Molecular Alignment of a Meta-Aramid on Carbon Nanotubes by In-Situ Interfacial Polymerization.

Cécile A. C. Chazot,* Behzad Damirchi, Adri C. T. van Duin, A. John Hart*

Cécile A. C. Chazot - Department of Materials Science and Engineering, Department of Mechanical Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, Massachusetts 02139, United States.

Behzad Damirchi - Department of Mechanical Engineering, Pennsylvania State University, 245 Research East, University Park, Pennsylvania, 16802, United States. †

Adri C. T. Van Duin - Department of Mechanical Engineering, Pennsylvania State University, 240 Research East, University Park, Pennsylvania, 16802, United States.

A. John Hart – Department of Mechanical Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, Massachusetts 02139, United States.

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ABSTRACT
Molecularly organized nanocomposites of polymers and carbon nanotubes (CNTs) have great promise as high-performance materials; in particular, conformal deposition of polymers can control interfacial properties for mechanical load transfer, electrical or thermal transport, or electro/chemical transduction. However, controllability of polymer-CNT interaction remains a challenge with common processing methods that combine CNTs and polymers in melt or in solution, often leading to non-uniform polymer distribution and/or aggregation of CNTs. Here, we demonstrate CNTs within net-shape sheets can be controllably coated with a thin, conformal coating of meta-aramid by simultaneous capillary infiltration and interfacial polymerization. We determine that π interaction between the polymer and CNTs results in chain alignment parallel to the CNT outer wall. Subsequent nucleation and growth of the precipitated aramid forms a smooth continuous layered sheath around the CNTs. These findings motivate future investigation of mechanical and interfacial properties of the resulting CNT composites, and adaptation of the in-situ polymerization method to other substrates.

INTRODUCTION
Ordered polymer nanocomposites provide a route to high-performance lightweight structural materials with applications including aircraft and spacecraft structures, sporting equipment, and filtration membranes.\(^1\) Research and development on nanocomposite processing has been devoted to the combination of fillers such as high-crystallinity natural\(^2,3\) or synthetic\(^4,5\) polymer fibers or carbon nanotubes (CNTs)\(^6,7\) with polymers such as poly(vinyl alcohol)\(^3,8,9\) or commercial resin systems.\(^10\) The molecular organization and potential ordering of the polymer at the surface of the nanofillers has also been a particular center of attention as a mean to affect overall composite performance.\(^6\)

Due to their outstanding mechanical and electrical properties and low mass density, the composites of CNTs with engineered polymers offer great potential as next-generation lightweight structural materials.\(^11\) CNT-polymer composites have most commonly been made by resin infiltration of CNT mats or fibers, thereby translating manufacturing techniques widely used for carbon-fiber composites.\(^6\) Nevertheless, as the size of the reinforcement fibers reduces to the nanoscale, the complex interplay among transport-limited polymer infiltration, curing kinetics, and CNT-polymer molecular interaction greatly impacts the local polymer content and properties, and
Therefore, mechanical properties of the final composites. For example, the polymer chemistry and CNT surface chemistry can greatly influence polymer chain packing and organization, through localized chain alignment,\textsuperscript{12} wrapping around the CNTs\textsuperscript{13} or even long-range crystallization.\textsuperscript{7,9} The chemical interaction between the polymer and CNTs also affects wetting of the resin precursor on the CNTs, the resin infiltration speed,\textsuperscript{10,14} polymerization or curing reaction kinetics,\textsuperscript{15} and final composite mechanical properties through load transfer.\textsuperscript{16} However, the multi-component nature of the polymer precursors impedes the assessment of each compound’s contribution to the resin-CNT interaction. This intricacy constitutes a major challenge to experimentally understand and accurately compute the influence of CNTs on polymer formation. The study of such complex systems also prevents the extraction of guiding principles towards the design of polymer systems specifically for high-performance CNT-based composites. Moreover, resin cure cycles typically involve high temperature treatment for prolonged periods of times, from hours to days,\textsuperscript{17,18} making systematic studies of the CNT-polymer interaction particularly time- and energy-consuming.

Here we show that conformal polymer coatings on CNTs can be obtained by in-situ interfacial polymerization (ISIP) of meta-aramids,\textsuperscript{19,20} through short-range molecular ordering of the polymer chains and long-range nucleation and growth of polymer particles. ISIP of meta-aramid poly(m-phenylene isophthalamide) (PMPI) relies on the reaction of m-phenylene diamine (MPD) dissolved in water with isophthaloyl chloride (IPC) dissolved in an organic solvent (in this case, cyclohexanone) (Figure 1). A nanoporous aligned CNT sheet (Figure 1a), obtained commercially and produced by floating-catalyst chemical vapor deposition (CVD), is immersed in the aqueous and organic solutions subsequently (Figure 1b,c). Upon contact with the organic phase, the liquid wicks into the network by capillary action, displacing the aqueous phase (Figure 1c). MPD is transported across the moving liquid-liquid interface into the organic phase where it immediately reacts with IPC, producing PMPI chains and releasing hydrochloric acid as a by-product. As the PMPI chains grow in length, they precipitate out of the organic phase to form solid polymer onto the CNTs. Therefore, ISIP relies on the simultaneous liquid-liquid interface motion and polymer precipitation inside the nanoporous network of CNTs. Last, rinsing and solvent evaporation result in a CNT-PMPI composite free of impurities and unreacted monomers (Figure 1d). By adjusting monomer concentration, the mass of polymer produced as well as the morphology of the polymer coating onto the CNTs can be adjusted.\textsuperscript{20}
Figure 1. Schematic rendering of the ISIP process, forming a polymer-CNT composite sheet. (a) Neat aligned CNT mat. (b) CNT nanoporous network imbibed with water solution. (c) Immersion in the organic solution results in simultaneous liquid displacement and polymer formation on the organic side of the interface. (d) After rinsing and drying a CNT-aramid composite mat is obtained.
In recent work, we showed that PMPI forms in droplets or conformal sheaths around the CNTs, depending on the relative rates of transport and reaction kinetics.\textsuperscript{20} The polymer formed is amorphous in nature, demonstrating that commonly observed crystallization of meta-aramids through interchain hydrogen interaction\textsuperscript{21} does not contribute to ordering of PMPI in the vicinity of CNTs. Instead, we expect from literature that the prominent $\pi$-interaction\textsuperscript{22} involving PMPI meta-phenyl rings and the outer wall of CNTs induces short-range molecular ordering of polymer chains onto the CNT. This sub-nanometer ordered polymer layer is hypothesized to serve as a template for further precipitation by heterogeneous nucleation of amorphous PMPI particles onto the CNTs, governing long-range polymer sheath formation.

RESULTS AND DISCUSSION

First, we aim to experimentally identify the governing phenomenon for the formation of conformal polymer sheaths. Precipitation kinetics and its dependence on monomer concentration in the ISIP process is assessed by scanning electron microscopy (SEM) observation of the polymer morphology in presence of CNTs (Figure 2). At low monomer concentrations (0.098 mol.L\textsuperscript{-1}), PMPI precipitates onto the CNT surface as nanometer-scale droplets and thin polymer sheaths (Figure 2a,b). For experiments performed at higher concentrations, from 0.098 mol.L\textsuperscript{-1} to 0.248 mol.L\textsuperscript{-1}, the polymer sheath diameter increases from $\sim$20 nm to $\sim$40 nm (Figure 2b,c). Additionally, while some CNTs are observed to have smooth conformal sheaths, others feature bead-like deposits adhered to an underlying smooth layer (Figure a-c). Last, at high monomer concentration (greater than 0.2 mol.L\textsuperscript{-1}), and in regions where the CNT density is low in the network, smooth conformal polymer coatings with thicknesses in the order of 100 nm with a layered structure can be observed around individual CNTs (Figure 2d). These observations are strong evidence that PMPI forms by heterogeneous nucleation and growth of polymer particles at the surface of the CNTs or on the already-formed polymer sheaths (Figure 2e). Initial polymer nuclei form around individual CNTs and, as more polymer is formed, these coalesce as a continuous sheath which serves as the surface for subsequent precipitation. The layered structure of the larger sheaths observed at high concentration is consistent with this interpretation.
Figure 2. PMPI precipitation mechanism onto CNTs in ISIP. (a-d) SEM images of CNT-PMPI composites obtained for monomer concentrations of (a-b) 0.098 mol.L\(^{-1}\), (c) 0.248 mol.L\(^{-1}\) and (d) 0.281 mol.L\(^{-1}\). (e) Schematic describing the steps of nucleation and growth of PMPI onto CNTs and forming polymer sheaths. Nuclei form on a CNT before coalescing to form a smooth polymer sheath which subsequently supports polymer nuclei formation. Step 2 is repeated (s.x where x≥2) to form multiple polymer layers around the CNT. Specific steps of the polymer nucleation and growth process are also indicated on (a-d) where k, l and m are integers and k<l<<m.

The CNT-polymer chemical interaction and its influence on polymer ordering within the conformal sheaths is assessed by molecular dynamics (MD) simulations, as described in Supplementary Information section S1 (Figure 3). The reaction between MPD and IPC is assumed to occur in a thin reaction zone on the organic side of the cyclohexanone-water interface (Figure 3a).\(^{19}\) To assess π-π stacking\(^{22}\) between the surface of the CNTs and the aromatic rings of the PMPI backbone, we define the normal vector to each meta phenyl group of the polymer as the cross product of the two vectors generated using three nonadjacent aromatic carbon atoms (Figure 3b). The angle \(\phi\) between this normal vector and the normal to the CNT surface is used to calculate the Herman’s orientation function \(f\)\(^{23}\) which quantifies the alignment of the polymer aromatic rings in relation to the CNT surface:
\[ f = \frac{3((\cos^2 \varphi) - 1)}{2} \]

Here the angle brackets stand for the average value over all angles, and

\[ \langle \cos^2 \varphi \rangle = \frac{\int_{0}^{\pi} \cos^2 \varphi \sin \varphi \, d\varphi}{\int_{0}^{\pi} \sin \varphi \, d\varphi} \]

A value of \( f = 1 \) describes parallel \( \pi-\pi \) stacking of all the aromatic rings with the CNT surface in a face-to-face or parallel staggered configuration,\(^{22}\) while a value of \( f = 0 \) corresponds to random alignment of the polymer chains along the CNT surface. If \( f = -0.5 \), the vectors are perpendicular and the interaction of polymer chains with the CNTs can be described as \( \pi-\pi \) T-shaped or \( \pi-\pi \) Y-shaped stacking (also called edge-to-face orientation).\(^{22}\)

We use a ReaxFF reactive force field framework\(^{24}\) with chemical acceleration tools to enable reaction barrier sampling at time-scales accessible to ReaxFF-based molecular dynamics\(^{25-29}\). With this, we capture the polymerization reaction between MPD and IPC in the organic phase as well as the CNT-PMPI interaction. The simulation features three successive stages, shown in Figure 3c. The influence of monomer concentration on polymer-CNT interaction is studied by varying the number of monomer molecules (50 or 250 of each monomer) present in the system at \( t=0 \) while keeping the initial number of cyclohexanone molecules constant and equal to 500 (Table S1). During stage 1 (0-500ps), monomers are allowed to react with one another in presence of cyclohexanone molecules. Stage 2 and stage 3 (both 500ps in duration) correspond to removal of the organic solvent; at the end of stage 1, 250 cyclohexanone molecules are removed, and 250 more are removed between stage 2 and stage 3 (Table S1).

We monitor the value of the Herman’s orientation function \( f \) and the normal vector to the polymer chain aromatic groups in stage 3 (Figure 3d). At low concentration (50 of each monomer at \( t=0 \) in the MD simulation), polymer chains form an ordered layer along the CNT surface (Figure 3d). With a \( f \) value between 0.7 and 0.8, the polymer chains’ aromatic groups interact with the CNT outer wall via parallel \( \pi-\pi \) stacking, resulting in parallel alignment of the polymer chains with the CNT surface (Figure 3e). With increasing monomer concentration (250 of each monomer at \( t=0 \)), \( f \) is only 0.2-0.3, and the PMPI chains are randomly aligned in the absence of \( \pi-\pi \) stacking away from the CNTs. This low \( f \) value is consistent with the experimental observation that PMPI
synthesized by ISIP is amorphous and show no visible long-range order. Nevertheless, the spatial distribution of the normal vector to the aromatic rings of the polymer backbone (Figure 3f) indicates that chains located in the vicinity of the CNT still align along the surface of the nanotube via parallel π-π stacking, while chains closer to the edge of the box form randomly aligned bridge between CNTs, as captured by periodic boundary conditions in the simulation. It is worth noting that experimental observation of sheath formation was performed on a mixture of cylindrical and self-collapsed (i.e., flattened) CNTs. Indeed, large-diameter single- and double-walled CNTs produced by CVD show collapsed geometry due their radial flexibility or defect density. Importantly, the MD simulation predicts the same alignment trend for collapsed CNTs (Figure S1).
solvent molecules begin removed (Stage 2), and then all the solvent molecules being removed (Stage 3). The color scheme is that of (a) and (b) and orange color represent cyclohexanone. (d) Herman’s orientation function during stage 3 for a simulation featuring 50 and 250 of each monomer. Both simulations feature 500 cyclohexanone molecules. (e-f) Polymer morphology and spatial distribution of the normal vector to the PMPI aromatic rings at t=1500ps for an initial number of each monomer molecules of (e) 50 and (f) 250.

Now, we experimentally evaluate the nature of the CNT-polymer molecular interaction and its evolution as PMPI forms increasing number of sheath layers around the CNTs. We combine Fourier transform infrared (FT-IR) and Raman spectroscopy, which together let us identify the vibrational modes affected by the CNT-polymer interaction.34,35

FT-IR spectroscopy is used first as PMPI exhibit a strong FT-IR signal while CNTs feature vibrational modes which are only Raman-active for the most part (Figure 4).35 This results in the straightforward isolation of the contribution from PMPI to the FT-IR spectrum, and its comparison against the neat polymer and neat CNT signals (Figure 4a). The detailed peak assignment for the FT-IR signature of PMPI can be found in Table S2.34,36 To analyze the evolution of PMPI-CNT π-interaction with increasing polymer content, we observe the location of selected peaks characteristic of in-plane (Figure 4b,c) and out-of-plane (Figure 4d) vibrational modes relevant to the meta-aromatic groups and the secondary amines of PMPI. For CNT-PMPI composites, the location of peaks representing PMPI meta-phenyl ring semicircle stretch mode (Figure 4b) and in-plane C-H bend-stretching mode (Figure 4c) noticeably shift towards higher wavenumber with increasing monomer concentrations. For composites obtained with monomer concentrations greater 0.2 mol.L⁻¹, the peak location stabilizes at a value slightly smaller than that found on the neat PMPI FT-IR spectrum. The blue shift and asymptotic behavior of the in-plane aromatic vibrational peaks suggest that in-plane motion of the PMPI meta-phenyl ring is hindered when the polymer forms in the vicinity of the CNT outer wall, and that this hindrance decays as more polymer is formed, further from the CNT surface. Additionally, splitting of the peaks relevant to the two vibrations involved in semicircle stretch is observed for PMPI formed in presence of CNTs. The observed peak shift and splitting are consistent with a strong π-π interaction35,37,38 that weakens with increasing polymer content as predicted by the MD simulation. Indeed, the strong π interaction between the CNTs and polymer repeat units results in an attractive electrostatic force that disrupts the degrees of freedom involved in PMPI’s characteristic vibrational modes. In the
case of parallel $\pi$-$\pi$ stacking for instance, the electrostatic quadrupole interactions between $\pi$ orbitals of the CNT outer wall and PMPI aromatic groups results is disruption of the motion of the atoms involved in the in-plane vibrational modes, therefore affecting the frequency and relative intensity of the characteristic peaks.

Additional evidence of $\pi$ interaction can be found in the FT-IR peaks relevant to out-of-plane vibrational modes (Figure 4d). First, red-shift of the Amide V band provides evidence of interaction between the CNTs and the electron-rich -NH- group. This interaction appears to decrease rapidly with increasing thickness of the polymer sheath, as the peak position reaches a steady value of 721 cm$^{-1}$ for monomer concentrations greater than 0.1 mol.L$^{-1}$, a value 5 cm$^{-1}$ greater than that of neat PMPI. While the Amide V band shows this red-shift, in-plane vibrational modes related to the polymer amide linkage, such as Amide I and II vibrations corresponding to stretching of C=O and C-N bonds respectively, show no significant frequency shift. This can be explained by an additional NH-$\pi$ interaction$^{22,35,38}$ between PMPI and CNTs. The red-shift of the Amide V band occurs in a narrower range of concentrations compared to the aromatic parallel vibrational modes. This reflects the weaker interaction energy involved in NH-$\pi$ compared to $\pi$-$\pi$ stacking.$^{38}$ Moreover, the FT-IR peak corresponding to the meta-phenyl group’s sextant out-of-plane bending mode shows a concentration-independent blue shift of 10 cm$^{-1}$ compared to neat PMPI. The combination of concentration-dependent peak location for in-plane aromatic vibrational modes with concentration-independent significant peak shift for out-of-plane aromatic vibrational modes hints at parallel $\pi$-$\pi$ stacking between CNTs and PMPI. In edge-to-face $\pi$ interaction, one positively charged structure of each PMPI ring is juxtaposed against the $\pi$ electron cloud of the CNT ring,$^{6,22}$ leaving most of the other bonds free to vibrate, particularly through in-plane motions. On the other hand, in parallel $\pi$-$\pi$ stacking, multiple aromatic sub-groups in each PMPI ring feature electrostatic interaction with the $\pi$ electron cloud of the CNT outer wall. Parallel $\pi$-$\pi$ stacking results in greater restraint of both the in-plane and out-of-plane aromatic bonds’ vibration, therefore altering the shape of the relevant FT-IR peaks.
Figure 4. FTIR investigation of CNT-PMPI interaction and its dependence on ISIP monomer concentration. (a) Raw FTIR spectra of neat CNT sheets, neat PMPI, and CNT-PMPI composites obtained at various monomer concentration \([M_0]\). Grey boxes show wavenumbers which are represented on b, c and d. (b-c) Zoomed view in spectral regions exhibiting peaks characteristic of in-plane vibrational modes of the meta-phenyl group of PMPI. (b) Analysis of the absorbance spectra in the 1450-1500 cm\(^{-1}\) range and location shift with monomer concentration of the peak characteristic of PMPI meta-phenyl ring semicircle stretch modes. The grey dashed horizontal line is placed at the location of the same vibrational peak on the neat PMPI spectrum. The C and S letters are used to identify which part of the aromatic group is contracted (C) and stretched (S). (c) Analysis of the absorbance spectra in the 1360-1450 cm\(^{-1}\) range and location shift with monomer concentration of the peak characteristic of PMPI meta-phenyl ring in-plane C-H bend-stretching mode. (d) Analysis of the absorbance spectra in the 650-750 cm\(^{-1}\) range and evolution with monomer concentration of out-of-plane bending vibrational mode related to the amide linkage (Amide V band) and the meta-phenyl group of PMPI. The amide linkage peak at ~720 cm\(^{-1}\).
for CNT-PMPI composites and 716 cm$^{-1}$ is representative of the N-H out-of-plane bending vibration while the aromatic peak at $\sim$693 cm$^{-1}$ for the composites and 682 cm$^{-1}$ for neat PMPI is characteristic of sextant out-of-plane bending mode of the meta-phenyl group.

To further confirm that CNT-PMPI interaction involves parallel $\pi$-$\pi$ stacking, we investigate the polymer’s Raman-active vibrational modes in CNT-PMPI composites and compare them to that of neat PMPI. To isolate PMPI contribution from the strong CNT signal in composites, we implemented a custom principal component analysis (PCA) algorithm which enables the isolation of the partial Raman spectra (PRS, also called component spectra) involved in the overall measured signal (Figure 5).\textsuperscript{39} PCA relies on the acquisition of multiple spectra of the analyzed material (Figure 5a) which are then processed to reveal the PRS. The sum of the PRS describes the overall Raman spectrum of the characterized material. The neat CNT sheet signal is found to be accurately described with 5 components, while the CNT-PMPI composite Raman response was found to feature 6 components. Because of their characteristic peaks corresponding to the D, G and G’ bands,\textsuperscript{35} the first 5 PRS of both the neat sheet and PMPI-CNT composite are attributed to the CNTs themselves (Figure 5b). Comparing the first 5 PRS of the CNT-PMPI composite with that of the neat CNT sheet, no significant difference is observed, and the modest deviation for components 3 and 4 can be attributed to variation of the starting CNT material as well as the difference in number of spectra acquired between the two acquisitions. The sixth component obtained for the CNT-PMPI composite is attributed to the contribution of the polymer and compared to that of neat PMPI (Figure 5c).\textsuperscript{34,40} The detailed peak assignment for the Raman signature of PMPI can be found in Table S3. The peak associated with the meta-phenyl ring trigonal ring breathing vibration (located at 1001 cm$^{-1}$) which is also the highest intensity mode for neat PMPI completely disappear from the sixth PRS of the CNT-PMPI composite. This confirms that the polymer interacts with the CNT outer wall through parallel $\pi$-$\pi$ stacking which impinges the in-plane ring breathing vibration through electrostatic interaction. In parallel $\pi$-$\pi$ stacking, ring breathing of the meta-phenyl group is opposed by the attractive force between the positively charged groups of one aromatic group with the negatively charged $\pi$-electron-clouds of a graphene ring. This results in the annihilation of the associated vibrational peak from the PRS attributed to PMPI.
Figure 5. Raman spectroscopy investigation of CNT-PMPI interaction. (a) Representative spectra taken at different locations on the cross-section of a CNT-PMPI composite made by ISIP using a monomer concentration of 0.150 mol.L⁻¹. (b) First five component spectra are attributed to the CNTs in the composites. Solid lines correspond to components 1-5 extracted on the CNT-PMPI composite while dashed lines correspond to the five components picked up on the Raman response of a neat CNT sheet. (c) Sixth component spectrum obtained on the CNT-PMPI composite and comparison to the Raman spectrum of neat PMPI synthesized by interfacial-polymerization. Inset schematic represents the trigonal ring breathing mode associated with the 1000 cm⁻¹ peak.

Most prior observations of polymer nanostructure formation in the presence of CNTs such as “shish-kebab”⁴¹,⁴² or smooth sheaths⁴³ rely on polymer crystallization through heterogeneous nucleation, as this has been observed for polyethylene,⁴¹ or poly(vinyl alcohol).⁴³,⁴⁴ Sheath formation and short-range ordering without increased crystallinity has been observed with CNTs for three other polymer systems, in addition to PMPI studied here: polycarbonate (PC),⁴⁵ polyvinylidene fluoride (PVDF)⁴⁶ and polypyrrole (PPy).⁴⁷ All these polymers feature a flexible
backbone and functional groups which can interact with the CNTs’ outer wall through π-interactions (Figure S2).\(^{38,48}\) PC sheaths exhibited a multilayer structure with a strong short-range adhesion of the polymer chains onto the CNTs’ surface, similarly to our observation for PMPI.\(^{45}\) The thickness of the polymer sheaths was only controlled in the case of PPy which was polymerized by chemical oxidation in presence of CNTs.\(^{47}\) Both PPy polymerization and ISIP of PMPI enabled the polymer coating thickness to be varied in the 10-100 nm range by adjusting monomer concentration. However, ISIP allowed for much shorter processing times (e.g., ISIP reaction time in the order of minutes, compared to hours for chemical oxidation polymerization of PPy). Moreover, chemical oxidative polymerization limits greatly the range of available polymer systems to often electrically conductive and thermally unstable polymers.\(^{49}\) In contrast, ISIP offers a rapid and flexible transport-reaction scheme for the formation of CNT-based composites with chain-oriented amorphous PMPI or other suitable polycondensation polymers such as polyamides, polyesters, or polyureas.\(^{19}\) The high thermal stability of amorphous PMPI (up to about 400°C), its superior mechanical properties, as well as its short-range ordering at the surface of the CNTs makes it particularly suitable for a broad range of high-temperature applications such as structural aerospace components, filtration membranes or high thermal conductivity materials for heat exchangers or power electronics.\(^{50}\)

**CONCLUSION**

In conclusion, we demonstrated that the conformal amorphous polymer sheaths observed in CNT-meta-aramid composites produced by ISIP is the result of the interplay between polymer precipitation kinetics and parallel π-π stacking that involves the aromatic groups from the polymer backbone and the CNTs outer wall graphene rings. Using ReaxFF reactive force field MD simulation, we predicted that while polymer chains at the immediate vicinity of a CNT feature strong alignment and parallel π-π stacking with the CNT outer wall, PMPI formed further from the nanotube is amorphous in nature. Subsequently, the formation of a smooth polymer coating around individual CNTs is shown to be a consequence of PMPI precipitation kinetics by successive heterogeneous nucleation and multilayer sheath formation. Last, we combined FT-IR and Raman spectroscopy to experimentally demonstrate the existence of both parallel π-π stacking and NH-π interaction between CNTs and PMPI, as well as their decrease with increasing number of conformal layers in the meta-aramid sheath. This new understanding of CNT-polymer interaction
combined with its implementation in the rapid and flexible ISIP synthesis scheme could guide the
design and selection of polymer systems for the development and manufacturing of high-
performance CNT-based composites.

ASSOCIATED CONTENT

Supporting Information.
The following files are available free of charge.
MD simulation framework, Materials and synthesis, Characterization methods, Spectroscopy and
FTIR/Raman peak assignment, Comparison of PMPI obtained by ISIP with other polymer
systems. (PDF)

AUTHOR INFORMATION

Corresponding Authors
*Cécile A. C. Chazot - Department of Materials Science and Engineering, Department of
Mechanical Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue,
Cambridge, Massachusetts 02139, United States. Email: cchazot@mit.edu
*A. John Hart – Mechanical Engineering Department, Massachusetts Institute of Technology, 77
Massachusetts Avenue, Cambridge, Massachusetts 02139, United States. Email: ajhart@mit.edu

Present Address
† Behzad Damirchi - Department of Computational and Quantitative Medicine, Biomedical
Research Center, City of Hope, 1218 Fifth Ave, Monrovia, CA 91016, United States

Author Contributions
C.A.C.C. and A.J.H. conceived the in-situ interfacial polymerization method. B.D. and A.C.T.D
conceived the MD simulation approach. B.D. conducted the MD study. C.A.C.C designed and
conducted the experiments and material characterization. C.A.C.C. wrote the first draft of the
manuscript. All authors contributed to the discussion of the results and the preparation of the
final version of the manuscript. C.A.C.C. coordinated the collaboration, and A.J.H. and A.C.T.D.
supervised the work.
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**Notes**

The authors declare no competing financial interest.

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**ABBREVIATIONS**

ISIP, in-situ interfacial polymerization; PMPI, poly(m-phenylene isophthalamide); CNT, carbon nanotube; MD, molecular dynamic; PCFF-IFF, polymer consistent force field–interface force field; CVD, chemical vapor deposition; MPD, m-phenylene diamine; IPC, isophthaloyl chloride; SEM, Scanning Electron Microscopy; FT-IR, Fourier-transform infrared spectroscopy; PCA, principal component analysis; PRS, partial Raman spectrum/spectra.

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