multi-walled carbon nanotubes for paint applications

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1. Introduction

Multi-walled carbon nanotubes (MWNTs) have received much interest since they were brought to the attention of the wider research community in 1991 by Iijima [1] and although commercially available, have yet to become widely utilised by industry in large volume applications. MWNTs have extraordinary mechanical [2], electrical [3] and thermal [4] properties and are now being used in many research areas and commercial products; ranging from coatings [5–7] and composites [2,8,9] to batteries [10–12] and filters [13]. This work is part of a wider programme to assess the potential of MWNTs as pigmentation for matt black static dissipative coatings for the aerospace industry. For this reason it has been important to assess the characteristics of a selection of MWNTs that are available in kilogram quantities, at prices that would not make the cost of the end product prohibitively expensive. Although there are papers addressing the characterisation of various grades of MWNTs [14,15], there is a need for the characterisation of a wider range of commercially grown MWNTs using the same characterisation methodology.

In this study MWNTs, commercially grown via chemical vapour deposition (CVD) methods by Nanocyl™, Arkema and Bayer MaterialScience, are comprehensively characterised by scanning electron microscopy (SEM), energy dispersive X-ray spectroscopy (EDX), X-ray photoelectron spectroscopy (XPS), Raman spectroscopy, thermogravimetric analysis (TGA), surface area analysis by the Brunauer–Emmett–Teller (BET) method and non-local density functional theory (NLDFT). This characterisation will aid in the formulation of coatings and composites using MWNTs where attributes such as agglomerate structure and surface area affect the optimum dispersion method, processing time and dispersant loading.

MWNTs are not only available in their raw state and are often used in a purified or functionalised state. The most common purification and functionalisation methods for MWNTs involve further wet processing in acids [16,17] and length reduction of MWNTs has been achieved in various ways including oxidation with acids [18] and ball milling [16,19]. All of these techniques involve wet processing followed by washing and then drying of the nanotubes; in this study the restructuring of the MWNT agglomerates after processing is evaluated.

The most basic industrial coatings are a fluid blend of binder, solvents, additives and pigments. The inclusion of MWNTs into a...
coating can introduce electrical conductivity, physical reinforcement and other properties to the coating. These properties have a high dependency on the MWNTs remaining reasonably untangled and uniformly dispersed during storage and coating curing. All of these properties are directly influenced by the physical properties of the MWNTs that are investigated within this study.

2. Materials

MWNTs were sourced from Nanocyl™ S.A. (Belgium), Arkema (France) and Bayer MaterialScience AG (Germany). Details of the various grades are summarised in Table 1 along with properties as detailed in the suppliers’ technical data sheets [20–25].

3. Techniques

SEM was performed on a JEOL JSM–7100F field emission SEM equipped with a Thermo Scientific triple analysis system featuring an UltraDry EDX detector, used in this work, a MagnaRay wavelength dispersive X-ray (WDX) system and a QuasOr camera for electron backscatter diffraction (EBSD). An accelerating voltage of 8 kV was employed for SEM images and 15 kV for EDX analysis. For SEM, samples of each MWNT powder were sprinkled onto carbon adhesive tabs and a 2 nm gold coating was applied to reduce charging effects. For X-ray analysis, samples of each MWNT powder were sprinkled onto a fragment of silicon wafer. Three locations from three separate agglomerates were analysed by EDX for each MWNT sample at a magnification of 5000×. The mean result was then used to confirm the elements present. The substrate signal (SiKα) was excluded from the quantification.

XPS was performed using a Theta Probe spectrometer (Thermo Scientific) operating in constant analyzer energy (CAE) mode using a monochromated Al-Kα anode. Survey spectra were obtained from 0 to 1350 eV using a pass energy of 300 eV and a step size of 0.4 eV. High resolution spectra of the C1s, O1s and, where appropriate, N1s, Ca2p and Cl2p regions were obtained using a pass energy of 20 eV and a step size of 0.2 eV. Quantitative surface analysis was calculated from the high resolution core level spectra by peak fitting, following the removal of a non-linear Shirley background, using the manufacturer’s standard Avantage software.

Raman spectroscopy was performed on a Renishaw inVia micro-Raman spectrometer with a laser wavelength of 633 nm and a power of 1.67 mW measured at the sample. A 50× objective was used and the scan time of 10 s. Ten spectra were collected from different agglomerates for each MWNT sample. Peak fitting was performed using Origin Pro software.

TGA was performed on a TA Q500 (TA Instruments) from room temperature to 800 °C with a heating rate of 5 °C/min. The sample size was ca. 5 mg and the analysis was performed in air, and a balance gas of nitrogen, with flow rates of 60 ml/min and 40 ml/min respectively.

Nitrogen adsorption isotherms were measured using a Gemini V surface area analyser (Micromeritics) at 77 K. Specific surface areas (SSA) were calculated using the BET method [26] and the PSDs were calculated using the infinite slit NLDFh model [27,28] on the adsorption isotherm. The analysis software used was DataMaster from Micromeritics. Before analysis, samples were out-gassed under N2 for 24 h at 150 °C using FlowPrep 060 (Micromeritics).

4. Results and discussion

4.1. Appearance by SEM

Low magnification SEM micrographs reveal differences in the MWNT agglomerates that make up the supplied powder. NC7000 consist of ropes or yarns, ca. 2–3 μm in diameter, of highly entangled tubes showing some degree of alignment. These ropes then tangle to form larger, more loosely entangled agglomerates ranging in size from ca. 10 μm to 1 mm, individual ropes up to 40 μm in length from support to tip were observed. Both C150P and C100 resemble more ‘birds-nest’ type agglomerates showing no sign of ropes or alignment. These tubes are also highly entangled, with agglomerate dimensions in the same range as NC7000, but with the majority towards the higher end of the scale, with C100 being most consistent in size. SEM micrographs showing typical agglomerates from each sample are shown in Fig. 1.

For samples that have undergone further processing, such as purification, shortening or functionalisation, the agglomerate structure changes, as shown in Fig. 2. Nanocyl™ NC3100, a purified grade based of NC7000 appears to have similar sized agglomerates and still has signs of a rope structure, although these are much less clear than in NC7000 agglomerates, and the individual tubes seem more tightly entangled. NC3101, a carboxylic acid functionalised grade based on NC3100, appears to have a similar structure to NC3100, although the tightening of the agglomerates seems even more severe. NC3151, a shortened and then acid functionalised grade based on NC3100, consists of much smaller primary agglomerates and again, the individual tubes are even more tightly tangled together. Bayer C150HP, a purified grade of C150P, follows a similar change as that from NC7000 to NC3100 in that the agglomerates seem similar in size, but the entanglement has tightened.

Based on these observations there is an indication that wet processing steps lead to a tightening of nanotube agglomerates and that the process Nanocyl™ use to shorten their MWNTs leads to a reduction in agglomerate size.

4.2. Bulk composition by EDX

It is well known that most MWNTs carry a small concentration of catalyst residue within their structure. EDX was used to determine the composition of the metal catalysts used in the synthesis of the MWNTs and to identify any contaminants present. The spectra obtained for NC7000 is shown in Fig. 3, catalysts residues are readily observed with an intense AlKα signal and weak signals of FeKα and CoKα.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>A Summary of the MWNTs investigated and properties supplied by the manufacturers.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Product name</td>
<td>Description</td>
</tr>
<tr>
<td>Nanocyl™ NC7000</td>
<td>Industrial grade</td>
</tr>
<tr>
<td>Nanocyl™ NC3100</td>
<td>Purified grade</td>
</tr>
<tr>
<td>Nanocyl™ NC3150</td>
<td>Purified and shortened</td>
</tr>
<tr>
<td>Nanocyl™ NC3101</td>
<td>Purified and COOH functionalised grade</td>
</tr>
<tr>
<td>Nanocyl™ NC3151</td>
<td>Purified, shortened and COOH functionalised grade</td>
</tr>
<tr>
<td>Nanocyl™ NC3152</td>
<td>Purified, shortened and NH3 functionalised grade</td>
</tr>
<tr>
<td>Arkema C100</td>
<td>Industrial grade</td>
</tr>
<tr>
<td>Bayer C150P</td>
<td>Industrial grade</td>
</tr>
<tr>
<td>Bayer C150HP</td>
<td>Purified grade</td>
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Table 2

<table>
<thead>
<tr>
<th>Main impurities (wt%)</th>
<th>Trace impurities</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC7000</td>
<td>Al (5.9) Fe (0.5) Co (0.2)</td>
</tr>
<tr>
<td>NC100</td>
<td>Fe (0.2) Co (0.1)</td>
</tr>
<tr>
<td>NC3150</td>
<td>Al (0.1) Fe (0.2) Co (0.1)</td>
</tr>
<tr>
<td>NC3101</td>
<td>Fe (0.5) Co (0.1) Cl (0.9) Mg (0.3) Ca (2.6) Mn (0.1)</td>
</tr>
<tr>
<td>NC3151</td>
<td>Al (0.1) Fe (0.2) Cl (2.4) Mg (0.3) Na (0.1) Mn (0.4)</td>
</tr>
<tr>
<td>NC3152</td>
<td>Ca (1.5)</td>
</tr>
<tr>
<td>C100</td>
<td>Al (1.2) Fe (4.5)</td>
</tr>
<tr>
<td>C150P</td>
<td>Al (0.5) Mg (0.5) Mn (0.5) Co (0.5)</td>
</tr>
<tr>
<td>C150HP</td>
<td>Co (1.2) Cl (0.3)</td>
</tr>
</tbody>
</table>

Table 2 summarises the elements detected for each CNT powder. All quantities are based on the mean analysis of three separate area scans on three different agglomerates. Only elements detected in all three analysis areas are shown as main impurities, with mean values higher than 0.1% by weight. It should be noted that some elements are present very close to the sensitivity limit of EDX and that the samples were not flat or dense, making them less than ideal samples for EDX analysis. NC7000 shows a large amount of aluminium and smaller quantities of both iron and cobalt. C150P contains mainly cobalt, manganese, magnesium and aluminium and C100 mainly iron and aluminium. MWNTs are routinely grown from iron, cobalt and manganese catalysts and aluminium oxide and magnesium oxide are used as catalyst supports [29–32], which explains the presence of these elements. These elements are also mentioned in various patents from Nanocyl™ [33,34], Bayer MaterialScience [35,36], and Arkema [37].

When purified to NC3100 the large quantity of aluminium oxide present in NC7000 reduces to just trace amounts, as does the cobalt concentration. Traces of sulphur are also present, indicating that the purification step used by Nanocyl™ probably involves the use of sulphuric acid. Purification of both Nanocyl™ and Bayer industrial grades results in a large reduction in catalyst and catalyst support, but appears more effective at removing the support than the metal catalyst particles; thereby indicating some catalyst may be located inside the nanotubes, as reported by Pumera [38]. Nanotubes that have undergone further processing contain additional contaminants including chlorine, sulphur, calcium and magnesium.

EDX elemental mapping shows the distributions of the metal catalysts and catalyst support within the agglomerates and the results for the three industrial grades are shown in Fig. 4. The overlapping of the oxygen and aluminium atoms confirms the support material for all three industrial grades and iron (NC7000 and C100) and cobalt (C150P) maps confirm the finely dispersed catalyst particles.

Fig. 1. SEM micrographs showing the agglomerate size and structure of the industrial grade MWNTs; NC7000 (a, b), C100 (c, d) and C150P (e, f).
A good review of nanotube growth is given by Zhang et al. [39] who divided nanotube growth into two types: either aligned or agglomerated. Typical powder catalysts used in nanotube synthesis consist of metal nanoparticles on a support. When a source of carbon is introduced (for example methane), carbon nanotubes grow and as they grow they crush and eventually rupture the catalyst particles, dispersing them throughout the nanotube agglomerate that is formed. This is the method of growth used to produce Baytubes® [40] and explains the ‘birds nest structure’ observed by SEM and the finely dispersed catalyst seen in the EDX maps for Bayer C150P (Fig. 4). As the SEM micrographs and EDX maps for C100 are very similar to C150P, it is likely they also grow in this way. Nanocyl® NC7000 contains distinctive support particles of aluminium oxide which have not been crushed and finely dispersed throughout the agglomerates and the EDX maps show ropes growing from both sides of these well defined support particles, making them very different from both C100 and C150P. The SEM micrograph and EDX maps shown in Fig. 5 clearly show the primary structure of NC7000 as highly entangled ropes grown from an aluminium oxide support. The highly entangled nature of the ropes may be explained by the surface of the aluminium oxide support; Atthipalli et al. [41] showed that the surface roughness of the
Fig. 3. EDX spectra for NC7000.

Fig. 4. EDX elemental maps for the three industrial grades of MWNTs.
support used to grow carbon nanotubes reduces alignment and a patent from Nanocyl™ [33] describing catalyst production refers to milling and sieving of support powder that would not lead to a smooth surface.

4.3. Surface analysis by XPS

XPS reveals the surface chemistry of the MWNTs. The survey spectra for NC7000, NC3150 and NC3151 are shown in Fig. 6. The degree of graphitisation and oxidation can be measured using the C1s peak and O1s peak, respectively.

The spectra are dominated by the C1s peak at 284.8 eV with a small O1s peak at 532 eV that increases and broadens in the acid functionalised grades. The O1s peak at 532 eV corresponds to the carbonyl bond in an acid or ester [42,43]. The results of the quantitative surface analysis are summarised in Table 3.

All MWNT grades have oxygen on the surface, likely as a result of the catalytic growth process; the quantity of oxygen is relatively low in the raw industrial grades, although for NC7000 is slightly higher than C100 and C150P. While this could be a result of the high level of aluminium oxide in NC7000, aluminium was not detected by XPS. There is no increase in oxygen when comparing NC7000 to NC3100, which would indicate that the purification process used by Nanocyl™ does not impact the degree of oxidation of the nanotubes. Shortening of the MWNTs may increase the degree of oxidation of the MWNT, which is expected as the tubes are broken, but this cannot be shown definitively as any small increase or decrease could be from batch-to-batch variation. For the acid functionalised grades from Nanocyl™, there is a clear increase in oxygen surface concentration to 11.6% and 12.9% for purified and purified-shortened grades respectively, confirming the addition of oxygen containing groups on the nanotube surface.

4.4. Raman spectroscopy

Raman spectroscopy is widely used to provide information relating to the quality of nanotubes, although most work relates to single walled nanotubes with MWNTs being less well understood. The main features of a Raman spectra for a graphitic material are described in detail by Dresselhaus et al. [44] and Farrari et al. [45] but for MWNTs the spectra consist mainly of the D band (≈1350 cm⁻¹), G band (≈1580 cm⁻¹), and 2D (≈2700 cm⁻¹) band, sometimes called G' band. The disorder induced D band relates to vibrations from defects present in a carbon structure. The G band relates to the in-plane vibration of an ideal sp² bonded carbon structure and the 2D band is the first overtone of the D band and is always present in a graphitic sample and is used to infer long range order in a carbon based structure.

The ratio of I_D/I_G has long been used as an indication of the quality of nanotubes with an increase in the ratio, meaning more defective nanotubes or the presence of more amorphous carbon in the sample. Behler et al. [46] observed a change in the position and intensity of the D band with changing excitation wavelength, making it difficult to standardise and compare I_D/I_G ratios obtained using different wavelengths. DiLeo et al. [47] proposed that defects would reduce the double resonance process that results in the decreasing of the 2D band, making the I_D/I_G ratio a better indicator of crystallinity in MWNTs. Another advantage to using the 2D band, as opposed the D band, is that the 2D band is always present and satisfies the Raman selection rule [48] where the D band is forbidden and requires defects to be present [49]. Other researchers have used the ratio of the areas of the peaks rather than the intensities to infer quality [50].

Fig. 7 shows a typical Raman spectra for NC7000 with a relatively intense D band and a G band of lower intensity, typical of a highly defective structure [51].

Table 4 summarises the ratios of the intensities as peak heights (I) and areas (A) of the D band to G band and the 2D band to the G band.

All ratios are similar to each other, with only C100 showing a slightly lower D/G band ratio and slightly higher 2D/G band ratio, inferring lower numbers of defects and higher crystallinity. As discussed, it can be difficult to directly compare Raman spectra and thus band ratios obtained using different equipment, but these similarities are in agreement with Tessonnier et al. [14] who showed NC3100 and C150P had similar I_D/I_G ratios of 2.1 and 1.8.
respectively. Due to the similarity of the D/G band and 2D/G band ratios, it can be concluded that the purification processes used by both Nanocyl™ and Bayer MaterialScience and the functionalisation process used by Nanocyl™ do not increase the number of defects within the nanotubes.

4.5. Surface area and pore size distribution

SSA is an important parameter for a pigment as it provides an indication to the amount of binder, or dispersant, that is required to fully wet the particle surface. The pore size distribution was investigated to detect the differences in agglomerate structure observed via SEM.

All nanotubes exhibit Type IV BET isotherms with Type II characteristics, according to the IUPAC classification[52,53]. This indicates a mesoporous material with possible small macropores. Isotherms for various Nanocyl™ grades are shown in Fig. 8.

The SSA and total pore volume for each nanotube powder are summarised in Table 5.

Industrial grade nanotubes from Nanocyl™ have the highest SSA at 475 m² g⁻¹ followed by Arkema with 249 m² g⁻¹ and lastly Bayer with 210 m² g⁻¹. Peigney et al. [54] showed that nanotube surface area was related to nanotube diameter and number of walls and this is likely the source of the difference between these samples. This is in agreement with the manufacturers data, showing C100 and C150P having similar diameters and numbers of walls. The similarity in diameter was confirmed by Rosca et al. [8]. Nanocyl™ report a narrower diameter for NC7000, but do not specify this as the external diameter [20]. Tessonnier et al. [14] measured the SSA of NC3100, which are based on NC7000, and found it to be significantly higher than C150P and also observed that C150P had more wall terminations and more regular closing of the internal channels, drastically reducing the accessible internal surface area and thus explaining the higher SSA measured for NC7000 in this study. The SSA for NC3100 measured in this study was very similar to the 334 m² g⁻¹ measured by Tessonnier et al. but the SSA of C150P measured in this study is lower than the 228 m² g⁻¹ measured by Tessonnier et al. This is likely due to batch-to-batch variation, as an increased amount of residual catalyst would lead to an overestimation of the SSA of a MWNT sample. The residual catalyst measured by Tessonnier et al. using TGA for C150P was 4% compared to 2.9% in this study, which would account for the higher SSA.

When comparing Nanocyl™ purified MWNTs to their industrial grade equivalent, there is a reduction in SSA from 476 m² g⁻¹ to 323 m² g⁻¹ for NC7000 and NC3100, respectively. This decrease is likely due to either the removal of the catalyst support, which will have had a high surface area, or batch variation. Purified and shortened MWNTs from Nanocyl™ (NC3150) have a higher surface area than purified MWNTs, which is due to the shortening of the tubes making the internal channels within the tubes more accessible. The small increase in SSA when comparing C150P to C150HP, which has undergone a purification process, could be for the same reason. The functionalised grades from Nanocyl™ have lower SSAs than their un-functionalised equivalents, but they still show the same rise in SSA when the tubes are shortened.

The PSD for the industrial grade MWNTs are shown in Fig. 9 and show a bi-modal distribution. The first distribution (3–20 nm) falls within the diameter dimensions for the MWNTs when considering inner tube diameter, inner channel diameter and outer tube diameter and likely corresponds to pores attributed to individual tubes. The second distribution corresponds to spaces between tubes and is a result of the bundled nature of the nanotube agglomerates.
If the pore volume contributions are separated in relation to these distributions, an insight into the pore volume from within the tubes and from the agglomerates can be achieved.

Taking the Nanocyl™ range as an example, Fig. 10 shows a clear shift in the second distribution towards smaller pores after the nanotubes have been purified, shortened and functionalised, indicating a restructuring in the agglomerates in agreement with the observation from the SEM micrographs. The most common purification methods involve acid solutions [16,17], indicating the tubes are wet-processed and subsequently dried, which would explain this change in the agglomerate structure. For shortened tubes this change is even more pronounced and the shift leads to a combining of the two distributions, which is due to shorter tubes being able to pack more densely. Functionalisation, which involves further processing, follows this trend and the distributions not only combine but also shift even further towards smaller pores in the shortened grades.

Fig. 11 shows the cumulative pore volume for the two pore size ranges, for the MWNTs considered in this work. The volume in the pores with diameters <20 nm remains relatively consistent across nanotube samples, although there is an increase from NC7000 to NC3150, which again can be attributed to either tube opening or enlargement of defects. The same rise is present for NC3151 and NC3101, but not in NC3100, implying that the functionalisation process opens the tube ends or enlarges defects regardless of whether the tubes have been shortened or not. This same increase is observed when comparing C150P and C150HP, indicating either tube opening or defect enlargement or a combination of the two during Bayer’s purification process. When considering the cumulative volume from larger pores in the agglomerate structure, it is clear that with increased processing steps, pore volume drops considerably. This is true for both Nanocyl™ and Bayer grades and shows the tightening of the agglomerates reducing the volume between the tubes, in agreement with SEM micrographs. Using this method to evaluate how tightly entangled agglomerates are, the PSD data indicate that NC7000 is more loosely agglomerated than C100, which is more loosely agglomerated than C150P, in agreement with Krause et al. [15] who found C150P required five times more energy to disperse than NC7000. In contrast, Tessonnier et al. [14] predicted that C150P was more loosely agglomerated than NC3100, which is based on NC7000, as less volume from larger pores was present. It is worth noting that NC7000 has two agglomerate structures: primary tightly agglomerated ropes and more loosely agglomerated bundles of ropes. The pore size distribution is an amalgamation of these two structures.

4.6 Thermogravimetric analysis

TGA is often used to characterise the oxidation behaviour of MWNTs, as well as in the determination of the residual catalyst content [16,17,19,55]. Table 6 summarises the minimum residual catalyst for each sample after 800 °C; the temperature to 50% mass (TSD) and the temperature of maximum combustion rate (TMAX) taken from the derivative of the TGA curve. These values give an indication of the rate of oxidation and the purity and uniformity of the samples.

It is well documented that oxidation temperature relates to residual catalyst and defects present in the structure of carbon nanotubes [56]. From the Raman analysis it was shown that all powders are similar in terms of defects with all having relatively high levels. MWNTs from Bayer combat at a lower temperature than those from Nanocyl™, with NC7000 and C150P having TMAX of 579 °C and 543 °C respectively. This can be seen more clearly in Fig. 12. For both manufacturers, purification increases oxidation temperature but the increase in TMAX is larger for Bayer than for Nanocyl™ nanotubes; NC7000 to NC3100 has an increase in TMAX of 19 °C where C150P to C150HP has an increase of 38 °C. Although the residual catalyst for NC7000 is higher than all other samples, EDX has shown that the majority of that is aluminium oxide catalyst support rather than metal catalyst particles. For C150P, more of

![Table 6](image)

<table>
<thead>
<tr>
<th>Residual catalyst (as % of initial mass), TSD and TMAX of the MWNT powders.</th>
<th>Residual (%)</th>
<th>TSD (°C)</th>
<th>TMAX (°C)</th>
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<tbody>
<tr>
<td>NC7000</td>
<td>11.9</td>
<td>569</td>
<td>579</td>
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<tr>
<td>NC3100</td>
<td>2.5</td>
<td>585</td>
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<td>NC3150</td>
<td>2.5</td>
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<tr>
<td>C150HP</td>
<td>1.8</td>
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</tr>
</tbody>
</table>

![Fig. 10](image)

**Fig. 10.** PSDs for the various grades of MWNTs from Nanocyl™ showing a reduction in pore size with increased processing.

![Fig. 11](image)

**Fig. 11.** Pore volumes for each distribution showing a decrease in volume with further processing.
the residual is metal catalyst (manganese and cobalt), which would explain the lower oxidation temperature. C100 oxidises at a temperature much closer to that of NC7000 than that of C150P even though C100 does have a large amount of residual iron catalyst.

The EDX analysis for C100 indicates no cobalt catalyst and both NC7000 and C150P use cobalt as a co-catalyst with C150P showing higher amounts than NC7000, which implies that the oxidation temperature is more affected by the amount of residual cobalt in the samples than residual support material or iron. For the further processed Nanocyl™ grades, there is a reduction in $T_{\text{max}}$ of 10 °C from NC3100 to NC3150, which is likely caused by the increase in tube ends and increase in accessible surface area shown by nitrogen sorption data. Functionalised grades show a drop in mass below 300 °C, which is the oxidation of the functional groups [57], leaving exposed defects and hotspots reducing the $T_{\text{max}}$ by 80 °C and 100 °C for NC3101 and NC3151 from their un-functionalised equivalents.

EDX suggests that differences in structure of the agglomerates come from differences in growth conditions, with NC7000 being grown from platelets of aluminium oxide, while those of C100 and C150P are grown from a support that disintegrates as the tubes grow. EDX confirms the catalysts and support used during growth and shows that although NC7000 has the highest level of impurities, the majority is aluminium oxide support rather than transition metal catalyst. The high level of catalyst, as well as being shown to affect the thermal properties of the MWNTs, will also have to be considered when incorporating MWNTs in resin systems. The relatively large support platelets of aluminium oxide observed in NC7000 will be present in any cured resin system and must be stabilised. The small sizes of the disintegrated catalyst support of C100 and C150P will likely have little effect on the coating or composite they are dispersed into. If the aluminium oxide particles of NC7000 do give undesired properties to cured resin systems, the purified grade (NC3100) would be a good alternative.

Raman spectroscopy gives similar D/G and 2D/G ratios implying all grades are similar to each other in terms of quality. There is no indication that the purification processes used by Nanocyl™ or Bayer MaterialScience, or the functionalisation process used by Nanocyl™, increase the number of defects within the nanotubes.

TGA shows a higher thermal stability for Nanocyl™ grades when compared to Arkema or Bayer and this is thought to be due to the higher level of residual transition metal catalyst in the latter samples. TGA and EDX also confirmed the purified grades from Nanocyl™ (NC3100) and Bayer (C150HP) contain lower levels of residual catalyst and that the purification process appears more effective on the catalyst support than the transition metal catalyst itself. These differences in thermal stability could cause different changes in the thermal stability of any coating they are dispersed into. These results also highlight thermal stability is not always directly related to purity and that an understanding of the nature of any impurities is important.

A tightening of the agglomerate structures during processing, seen by SEM, is confirmed using BET and NLDFT and shows massive decreases in the volume of the larger pores, attributed to the agglomerate structure, following wet chemical treatment. As a result, an indication of agglomerate structure can be obtained on a larger sample size using BET and NLDFT than with SEM.

XPS shows that all grades have a low concentration of oxygen on the surface from the CVD growth process and that COOH functionalised grades from Nanocyl™ (NC3101 and NC3151) have increases of circa 10% oxygen on their surfaces. This functional surface should lead to better compatibility and stability in specific resin systems.

6. Conclusions

Commercially CVD grown MWNTs are supplied with various agglomerate structures and these structures are dependent on the growth mechanisms used. Wet processing of MWNTs changes agglomerate structure and leads to a tightening of the MWNT bundles. Residual catalyst left from nanotube growth has an effect on the thermal stability of the MWNT powder, with residual transition metal having a larger impact on oxidation temperature than the support material. All samples investigated in this body of work were shown to be of similar quality with regards to crystallinity.

Acknowledgements

This research was supported by the Engineering and Physical Science Research Council (EPSRC) through the Centre for Doctoral Training (CDT) in Micro- and NanoMaterials and Technologies (MiNMat) at the University of Surrey and Indestructible Paint Ltd [EPSRC Grant reference number: EP/G037388/1].

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