Thermosetting nanocomposites with high carbon nanotube loadings processed by a scalable powder based method

Tomi M. Herceg\textsuperscript{a,b,c}, Sung-Ho Yoon\textsuperscript{c,d}, M. Shukur Zainol Abidin\textsuperscript{c,e}, Emile S. Greenhalgh\textsuperscript{c,e}, Alexander Bismarck\textsuperscript{b,c,f}, Milo S.P. Shaffer\textsuperscript{a,c}

\textsuperscript{a} Department of Chemistry, Imperial College London, UK
\textsuperscript{b} Department of Chemical Engineering, Imperial College London, UK
\textsuperscript{c} The Composites Centre, Imperial College London, UK
\textsuperscript{d} Solid Mechanics and Materials Engineering, Oxford, UK
\textsuperscript{e} Department of Aeronautics, Imperial College London, UK
\textsuperscript{f} Institute for Materials Chemistry and Research, University of Vienna, Austria

Abstract
A powder based processing route was developed to allow manufacturing of thermosetting nanocomposites with high (20 wt\%) carbon nanotube (CNT) loading fractions. Adaptation of high shear mixing methods, as used in thermoplastic processing, ensured that the CNTs were well distributed and dispersed even at the highest loadings. By minimising flow distances, compression moulding of powders ensured that the CNTs did not agglomerate during consolidation, and yielded a percolated CNT network in a nanocomposite with excellent electrical and thermal conductivities of 67 S m\(^{-1}\) and 0.77 W m\(^{-1}\) K\(^{-1}\), respectively. Unusually, the CNTs provided effective mechanical reinforcement at even the highest loadings; embrittlement is minimised by avoiding large scale inhomogeneities and the maximum measured Young’s modulus (5.4 GPa) and yield strength (90 MPa) could make the nanocomposite an attractive matrix for continuous fibre composites. The macromechanical measurements were interpolated using micromechanical models that were previously successfully applied at the nanoscale.

1. Introduction
Carbon nanotubes (CNTs) have been scrutinized for over a decade as a constituent in thermoset based nanocomposites (NCs) due to their exceptional intrinsic multifunctional
properties [1]. One of the major challenges in the field has been to distribute and disperse high loading fractions of CNTs in epoxy resins by a route that is scalable to high throughput. The choice of processing route is critical to determining the microstructure and hence properties of the composite; agglomeration of CNTs is often particularly problematic [2].

Large CNT loadings have been relatively easily achieved in thermoplastics by shear mixing [3, 4], as they benefit from broad processing windows. For thermosets, traditional dispersion techniques such as simple stirring and sonication have produced good quality NCs for low CNT loadings, but have been ineffective at CNT volume fractions beyond a few percent [5-7]. At high loadings, techniques such as layer-by-layer (LBL) deposition [8], resin infiltration of buckypapers [9], drawing, aligning and stacking CNT sheets [10] or resin infusing continuous and aligned CNT forests [11, 12] have been successfully employed to manufacture high quality thermosetting composites. In one promising example, highly aligned and crystalline CNT arrays were infused with epoxy and shown to be an effective reinforcement (16.5 wt%), improving stiffness from 2.5 to 20.5 GPa, strength from 89 to 231 MPa, and electrical conductivity ($\sigma_{DC}$) similar to that of amorphous carbon ($\sim 10^3$ Sm$^{-1}$) [10].

Three roll milling has been identified as a promising approach for dispersing high fractions of CNTs in epoxy resins in large volumes [13]. This technique has been successfully used to produce NCs containing up to 5 wt% multi walled carbon nanotubes (MWCNTs) that also have enhanced mechanical, electrical and thermal properties [14]. The highest CNT loading reported in epoxy with this method is 8 wt% [15], but since the aim was maximum $\sigma_{DC}$, the NC was only moderately sheared to allow the formation of a dense, interconnected CNT network, which is not optimal for mechanical performance. However, in principle, mixing CNTs and thermosets using large shear forces as done in thermoplastic based NCs could yield good CNT dispersions at high loadings.
This work draws on the potential of readily scalable, shear mixing techniques to make excellent isotropic dispersions of CNTs in epoxy matrices. The aim was to avoid agglomeration on the 10-100 μm scale, which is typically the limitation to introducing large CNT fractions in the resin. The approach uses thermoplastic processing techniques, specifically extrusion; the excellent dispersion was maintained during consolidation by minimising the flow distance using a powder-based, vacuum-assisted compression moulding technique. The resulting NCs were characterised mechanically, electrically and thermally to determine the success of the processing route and the resulting properties.

2. Experimental

2.1 Materials and nanocomposite preparation
The CNTs (NC 7000, Nanocyl™) were industrial grade, highly entangled and catalytically grown. According to the manufacturer’s datasheet, the CNTs were about 1.5 μm long, 10 nm in diameter and contained 10 wt% metal oxide catalyst; the catalyst content was confirmed by thermogravimetric analysis (TGA) in air. Raman spectroscopy confirmed [16] a large proportion of defects in the otherwise graphitic walls ($I_G/I_D = 0.86$). The matrix consists of a Bisphenol-A and Bisphenol-F epoxy blend (EPIKOTE™ 1001) cured using a dicyandiamide (Dyhard® 100S) hardener with an average particle size of 10 μm. Both components were generously provided by Hexcel Composites (Duxford, UK).

When choosing this resin system, there were two considerations. The first was to upgrade the properties of a low-cost resin so that it could be used for structural applications. The second was that the NC should be moulded as a fine powder in order to minimise flow distances during consolidation thus minimising re-agglomeration and difficulties associated with high viscosities. EPIKOTE™ 1001 was chosen because it is a relatively inexpensive coating resin and remained as a powder at ambient temperatures rather than fusing. Dicyandiamide
(DICY), a latent curing agent, provides a broad processing window so that CNTs can be initially mixed into the epoxy at high temperature and shear without the undesired onset of gelation [17].

DICY was weighed out at 4% by weight of resin and 2.4, 4.8, 9.1, 13 and 20% CNTs by weight of the composite. Control samples were prepared without any CNTs. The raw materials were mixed using a co-rotating twin screw extruder (PRISM TSE 16 TC, Thermo Scientific) in two passes at speeds of 100-200 RPM (residence time of 2-5 minutes) and barrel temperature of 70-125°C to maintain constant torque.

The NC product was then ground into a powder using a cryogenic ball mill (Cryomill, Retsch). The milling process was adjusted by varying the duration and frequency of the grinding cycles to reduce the median powder particle size, and obtain a narrow particle size distribution. The powder particle sizes were measured from surfactant stabilised suspensions (0.5% Triton X-100 by weight of water) in deionised water using a laser diffractometer (Mastersizer 2000, Malvern Instruments), which produces a distribution of particle diameters based on refraction angles, with larger particles having smaller refraction angles.

The powder was consolidated by compression moulding into dogbones according to the ASTM D638 Type V specification with a 3 x 3 mm² gauge section. The mould assembly containing the powder was covered with peel ply, breather cloth, and a vacuum bagging sheet sealed to a metal plate with tacky tape. Vacuum was applied for 30 min to remove air, then held while the mould was heated to 125°C on the platen of a hydraulic hot press (Moore Presses) to allow the powder to melt. Once the desired temperature was reached, 14.5 bar of pressure was applied and the mould was heated to 150°C to cure for 12 h. The control sample was cured following the same temperature profile, but in a vacuum oven and without a
plunger since the low viscosity epoxy melt flowed easily; overflow during degassing was thus prevented.

2.2 Nanotube and nanocomposite microscopical characterisation

SEM (LEO Gemini 1525 FEG, Carl Zeiss) and TEM (2100, JEOL) of over 100 CNTs were used to determine their length and diameter distributions, respectively. Samples for TEM were prepared by dipping a holey carbon grid into an ethanol solution containing as-received nanotubes dispersed by bath sonication (USC300T, VWR), and imaged at an accelerating voltage of 200 kV in bright field. Samples for measuring CNT lengths by SEM were prepared by dissolving in acetone the uncured resin from NCs that had been extruded and cryomilled, filtering (0.1 μm hydrophilic PTFE membrane, Millipore) and washing the nanotubes with acetone (ACS, VWR), suspending them in dimethylformamide (DMF) (Rectapur, VWR) by bath sonication and pipetting the suspension onto polished SEM stubs. Images were collected at an accelerating voltage of 10 kV, and analysed with the ImageJ software.

SEM of fractured tensile samples was also used to characterise the dispersion and distribution of the CNTs in the epoxy system and measure their pull-out length. Fractured samples were mounted onto aluminium stubs and sputter coated with a 5 nm thick layer of chromium to prevent charging. The specimens with the highest CNT loading were sufficiently conductive and were not sputter coated so as to allow observation of the CNT-resin interface. Samples were imaged at an accelerating voltage of 3 kV.

The morphology of the NC microstructure was characterised by differential interference contrast (DIC) optical reflective microscopy (AX10 with AxioVision, Zeiss) using fragments from the grips of tensile test specimens that were mounted in epoxy, then ground and polished to 1μm with a rotary polishing machine (Motopol 12, Buehler).
2.3 Chemical and physical characterisation
The composition of the NCs was determined by TGA (Pyris 1, PerkinElmer) in N₂ at a ramp rate of 5°C/min from 100 to 850°C using segments taken from spent mechanical test specimens. Densities of the NC dogbones were measured by the gas displacement method with a pycnometer (Accupyc 1330, Micrometrics GmbH).

The curing behaviour of the NC was investigated using differential scanning calorimetry (DSC) (TA Q2000, TA Instruments). The extent of reaction was determined from integrating the heat flow against time as measured from isothermal scans at 150°C [18] and normalising it to the integrated heat flow of the control sample. The heat flows were normalised for the mass fraction of resin, as the CNTs are not believed to participate in the crosslinking reaction. The glass transition temperature (T_g) of the cured NCs was measured from dynamic DSC scans at 10°C/min between 20 and 150°C.

Samples for conductivity measurements were sectioned from the grip regions of spent mechanical test specimens and polished to remove gripping damage. The electrical conductivities in three axes were deduced from volume resistances measured with a two point probe and an ohm meter (DM9C, Amprobe). Silver paste was applied to the probe and the contact points on each sample to minimise contact resistance. The thermal conductivity (k) was calculated from the thermal diffusivity measured using a Xenon light flash system (LFA 447 Nanoflash®, Netzsch Instruments, Inc.).

2.4 Nanocomposite mechanical characterisation
The NC dog bone samples were polished to remove any surface imperfections present from the moulding process. Strain gauges (FLK-1-11, TML) were glued to the working region of the dog bones with cyanoacrylate glue before tensile testing in an Instron 4505 screw driven machine at a speed of 0.5mm/min. At least five samples were tested for each CNT loading.
The fracture surfaces were inspected visually for gross defects that would have adversely impacted strength and strain-to-failure results.

Preliminary fracture tests were performed on the neat resin according to ASTM D5045 in the single-edge notch bend (SENB) configuration using three 40 mm x 10 mm x 5 mm specimens that were initially machined with a 45° disc cutter to 0.45 of the width and further notched with a razor blade. The final crack length was approximately half the width and accurately measured by microscopically inspecting the fractured specimens. The compressive load was applied using a 3-point bending jig at 0.5 mm min⁻¹.

3. Results and discussion

3.1 Nanocomposite production and composition

The CNTs and thermosetting matrix were mixed by extrusion as it provides high shear, and is a continuous process that is readily scalable in volume. With careful monitoring of temperatures along the barrel of the extruder, runaway exotherms were avoided. The consistency of the extruded NC changed from a thin paste to pellets, with increasing CNT content. The highest CNT loading successfully extruded was 20 wt%/ 13.6 vol%, more than double the highest reported to date (8 wt%) [15] for a thermosetting NC prepared by shear mixing. Typically, forming an extrudate with high CNT loadings into test specimens requires the NC to flow over large distances relative to the size of the CNT filler, producing agglomerates [19, 20]. As such, powdering the NC reduced the flow distances when consolidating under pressure, while curing under vacuum removed air pockets that would otherwise yield a composite with voids.

The extruded NC was initially milled using two long grinding cycles of two minutes with a long re-cooling period in between of two minutes (blue, dashed curve in Figure 1a). This method proved ineffective as it yielded powder with a large average particle size (~60 μm)
and broad distribution. Systematic variation of the grinding parameters showed that grinding cycles of 10 s and re-cooling for 40 s between them yielded powders (Figure 1b) with the smallest average particle size of 20 μm (red, solid curve in Figure 1a). Powder particles presumably experienced less frictional heating during shorter grinding cycles and were sufficiently cooled in between cycles to avoid reaching temperatures exceeding T_g and fusing into aggregates. In principle, more rapid cycles might be beneficial, but were not possible on the equipment used. The length distributions of the CNTs were measured after extrusion and grinding of the NCs (Figure 2). Interestingly, the lognormal distributions are equivalent within error across the range of CNT loadings, despite the variable viscosity, probably because the extrusion temperature was adjusted to maintain a similar torque during extrusion and/or as a result of the grinding step. Length distributions are not easily measured after curing, but it is reasonable to assume that there is little further change during consolidation, and an average CNT length (430 ± 250 nm) was assumed in further analysis.

![Figure 1. a) Particle size distribution and b) morphology of cryogenically milled powder containing 20 wt% CNTs.](image)

---

8
Macroscopically, the NC specimens were well formed across all CNT loadings (Figure 3a), with a good surface finish and no apparent defects. TGA analysis confirmed intended CNT weight fractions in the resin within the 1-2% error of the TGA; no changes in thermal stability were detected (Figure 3b). Whilst weight fractions can be measured during sample preparation and checked with TGA, CNT volume fractions are more relevant to most property comparisons. In principle, the conversion may not necessarily be straightforward, since the surface interactions involve a significant volume of the matrix, and the fate of the hollow nanotube core is uncertain; nevertheless, within the experimental error, CNT volume fractions in the NCs were adequately estimated using the measured matrix density of 1.2 g/cm³ and a CNT density of 1.9 g/cm³, which was deduced from a CNT density of 1.75 g/cm³ [21] corrected for the 10 wt% metal catalyst content. The CNTs themselves could not be sufficiently compacted to fill at least one third of the pycnometer chamber and ensure an accurate direct density measurement.
Figure 3. a) Well consolidated NC containing 13.6 vol% CNTs, b) TGA of NCs in an N₂ atmosphere used to deduce CNT weight fractions.

Figure 4. Measured densities of NC and as calculated by rule of mixtures and T_g measured by DSC. Weight fractions are derived from TGA data, and volume fraction estimated as described in the text.

Assuming complete consolidation and no voids, expected NC densities were estimated using the rule of mixtures from volume fractions; porosity was calculated from the expected and measured densities. The rule of mixture densities generally compared well with those measured, following the expected linear trend (Figure 4), suggesting that complete consolidation was achieved, with the exception of the NCs containing 13.6 vol% CNTs,
which were estimated to contain 2 vol% voids. DIC micrographs of polished sections appeared smooth at low CNT loadings (Figure 5a) while at the highest loading (Figure 5b) the microstructure appeared textured, with features on the scale of 50 – 100 μm, akin to that of the larger particles from the powder particle size distribution. It seems that the 13.6 vol% CNT NC particles were not fully consolidated even at high pressures, presumably due to the high viscoelasticity of heavily CNT loaded polymers [4]. Some of the resin was pushed out of CNT rich regions locally to fill gaps between powder particles (Figure 5c), but the highly viscoelastic resin could not effectively migrate to fill areas with larger gaps, giving rise to localised voids.

The curing of all NCs follows the typical exotherm for epoxy curing reactions, including the well documented [18] acceleration effect with increasing CNT loading. No significant variability was observed in the final extents of reaction, suggesting that the degree of crosslinking was not affected by the presence of large CNT volumes, which was confirmed by the constant Tg (Figure 4) across all NCs.
3.2 Nanotube dispersion

The ability of a process to disperse and distribute CNTs in a matrix is critical in reducing stress concentrations associated with agglomerates and thus achieving the full mechanical potential of NCs, particularly for mechanically entangled, industrial grade CNTs. For untreated MWCNTs in epoxy resins, agglomerates are typically a few micrometres in diameter and spaced tens of micrometres apart [15]. CNT agglomerates were not observed (Figure 6) and the CNTs, for the most part, appeared individualised (Figure 7). The matrix cones suggest a strong adhesion between the CNTs and the resin most likely associated with plastic deformation of the matrix during CNT pullout (Figure 7). Evidence of a good interface between the NC constituents is promising, especially with regards to stress transfer.
Figure 6. Low (left column) and high (right column) magnification SEM micrographs of NCs with different CNT loadings, as indicated. Samples depicted in the first three rows were sputter coated, while the last one was uncoated.
3.3 Nanocomposite electrical and thermal properties

The electrical properties and behaviour of NCs have been extensively studied and reviewed [22, 23]; both electrical percolation and the maximum DC conductivity have been shown to be strongly affected by CNT purity [24], aspect ratio [25], processing parameters [26, 27] and orientation [28-30]. The relationship between a perfect dispersion of fillers in a matrix and the percolation threshold has been formalised using the excluded volume concept [31, 32], and provides a means to qualify the CNT dispersion state, by comparing the calculated percolation threshold using the CNT aspect ratio to that measured.

$\sigma_{\text{DC}}$ was measured along three axes, two perpendicular (X and Y) and one parallel (Z) to the consolidation direction (inset of Figure 8). A $\sigma_{\text{DC}}$ of $67 \pm 5 \text{ S m}^{-1}$ was measured in both X and Y axes at the highest CNT loading of 13.6 vol%. This value is about four times lower than the highest previously reported [15] for an epoxy based NC with 8 wt% randomly dispersed CNTs. It should be noted that the CNTs used in that work had larger aspect ratios (290 compared to 43 in this work) and processing was optimized by finding the shear rate which produces the NC with maximum $\sigma_{\text{DC}}$.

The $\sigma_{\text{DC}}$ (Figure 8) in the X/Y and Z axes begin to diverge beyond CNT loadings of ~6 vol%. The difference would be consistent with CNT orientation induced during compression.
moulding not fully relaxing before curing due to increasing viscoelasticity of the NC with increasing CNT loading. As a result, the CNTs would have been preferentially aligned in the plane perpendicular to the direction of the applied force and the NC would have become increasingly anisotropic. This phenomenon is important to consider for characterizing both the percolating characteristics of the network and the micromechanical behaviour of the NC.

![Conductivity vs vol% CNT](image)

**Figure 8.** Electrical conductivity of NCs measured in three axes.

\[
\sigma_{DC} \propto |\Phi - \Phi_C|^t
\]

\[
\Phi_C = 1 - \exp\left(-\frac{cV}{\langle V_e \rangle}\right)
\]

The datasets in Figure 8 were fitted with a power-law function [23] (Equation 1): where \(\Phi\) is the nanotube volume fraction, \(\Phi_C\) the nanotube volume fraction at the percolation threshold and \(t\) parameter indicating network dimensionality. The fit yielded \(\Phi_C = 1.85\pm0.70\) vol\% in all axes, \(t = 1.37\pm0.40\) for the X and Y axis data, and \(t = 1.01\pm0.28\) for the Z axis data; a value of \(t\) near 1.3 has been shown to correspond to anisotropic networks, with elements preferentially aligned in a plane [33]. Nonetheless, values of \(t\) should be taken with caution as the degree of anisotropy is likely to vary. The general expression for calculating the percolation threshold according to the excluded volume concept is given by Equation 2:
where $c$ is the total excluded volume of the system, $V$ the volume of the filler particle and $<V_e>$ the excluded volume of an individual filler particle. [34] Using the average measured post-processed CNT length (430 ± 250 nm), diameter (10±3 nm), and $c = 1.4$ and $2.8$ for an isotropic and anisotropic system of capped cylinders [35], respectively, the excluded volume approach predicts $\Phi_C = 1.2$ and 2.4 vol%. The calculated CNT percolation threshold values are within the range of those deduced from the fitted $\sigma_{DC}$, supporting the interpretation from SEM micrographs that the processing method was effective in dispersing the CNTs in the matrix, which form a statistical network at low loadings.

![Figure 9. Thermal conductivity of NCs measured in the Z-axis.](image)

The $k$ of the NCs was measured in the $Z$-axis, parallel to the consolidation direction. The addition of 13.6 vol% CNTs to the resin improved the $k$ by 165% from 0.29 to 0.77 Wm$^{-1}$K$^{-1}$ (Figure 9). The trend is similar to the previous highest $k$ reported [14] for shear mixed CNT-epoxy NCs (0.33 Wm$^{-1}$K$^{-1}$ at 5 wt% CNTs), but greater in absolute terms due to the higher CNT volume fraction. A non-linear least squares fit of the data using the effective medium approach (EMA [36]) yields an interface thermal resistance value of $1.9\times10^{-8} \pm 10^{-9}$ m$^2$KW$^{-1}$, which falls within the $10^{-8}$ m$^2$KW$^{-1}$ order of magnitude theoretically calculated and measured [37, 38] for CNT-polymer interfaces. Furthermore, the $k$ of the matrix used in this study (0.29
Wm⁻¹K⁻¹) was appreciably higher than that typical for cured epoxy resins (0.19 Wm⁻¹K⁻¹ [38]) for which the relative improvement upon addition of a large fraction of CNTs should be even more significant. The current maximum measured value (0.77 Wm⁻¹K⁻¹) approaches the requisite range (1-2 Wm⁻¹K⁻¹ [39]) for thermal interface materials (TIM) such as aluminium filled silicones used in heat dissipation applications.

3.4 Nanocomposite mechanical properties

In the majority of existing studies on NCs with randomly dispersed CNTs, Young’s modulus (E_NC) and strength (σ_NC) improved up to CNT loadings of a few volume percent, after which moduli plateaued [40] and strengths sharply dropped [6, 41] due to inadequate processing resulting in CNT agglomerates that do not transfer stress efficiently and act as stress concentrators. Furthermore, NCs with large CNT loadings (> 5 wt%) are severely embrittled, with failure strains reduced by 100-200% [6].

In this work, E_NC increased monotonically with increasing CNT volume fraction (Figure 10a) and the maximum σ_NC were achieved at the highest loading (Figure 10b). The 2 vol% of voids in the NC containing 13.6 vol% CNTs are not thought to influence either modulus or strength significantly. These voids (Figure 11) were significantly smaller (10-30 μm) than an estimate of the critical defect size (c) for the resin (~500 μm), approximated from the Irwin definition of the stress intensity factor (K) [42] (Equation 3); a K of 1.18±0.05 MPa m¹/² was measured from preliminary fracture tests on the neat resin and σ is the failure strength of the neat resin (57.7±8.6 MPa).
The maximum $E_{NC}$ value of 5.4 GPa is significant not only because it is the largest reported to date for a CNT-epoxy NC produced by shear mixing, but also because of the possibility to upgrade the performance of inexpensive resins. The reduction in failure strain (Figure 12a), despite the improved strength and stiffness is expected, and also frequently observed in thermoplastics with high CNT loadings [3]; the stiff reinforcing CNTs break or pull out at lower strain, transferring additional load to the matrix, leading to overall composite failure. Given their large surface area, and potential entanglement, CNTs may also directly restrict plastic flow of the surrounding resin-rich regions, particularly increasing CNT loadings. The improved trends and large $E_{NC}$ values can be attributed to the good CNT dispersion and distribution maintained throughout processing, which allow for more effective reinforcement and stress transfer using the relatively defective and short CNTs.
The measured $E_{NC}$ as a function of CNT volume fraction were fitted with Halpin-Tsai [13] equations (Equation 4) for randomly oriented fibres, as well as the rule of mixtures with correction factors for length ($\eta_L^E$) and orientation ($\eta_O^E$) (Equation 5) as proposed by Cox [43] and Krenchel [44], respectively. A CNT modulus ($E_{CNT}$) of 46±3 GPa was determined by fitting experimental data with the Halpin-Tsai equations, and 65±5 GPa using the rule of mixtures, assuming the CNT network was anisotropic ($\eta_v = 0.375$). These values compare well with those reported in the literature for highly defective, wavy, industrial grade MWCNTs [45-47].

$$\frac{E_{NC}}{E_m} = \frac{3}{8} \left( 1 + \frac{\zeta \eta_{Long}^v f}{1 - \eta_{Long}^v f} \right) + \frac{5}{8} \left( 1 + 2 \eta_{Trans}^v f \right)$$

where $\eta_{Long} = \frac{(E_{CNT}/E_m) - 1}{(E_{CNT}/E_m) + \zeta}$, $\eta_{Trans} = \frac{(E_{CNT}/E_m) - 1}{(E_{CNT}/E_m) + 2}$, $\zeta = 2 \frac{l}{d}$

$$E_{NC} = \eta_O^E \eta_L^E E_{CNT} v_f + E_m (1 - v_f)$$

$$\sigma_{NC} = \eta_O^E \eta_L^E \sigma_{CNT} v_f + \sigma_m (1 - v_f)$$

$$\tau_{IFSS} = \sigma_{CNT} \frac{r}{l_c}$$

The NC fracture strength values at all loadings (triangles in Figure 12a) were also fitted with a modified rule of mixtures (Equation 6), with the orientation factor ($\eta_O^\sigma$) also taken to be
0.375 and the length factor \( \eta_L^G \) calculated to be 0.186 as prescribed in [48], using the CNT length distribution and critical length, as measured by SEM. A lower bound for the critical length (1130±120 nm) was estimated by doubling the average of the ten longest CNTs protruding from fracture surfaces. From the slope of the linear fit of the NC fracture strengths with Equation 6, a CNT strength of 2.9±0.6 GPa was calculated, which agrees well with experimental results (3.6 GPa) for CVD grown CNTs [49]. The CNT-matrix interfacial shear strength (IFSS) was estimated from Equation 7, where \( r \) is the CNT radius and \( l_c \) the critical length. An IFSS of 12±9 MPa was estimated, using the lower bound for critical length estimated from the SEM fractography, which agrees well with experimental results (20-40 MPa) obtained from pull-out tests [51].

4. Conclusions
A novel, readily scalable, powder based route for the fabrication of epoxy based nanocomposites with high CNT loadings (20 wt%/13.6 vol%) is presented. The process allowed the successful production of a dense material with excellent CNT dispersion and distribution, providing significantly improved mechanical properties (\( E = 5.4 \) GPa, \( \sigma = 90 \) MPa, \( \epsilon = 2.5\% \)). Although the nanocomposites with the highest CNT loading contained some porosity (2 vol%), the modulus and strength continued to increase. The high pressure consolidation process created an anisotropic CNT network, resulting in a material with large electrical (67 Sm\(^{-1}\)) and thermal (0.77 Wm\(^{-1}\)K\(^{-1}\)) conductivities relative to previous reports. Its mechanical and transport properties make the nanocomposite a candidate for multifunctional applications such as EMI shielding (in theory, 50 dB at 5 GHz [52]) and thermal dissipation where mechanical performance is also critical.

Even though this processing route allows fabrication of NCs without CNT agglomerates, one of the intrinsic limitations to composite performance remains the quality of the filler. Further
efforts should consider longer, straighter, more graphitic and generally higher quality CNTs. Along with materials selection, the powder particle size also appears to be critical to successful consolidation and, therefore, performance of the material; the process could be improved by reducing and narrowing the distribution of powder particle sizes. In addition to its exciting intrinsic properties, the powdered nanocomposite is potentially an interesting feedstock for processes such as powder coating and powder prepregging [53, 54] where multifunctional performance is desired, particularly for carbon based hierarchical composites [55].

Acknowledgements
The authors would like to thank Dstl for funding the research and Hexcel for supplying materials.

References
