Mechanical and electrical properties of MWCNTs - high early strength cement - mortars composite: Dispersion of CNTs and effect of chemical admixtures

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Abstract: The objectives of this research were to study (1) the dispersion of MWCNTs in an aqueous system by three commercial admixtures (CAds) for concrete, and (2) the effect of CAds and MWCNTs on indirect tensile strength and electrical conductivity of MWCNTs-high early strength (HE) cement-mortar composites. To achieve the objectives, we dispersed MWCNTs in an aqueous system with (1) hydroxylated polymers-based water reducing plasticizer (HPs), a nonionic compound, (2) Naphthalene based superplasticizer (SNF), an anionic compound, and (3) calcium chloride-based accelerating agent (CC) a neutral amphoteric salt. We prepared a total of 242 samples grouped in three sets: (1) Plain mortar [PM] (water + HE cement + Sand), (2) [PM+CAd], and (3) [PM+CAd+MWCNTs]. The three CAds dispersed MWCNTs in an aqueous solution. The CC and HPs admixtures have a two-time bigger dispersing power than the SNF. They demand half of SNF's ultrasound energy for optimal dispersion. Although the SNF (anionic) based superplasticizer resulted incompatible with the HE cement, it improved the indirect tensile strength of [PM+SNF+MWCNTs] composite. In contrast, the CC (amphoteric) based accelerating agent was compatible with the HE cement; the CC adsorption on the MWCNTs surface favors an improvement in the electrical conductivity of [PM+CAd+MWCNTs] composite.

Key words: MWCNT, cement-based materials, high early strength cement, chemical admixtures, indirect tensile strength.

INTRODUCTION

Carbon nanotubes (CNTs) exhibit extraordinary properties suitable to advance new materials for many engineering applications. CNTs are molecules of carbon atoms in a periodic hexagonal structure, constituting a folded graphene sheet that forms a hollow cylinder (Segawa et al. 2016). A single hollow cylinder of carbon atoms forms a single wall CNT (SWCNT). In contrast, hollow concentric cylinders form a multiwall CNT (MWCNT). Thirty years have passed since the CNTs discover (Iijima 1991). Nowadays, there is a considerable amount of scientific reports regarding CNT applications in the construction industry (Bautista-Gutierrez et al. 2019, Evangelista et al. 2019, Rashad 2017). Cement-based materials (CBMs) doped with CNTs display new and enhanced mechanical properties, self-control of crack propagations, and high electrical conductivity, among other improvements (Sanchez & Sobolev 2010). However, several unsolved issues still limiting the application of CNTs in the construction industry. The CNTs' price/Improvement ratio...
still high enough to limit practical applications. Therefore, new advances are needed lowering the CNTs’ price and increasing the magnitude of improvement (MOI). Although more research is needed to reduce CNTs’ prices, the chemical vapor deposition method has shown the most promise in its price/unit ratio (Siddique & Mehta 2014). The increase of the MOI still been a challenge, mainly because there are several types of portland cement and three interactions to be engineered; (1) the interaction between the dispersing agent’s molecule and the cement particles, (2) between cement particle and CNTs, and (3) between dispersing agent’s molecule and CNTs. When the dispersing agent is a commercial CAd for concrete, the interaction between the dispersing agent’s molecule and the cement particles is “known,” but few understood (Kurdowski 2014). It is the base of a huge industry directed to reduce the cost and increase the functionality of concrete construction. However, more studies are needed, considering the compatibility of the CAd and the type of cement.

The interaction of CNTs-cement components involves studying CNTs’ effect on cement components hydration (Alafogianni et al. 2019, Makar & Chan 2009). While the interaction between the dispersing agent’s molecule and CNTs has been a hot research topic during the last two decades. Few studies are regarding the use of CAd for concrete as CNTs dispersing agents. Fewer studies are dealing with the consequences of the interaction between the CAd molecules and the CNTs. Notably, the engineering of the interaction between CAd molecules and the CNTs remains an issue to overcome.

A central challenge overcoming this issue is the homogeneous dispersion of the CNTs in the CBM matrix. A well-accepted and practical method includes the use of ultrasound to disentangle CNTs bundles in an aqueous solution and the use of a dispersing agent to disperse CNTs (Konsta-Gdoutos et al. 2010, Yu et al. 2007), and the optical absorption spectroscopy to control CNTs dispersion degree (Alafogianni et al. 2016). Previous studies have shown that if the ultrasound source’s mechanical energy is very high, the physical integrity of the CNTs is affected. Thus, those properties that depend on aspect ratio will be affected. The electrical conductivity is affected by an increase in the number of CNTs needed to reach the percolation threshold. While if the energy delivered is too low, the degree of CNTs disentanglement is low.

Similarly, the CNT clusters become defects that adversely affect the properties of the CBM. The minimum energy required for CNTs disentanglement in an aqueous solution depends on the amount of CNTs, the type and the quantity of the dispersing agent, and the dispersing agent’s dispersion power. Many studies in the literature reported on the minimum energy required to disentangle CNTs in an aqueous solution with a dispersing agent dissolved in it. However, few studies reported the energy required to separate CNTs in aqueous solutions with CAd. The dispersing agent must be adsorbed on the surface of the CNTs to disperse CNTs homogeneously in solution. However, the dispersing agent’s molecular structure, molecular size, and the electrostatic characteristic of their functional groups largely determine the interaction between CNTs and the dispersing agent’s molecule and the improvement in the properties of the CBM. Despite many studies reports on the dispersion power of a large number of dispersing agents (Rashad 2017), few studies reported on CNTs dispersion power of CAd for concrete.

Previous researchers have shown evidence of CNTs’ ability to improve the mechanical properties of CBM. Evangelista et al. (2019) found
that the addition of 0.4 wt% of CNTs to CBM increases compression, tensile, flexural strength by 40%, 30%, and 15%, respectively. The addition of low amounts of CNTs enhances fracture behavior by bridging micro-cracks and delaying their propagation; increases fracture strength, toughness, and the flexural toughness index (Alafogianni et al. 2019). Hawreen & Bojas (2019) reported that the addition of CNTs caused a reduction in the initial and long-term shrinkage of CBM up to 54 % and 15 %, respectively. The authors reported that the CNTs-CBM composite shows a 17%-18% lower long-term creep than the plain CBM. Sedaghatdoost & Behfarnia (2018) reported that 0.1 wt % MWCNTs incorporation in CBM improved compression, tensile, and flexural strength by 35 %, 8 %, and 11.2 %, respectively. Lushnikova & Zaoui (2017) used molecular dynamics simulation to study the effect of CNTs incorporation into CBM. The authors reported that the shear modulus, bulk modulus, elastic constants, and Poisson ratio improved upon CNTs incorporation.

Campillo et al. (2003) reported that CNTs improved cement samples’ compressive strength for the first time. Makar et al. (2005) observed MWCNTs crack bridging and fiber pull-out and concluded that it is due to strong bonding between the CNTs and the cement matrix. Li et al. (2005) found that cement hydration products coated CNTs in composite materials while exhibited good compressive and flexural strength. They also found that MWCNTs improved the pore structure of the cement matrix. Sáez de Ibarra et al. (2006) reported that MWCNTs+CBM composites exhibited better mechanical properties than displayed by the SWCNTs+CBM composites. Kowald et al. (2008) compared untreated MWCNTs-, and oxidized MWCNTs-cements composites and concluded that untreated MWCNTs improve low hardened C-S-H content while oxidized MWCNTs increases contents of high hardened C-S-H. Hunashyal et al. (2011) found that a low amount of MWCNTs (between 0.25 and 0.75 wt %) increased the flexural strength by 30% to 45%, and increasing the MWCNTs content to 1 wt% decreased the flexural strength. Wang et al. (2013) found that treated CNTs improve the flexural strength and Portland cement composite’s compressive strength. They reported that MWCNTs acted as bridges and made networks across crack and voids and suggested a strong bonding between the CNTs and the cement matrix. Wang et al. (2015) stated as an existing problem, the issue of compatibility among CNTs and CBM as well as CNTs and dispersants used.

However, few reports are regarding the compatibility between cement and CAd, cement and CNTs, and between the CAd and CNTs. Fewer reports are considering the indirect tensile strength of CNTs-CBM composites. A considerable amount of reports deals only with ordinary Portland cement. They do not report on the compatibility issue between the type of cement and CAds. Moreover, few reports are on CAd-CNT interaction, which governs the relative improvement in the composite’s mechanical properties. The mechanical properties of CNTs+CBM composites depend mainly on the bond’s strength between the CAd and CNTs. The CAd-cement particle interaction depends on the compatibility of the CAd with the cement. It may depend on the interference of the CAd with the hydration of the different phases of the cement.

Electrical conductivity represents a material’s ability to conduct electric current to a given temperature. The electrical conductivity of CBM is a fundamental property in applications like electrical grounding, lightning protection, resistance heating, static charge dissipation, electromagnetic interference shielding, cathodic protection, and thermoelectric energy generation (Chung...
2004). The electrical conductivity of CNTs-CBM composite sparks an interest in engineering materials by CNTs addition in “smart” sensing applications. Previous reports indicate that the conductivity of CBM improved by adding CNTs (Yoo et al. 2019). There are many reports on the electrical conductivity of CNTs-CBM composites (Demarchi & Tagliaferro 2015, Rashad 2017). The most effective way to achieve high electrical conductivity values in CNTs-CBM composites is by increasing the load of CNTs. However, while an increase of the CNTs concentration improves the electrical conductivity, it simultaneously affects the thermal and mechanical properties of the composite (Jagdale et al. 2015). Therefore, there is a need for more research to increase electrical conductivity without significantly affecting other properties. There are few reports on how commercial CAds for concrete affects the electrical conductivity of CNTs-CBM composites. Advances in engineering CNTs-CBM composites, particularly in “smart” sensing technology, depend not only on CAd’s work but also in uncovering the functioning and the contribution of the main actors’ electrical conductivity (CAds, CNTs, cement, aggregates).

Interestingly, in CBM could exist different electrical conductivity regimens, electronic and ionic as well. CNTs could work as an electronic highway for electrons and as an ionic highway for ions trapped inside the CNTs. The ionic conduction by CNTs is a research topic in Lithium batteries, where electrodes made of CNTs are promising materials (Han et al. 2020). The first one is a central property in the “smart” sensing application of CNTs-CBM composites. However, the CNT’s surrounding environment and the ability to change the conductivity under stress/strain are important (Jagdale et al. 2015).

This research aimed to study: (1) the dispersion of MWCNTs by three commercial CAds for concrete in an aqueous system, and (2) the effect of CAds and MWCNTs on indirect tensile strength and electrical conductivity of MWCNTs-HE cement-mortar composites. MWCNTs were dispersed in an aqueous solution to study the CAd and CAd+MWCNTs’ effect on the [PM+CAd+MWCNTs] composite’s mechanical and electrical properties.

MATERIALS AND METHODS

Figure 1 shows an illustrative flowchart depicting the essential stages in this study: (1) Optimization of parameters for homogeneous dispersion of CNTs in aqueous solution. (2) Preparation of a HE cement-mortar sample. (3) Mechanical and electrical conductivity properties characterization. The first stages involve the optimization of sonication parameters and the weight of CAd for homogeneous dispersion of CNTs in an aqueous solution. The second stages comprise the preparation of three sets of disc-shaped samples and three sets of rectangular prism-shaped samples: (1) Plain mortar [PM]. (2) Plain mortar with a CAd [PM+CAd]. (3) Plain mortar with a CAd and MWCNTs added [PM+CAd+MWCNTs]. The third stages involve measuring the indirect tensile strength (Brazilian test) and the electrical conductivity through the direct-current four-pole method.

Materials

A hydraulic cement type HE (high early-strength cement) by Holcim Ecuador S.A. (Latacunga, Ecuador) was used to prepare MWCNTs-HE cement-mortar composite. Table I shows the chemical and mineral composition and the specific surface area of the HE cement.

Timesnano, a subsidiary of Chengdu Organic Chemical Co. Ltd., Chinese Academy of Sciences, PR China, provided the MWCNTs used in this study. Table II shows the properties of MWCNTs,
and Figure 2 shows the morphological attributes of MWCNTs.

ADITEC Ecuadorian Cia. Ltda (Durán, Guayas, Ecuador) provided the three CAds used in this study. CAd stabilized the MWCNTs dispersion and prevented the re-aggregation of the nanotubes in water (Table III).

**Mixture proportions**

Table IV shows the raw material proportions in the plain mortar mixture preparation. We prepared a PM and MWCNT-mortar composite following the Ecuadorian standards NTE-INEM 155:2009 (NTE-INEM 2009) and NTE-INEM 2563:2011(NTE-INEM 2011). Table IV shows the main aggregate characteristics, design specification, and the amount of raw material needed to fabricate one cubic meter of mortar.

This study used disc-shaped samples with a thickness of 25 mm for the indirect tensile test (Brazilian test) gotten after cutting cylindrical specimens (50 mm, 100 mm) prepared with fresh mortar. Similarly, the electrical conductivity test carried out on rectangular prism-shaped samples (80 mm x 20 mm x 20 mm) was made directly with fresh mortar in molds with the
same dimensions. After 24 h, the specimens were removed from their mold and stored in a pool of water kept at 20 °C until the test date.

Table V shows the sets of samples prepared with three different CAds, three different CAd/cement loading ratios, and three different CNTs/cement ratios. Therefore, this study involved testing 185 discs and 57 rectangular prisms.

**Dispersion of MWCNTs in aqueous solution**

Admixtures and ultrasound allowed the dispersion of MWCNTs in an aqueous solution. The CAd dispersibility was optimized through the CAd/MWCNTs loading ratio’s optimal value and delivering the minimum energy ($E_{min}$) to minimize CNT damage.

The optical absorbance served as an indicator of dispersion quality. We used a UV-VIS spectrometer (Jenwey® model 7305) and quartz cuvettes with an optical path of 10 mm measuring the absorbance. Distilled water was the aqueous medium in all the dispersion experiments.

An Ultrasonic Homogenizer Model JY96-IIN (Ningbo Scientz Biotechnology Co. Ltd, China) was used to deliver mechanical energy with a maximum power $P_{max}$ of 150 W in a volume of 10 to 100 ml ($V_m = 100$ ml). The Homogenizer works at 20-25 kHz and has a titanium alloy cylindrical probe with a diameter of 1/4". Equation (1) allowed the calculation of the mechanical energy delivered per ml of suspension. In equation (1): $P\%$ represents a power proportion of the maximum power. $DC\%$ stands up for the duty cycle. The immersion depth of the cylindrical probe in the flask was 4/5 of the total depth. The total working time, $t_{max}$, was 15 minutes in all the experiments.

\[
E\left(\frac{J}{ml}\right) = \frac{t_{min}}{v_{ml} P_{max} W P_{DC}} \left(\frac{60 J}{min W}\right)
\]  

(1)

The CAd’s concentration’s effect on dispersion quality was studied by preparing solutions with different CAd/MWCNTs loading ratios. Then we find the optimal CAd/MWCNTs loading ratio required to avoid CAd stagnation or saturation. A low amount of CAd leads to partial CNTs disentanglement, whereas CAd excess introduces micelles in bulk (Dassios et al. 2015).

Determination of the optimal CAd/MWCNTs loading ratio comprised: (1) 20 ml of water + 0.1 wt % of MWCNT in a flask. (2) Add 0.2 wt% of CAd. (3) set the ultrasonic Homogenizer to deliver 150 J/ml. (4) Measure optical absorption a 300 nm after the zero setting. (5) Repeat steps 1 to 4 while increasing the proportion of CAd in 0.2 wt %, each time until the absorbance value reaches
a plateau. The optimal CAd/MWCNTs ratio corresponds with a point where the absorbance reaches the plateau, or it is close to reaching its maximum value (Yu et al. 2007).

Determination of the minimum mechanical energy \( E_{\text{min}} \) to deliver to suspension in order to mitigate MWCNTs damage during exposure to acoustic waves (Chen et al. 2014) comprised: (1) 20 ml of water + 0.1 wt % of MWCNT in a flask. (2) Add the proportion of CAd, such as the ratio CAd/MWCNT is to equal the optimal proportion. (3) set the ultrasonic Homogenizer to deliver an energy of 30 J/ml. (4) Measure optical absorption a 300 nm after the zero setting. (5) Repeat steps 1 to 4 while increasing the energy in 30 J/ml, each time until the absorbance value reaches a plateau. The mechanical energy to deliver for full dispersion and minimum damage to MWCNT corresponds with a point where the absorbance reaches the plateau, or it is close to reaching its maximum value. The zero setting involved the absorbance of a suspension of \([\text{H}_2\text{O} + \text{CAd}]\) with the same amount of CAd used for preparing the \([\text{H}_2\text{O} + \text{CAd} + \text{MWCNT}]\) suspension (Grossiord et al. 2005). The beaker was positioned in a bath with ice to avoid temperature increases in the bulk solution. A temperature increases influence the dispersion efficiency. Therefore, the preparation of suspensions for the fabrication of the MWCNT-cement mortar composites comprises the delivery of the minimum mechanical energy to bulk suspension and the optimal CAd/MWCNTs loading ratio.

### Indirect tensile test or Brazilian test

Traditional Brazilian test demands cylindrical concrete samples, and its dimensions require the use of a relatively “large” quantity of material. However, nanomaterials are pricey even though they are used in low weight concentrations between 0.05 and 3%. The improved Brazilian test by García et al. (2017) represents a simple, low-cost, robust, and effective characterization technique for MWCNTs-CBM composites.

In the Brazilian test, a compression load concentrated on a pair of antipodal points induces tensile stress in the direction perpendicular to de applied load (Figure 3) (García et al. 2017). The test usually ends with a sudden and violent failure of the specimen

| Table II. Multiwall carbon nanotubes’ properties. |
| --- | --- | --- | --- |
| Inside diameter (nm) | 5-10 | Outside diameter (nm) | 20-40 |
| Length (μm) | 10-30 | Aspect Ratio | 250-1500 |
| Purity (wt%) | >90 | Elastic Modulus (TPa) | 0.8-0.9 b |
| Specific surface area (m²/g) | >80 | Tensile strength (GPa) | 11-63 b |
| Ash (wt%) | <8.0 | Elongation at break (%) | 12 b |
| Bulk density (g/cm³) | 0.16 | Electrical conductivity \( (\Omega m)^{-1} \) | \(-10^5 \) |
| True density (g/cm³) | -2.1 | Thermal conductivity \( W/(mK) \) | >3000 |
| Synthesis process | CVD (HiPCO NTs)a | ~2.1 | ~105 |

* Chemical vapor deposition; High-pressure disproportionation of CO. b (Chen et al. 2011; Hilding et al. 2003). c (D’Alessandro et al., 2016).
Table III. The chemical admixtures’ most relevant properties.

| Comercial Admixture | Based on:                        | Chemical structure of the main component | Type of molecule | Ref.                           |
|---------------------|----------------------------------|------------------------------------------|------------------|--------------------------------|
| 100N                | Hydroxylated polymers (HPs)      | ![chemical structure](image)              | Nonionic         | (Dodson 1990) (Blanch et al. 2010) |
| SF-107              | Sulfonated naphthalene formaldehyde (SNF) | ![chemical structure](image)              | Anionic          | (Dodson 1990) (Giraudeau et al. 2009) |
| FA-111              | Calcium chloride (CC)            | ![chemical structure](image)              | Amphoteric        | (Kurdowski 2014) (Dodson 1990) (Rixom & Mailvaganam 2001) |

Table IV. The mixture proportions for the plain mortar, main aggregate characteristics, design’s specification, and the amount of raw material needed to fabricate one cubic meter of mortar.

| Characteristics                              | Quantity/m³ | Water (l) 240 | 2.69 Fineness modulus (fine aggregate) |
|----------------------------------------------|-------------|---------------|---------------------------------------|
| Water (l)                                    | 240         | 240           | 2.59 Water absorption (%)              |
| Cement (kg)                                  | 462         |               | 0.52 Water/cement ratio                |
| Fine aggregate (kg)                          | 1447        |               | 0.33 Cement/fine aggregate ratio       |
| Compressive strength $f_c$ (MPa)             | 21.00       |               | 19.20 Compressive strength $f_c$ (MPa) at 7 days $^1$ |

$^1$The value is the average of three cylinders tested.
Table V. Sets of samples for the indirect tensile test and electrical conductivity test sample.

| Indirect tensile test —Brazilian test | Electrical conductivity test |
|--------------------------------------|-----------------------------|
| Sample | Admixture | MWCNTs | Disc | Sample | Admixture | MWCNTs | Rectangular |
|        | Type      | wt %   | a     | Disc   | Type      | wt %   | a     |          |
| PM     | -         | -      | 5     | PM     | -         | -      | 3     |
| PM+CAd | HPs       | 0.3    | -     | 5      | PM+CAd   | 0.24   | -     | 3       |
| PM+CAd+CNTs | 0.3 | 0.5 | 5     | PM+CAd+CNTs | 0.24 | 0.4 | 3       |
| PM+CAd+CNTs | 1 | 5     | PM+CAd | 0.48  | -     | 3       |
| PM+CAd+CNTs | 3 | 5     | PM+CAd+CNTs | 0.48 | 0.8 | 3       |
| PM+CAd | 0.6       | -      | 5     | PM+CAd  | 0.6     | -     | 3       |
| PM+CAd+CNTs | 0.6 | 0.5 | 5     | PM+CAd+CNTs | 0.6 | 1 | 3       |
| PM+CAd+CNTs | 1 | 5     | PM+CAd+CNTs | 1.8  | 0.5 | 5       |
| PM+CAd+CNTs | 3 | 5     | PM+CAd+CNTs | 1.8  | 0.5 | 5       |
| PM+CAd+CNTs | 0.55 | - | 5     | PM+CAd+CNTs | 0.55 | 0.5 | 3       |
| PM+CAd+CNTs | 1.1 | - | 5     | PM+CAd+CNTs | 1.1  | 1 | 3       |
| PM+CAd+CNTs | 3 | 5     | PM+CAd+CNTs | 3.3  | 0.5 | 5       |
| PM+CAd+CNTs | 1 | 5     | PM+CAd+CNTs | 2.1  | 0.5 | 5       |
| PM+CAd+CNTs | 3 | 5     | PM+CAd+CNTs | 0.7  | 0.5 | 5       |
| PM+CAd+CNTs | 1 | 5     | PM+CAd+CNTs | 0.7  | 1 | 3       |
| PM+CAd+CNTs | 2.1 | - | 5     | PM+CAd+CNTs | 2.1  | 0.5 | 5       |
| PM+CAd+CNTs | 3 | 5     | PM+CAd+CNTs | 1 | 5     |
| PM+CAd+CNTs | 3 | 5     | PM+CAd+CNTs | 3 | 5     |

*Values expressed as a cement weight percentage.
when the load reaches the value needed to spread an unstable crack (Carmona & Aguado 2012). The induced stress field depends on the specimen’s material properties and the loading geometry (Erarslan et al. 2012). García et al. (2017) reported that it is possible to reduce the loading geometry effect by using the experimental setup shown in Figure 3. Thus, equation (2) allowed the calculation of the indirect tensile strength ($\sigma_{IT}$).

$$\sigma_{IT} = \frac{2P_{\max}}{\pi Dt} \left[ \frac{4 \sin \alpha (\sin^2 \alpha - 3)}{3(8 \sin^3 \alpha + 24 \sin \alpha - 3 \sin 2\alpha - 6\alpha)} \right]$$

; $\alpha \geq 10^\circ$  \hspace{1cm} (2)

In equation (2), $D$ represents the diameter of the disc, $t$ the thickness, $P_{\max}$ is the maximum value of the applied load when the disc fails and $\alpha$ is the angle supported by the contact area ($\alpha = 12^\circ$).

The Brazilian test was performed on a universal testing machine (Versa Tester by Soil Test Inc., USA) with a load cell of 5 kN, in the controlled displacement mode at a speed of 0.02 inch/min ($8.47 \times 10^{-3}$ mm/s).

Electrical conductivity

We used the direct-current four-pole method for measuring electrical resistivity of rectangular prism-shaped samples (80 mm x 20 mm x 20 mm) with four metallic meshes embedded as electrodes conductivity (Han et al. 2007, Konsta-Gdoutos & Aza 2014). Electrical conductivity is the inverse of electrical resistivity. The mesh formed metal grids of $4 \times 4$ mm. The value of electrical conductivity ($\sigma$) calculated from equation (3) is independent of the area of B and C electrodes, as well as the dimensions of the mesh (Figure 4) (B. Han et al. 2007). Thus, the direct-current four-pole method eliminates the influence of contact resistance and provides higher accuracy (Konsta-Gdoutos & Aza 2014).

$$\sigma = \frac{I}{V} \left( \frac{L}{S} \right)$$

In equation (3), $V$ and $I$ represent the voltage between electrodes B and C and the current flowing through electrodes A and D, respectively (Figure 4). $S$ and $L$ represent the specimen’s cross-section and the distance between electrodes B and C, respectively. We measured the electrical conductivity at 20 °C.

The electrical conductivity measuring process comprises of direct-current four-pole method involves (1) application of voltage $V_o$ between electrodes A and D, (2) measurement of the voltage ($V$) between electrodes B and C, (3) measurement of the current $I$ flowing through electrodes A and D, and (4) calculation of the electrical conductivity from equation (6). We measured the electrical conductivity after seven days of curing at 20 °C.

Electron microscopy

The analysis of the MWCNTs morphological attributes was carried out using a TEM (JEM 2100F, JEOL Ltd. Tokyo, Japan). The sample preparation for TEM involved placing a few drops of solution on a carbon film Cu grid and dried in a vacuum desiccator for 48 h. We used the SEM (DSM969A, ZEISS, Oberkochen, Germany) to obtain images of the fracture surface of the disc-shaped sample after the Brazilian test at seven days.

Figure 3. Experimental setup for the Brazilian test.
Data processing and statistical analysis

Statistical data processing was performed using Minitab® 17 statistical software, State College, PA, USA. We generate descriptive data for the Indirect tensile strength and electrical conductivity; and carried out a Pearson’s correlation analysis to assess the strength of the correlation between independent and dependent variables.

RESULTS

Dispersion of MWCNTs in aqueous solution

Individual Cad (SNF, HPs, and CC) are active in the UV-vis region and exhibit a maximum value of absorbance around 300 nm in aqueous solution [H₂O+CAd]. Figure 5a shows the absorption spectra of [H₂O+CAd] solutions. Similarly, MWCNTs are active in the same region, and the [H₂O+CAd+MWCNTs] spectra show a maximum of absorbance around 300 nm. Figure 5b shows the absorption spectra of [H₂O+CAd+MWCNTs] solutions. Although not presented, the corresponding absorption spectrum’s intensity depends on the amount of MWCNTs individually dispersed in solution.

The Cad/MWCNTs ratios’ measured values were 1.1, 0.6, and 0.7 for SNF, CC, and HPs, respectively (Figure 6a). The CC and HPs are more efficient in dispersing MWCNTs than SNF. A Cad/MWCNTs ratio as low as 0.6 (CC) and 0.7 (HPs) is sufficient to separate the MWCNTs in aqueous media. The minimum energy required for optimal dispersion was 270, 150, and 150 J/ml for SNF, CC, and HPs, respectively (Figure 6b). Therefore, the CC and HPs required less energy for optimal dispersion than the SNF’s energy.

Indirect tensile strength

The indirect tensile strength is related to the compression strength, water-cement ratio, and age of the concrete, among other factors (Zain et al. 2002). Figure 7a to 7c show the results of the indirect tensile test performed on the [PM], [PM+CAd], and [PM+CAd+MWCNTs] samples. Symbols represent the mean value of three samples, and the vertical segments are the standard deviation. The indirect tensile strength of [PM] samples averaged 3.85 MPa.

Figure 7a shows the indirect tensile strength of [PM+HPs] and [PM+HPs+MWCNTs] samples. The indirect tensile strength does not change with
the amount of HPs, neither with the amount of MWCNTs. Similarly, the indirect tensile strength of [PM+CC] and [PM+CC+MWCNTs] samples remain similar independent of CC concentration and CC+MWCNTs (Figure 7b). Except for the samples with 0.6 wt% of HPs and 1 wt% MWCNTs, the average indirect tensile strength was 32 % bigger than the average of the [PM] samples. However, suppose the amount of HPs increases to 1.8 wt%, and MWCNTs increases to 3 wt%. In that case, the average value of the indirect tensile strength is 16% bigger than the PM samples’ average. Figure 7c shows the indirect tensile strength of the samples [PM], [PM+SNF], and [PM+SNF+MWCNTs] samples. The indirect tensile strength in the [PM+SNF] samples decreases while the SNF amount increases. However, the indirect tensile strength in the [PM+SNF+MWCNTs] increases while the SNF+MWCNTs amount increases.
Figure 8 illustrates the applied load evolution over time until collapse occurs and during the indirect tensile test on [PM+SNF] and [PM+SNF+MWCNTs] samples. The maximum load that the [PM+SNF] samples can stand before the collapse decreases in inverse proportion with SNF. Thus, the SNF has a deleterious effect on the HE cement-mortar samples. In contrast, the indirect tensile strength of the [PM+SNF+MWCNTs] samples increased as the amount of SNF and MWCNT increased, and samples show and reinforcement effect on the HE cement-mortar.

The zoomed-in images in Figure 8 highlight a loss in curve tendency to change smoothly. This marker is associated with the beginning of a crack. This marker also occurred in the test performed with [PM], but the disc’s fracture occurred immediately. In contrast, in the disc reinforced with SNF and MWCNTS, fracture occurred after the elapse of a considerable time upon the fault had started. The time between the onset of failure and fracture increased with increasing amounts of SNF and MWCNTs.

Figure 9 shows the micrographs of well-connected MWCNTs at the failure surface of the [PM+SNF+MWCNTs] sample. MWCNTs look like bridging different cement base materials units.

**Electrical conductivity**

The electrical conductivity as a function of time was measured when the applied voltage $V_o$ was equal to 30 V. The elapsed time before resistivity stabilized was 5 min (Figure 10a). Thus, we measured the voltage $V$ between electrodes B and C and the current $I$ flowing through electrodes A and D after 5 min for various values of $V_o$ in [PM+CAd] samples (Figure 10b) and [PM+CAd+MWCNTs] samples (Figure 10c). Figure 10b shows that the addition of CAd to the PM mixture decreases the electrical resistance (increase conductivity) of the [PM+CAd] mortar samples regardless of the CAd. While in Figure 10c,
it is apparent that the addition of HPs+MWCNTs to the PM mixture leads to samples with higher resistivity than the [PM] samples. However, the addition of CC+MWCNTs leads to samples with the smallest resistivity.

Figure 11 shows the electrical conductivity of [PM], [PM+CAd], and [PM+CAd+MWCNTs] samples. Symbols represent the mean value of three samples, and the vertical segments are the standard deviation. The mean value of the [PM] sample’s electrical conductivity is $21 \times 10^{-6}$ (Ω cm)$^{-1}$.

**DISCUSSION**

**Dispersion of multiwall carbon nanotubes in aqueous solution**

This study found that the CAd (SNF, HPs, and CC) for concrete are active in the UV-vis region in an aqueous solution $[H_2O+CAd]$ (Figure 5a). The three admixtures exhibit an absorption peak at 300 nm. Moreover, the three solutions $[H_2O+CAd+MWCNTs]$ show similar behavior no matter the CAd and the fact that we used the $[H_2O+CAd]$ as a blank (Figure 5b). There are four possible contributors to the absorption spectrum: (1) free CAd, (2) free MWCNTs, (3) free [CAd+MWCNT], and (4) Mie scattering from CNTs agglomeration with a size comparable to the incident wavelength (Alafogianni et al. 2016). The blank correction with the $[H_2O+CAd]$ reduces the contribution to the spectrum of CAd’s molecules free in solution. Free MWCNTs are unlikely in an aqueous solution after ultrasonication and enough CAd amount. Free big size CNTs agglomerations –around 300 nm– are unlikely after proper ultrasonic processing. In any case, the Mie scattering effect decreases the net absorption. It does not compromise the use of the absorption spectrum as a tool for dispersion quality control. Therefore, the main contributor to the spectrum comes from free [CAd+MWCNT] in solution. Results in figure 5b are in agreement with previous reports by (Zheng et al. 2019). Zheng et al. (2019) used similar MWCNTs from the same provider. They reported a characteristic peak at 300 nm. Li &
Lin (2019) reported a characteristic peak around 250 nm while using MWCNTs with similar lengths and a half diameter. This result hints that the peak position depends mainly on the diameter of the CNTs, as predicted by Kataura et al. (1999). The verification of this hypothesis could lead to a convenient tool for controlling CNTs’ diameter distribution.

The UV-vis spectra of all solutions exhibited a sharp peak from optical resonances in CNTs arising from excitons (Wang et al. 2005). At first, the community thought that these optical resonances arise from transitions between van Hove singularities in the valence and conductive bands of the nanotube’s electronic density of states (Blanch et al. 2010). Therefore, the intensity of absorbance is related to the concentration of MWCNTs in suspensions according to the Lambert-Beer’ law. Results in figure 5b suggest that the adsorption of CAd on MWCNTs leads to the complex [CAd+MWCNT] that is responsible for the stabilization of the dispersion. It includes a weak physical absorption of CAd molecules on the CNTs’ surfaces by either hydrophobic interaction, Coulombic attraction, or π–π stacking interaction (Dassios et al. 2015). Our study considered that the complex [CAd+MWCNT] does not include chemical absorption because it perturbs the CNTs’ electronic structure, potentially changing the characteristic peak wavelength.

Similarly, the absorbance at the peak specific wavelength is related to the degree of de-bundling of CNTs (Alafogianni et al. 2016), CAd concentration, and the energy needed for CNTs de-bundling. However, the dispersion capability of CAd depends on their concentration and an optimum CAd/MWCNTs ratio (Parveen et al. 2013). Besides concentration, the CAd type –nonionic, anionic, cationic, and amphoteric– and its structure significantly influence the dispersion

Figure 10. (a) The elapsed time before resistivity stabilized. (b) Effects of chemical admixtures on the electrical conductivity of PM. (c) Effects of CAd+MWCNTs on the electrical conductivity of PM.
of MWCNTs in water and subsequently within cement-based composites (Parveen et al. 2013).

We found that the CAd/MWCNT ratio for SNF, HPs, and CC was 1.1, 0.7, and 0.6, respectively (Figure 6a). Thus, the HPs (nonionic) and CC (amphoteric) compounds have more dispersion power than SNF (anionic). The optimal CAd/MWCNT ratio is important because the dispersion’s outcome strongly depends on the actual ratio value being below or above this umbral ratio. Suppose that the actual CAd/MWCNT loading ratio is above the optimal ratio. In that case, the dispersion rate of the MWCNTs will increase at the beginning, and the minimum energy required to achieve the maximum dispersion decrease. Under these conditions, the excess CAd is free in the suspension, leading to the flocculation of the MWCNTs, likely through attractive depletion interaction (Blanch et al. 2010, Rastogi et al. 2008). Suppose that the actual CAd/MWCNT loading ratio is below the optimal ratio. In that case, the reverse effect occurs, and the required energy rise. However, under certain circumstances, it is impossible to achieve the maximum dispersion; for example, in suspensions with large amounts of MWCNTs, the optimal amount of CAd required may exceed the critical micelle concentration (Islam et al. 2003). Furthermore, enough CAd is required to produce a good coating of the surface of the MWCNTs and prevent re-aggregation (Yu et al. 2007).

Similarly, we found the minimum energy required for the full dispersion of MWCNTs when SNF, HPs, and CC were 270 J/ml, 150 J/ml, and 150 J/ml, respectively (Figure 6b). The minimum energy required for optimal dispersion agrees with the trend observed for SNF, HPs, and CC’s dispersion power. The energy delivered to the system \([H_2O+CAd+MWCNTs]\) could produce two effects: the deagglomeration of the MWCNTs and the other being fragmentation of individual CNTs (Zheng et al. 2019). Energy above the optimal value affects the integrity of the MWCNTs,
generates defects, and alters their length distribution (Chen et al. 2014) and subsequently within cement-based composites. Similarly, the delivery of energy below the optimal value leads to a low level of disaggregation of MWCNTs and generates defects within cement-based composites.

One interesting finding is that CNTs characteristic peak is not affected by the CA nature –SNF (anionic), HPs (nonionic), and CC (amphoteric). Thus this result suggests that the characteristic peak is due to the photon interaction with MWCNT. Furthermore, this allowed the CA/MWNT ratio determination and the minimum energy for optimal dispersion. These results further support the idea that ultrasound energy drives the disentanglement of CNTs boundless. Also, CNTs disperse via CA adsorption, thermodynamically driven by CA concentrations (Alvarez-Primo et al. 2019).

**Indirect tensile strength**

The results of this study indicate that the indirect tensile strength of [PM+SNF] samples decreased with the SNF concentration (Figure 7c). However, in [PM+SNF+MWCNTs] samples, the indirect tensile strength variation pattern changed and increased as a function of MWCNTs concentration. The other samples’ indirect tensile strength remains approximately the same with or without HPs, CC, or MWCNTs (Figure 7a and 7b).

One finding is that the SNF is incompatible with cement-type HE. This result further supports the idea of at the beginning of cement hydration, C₃A is the most reactive phase in the fresh [PM+SNF] samples. The C₃A surface adsorbs a large amount of SNF. It delays the reaction of C₃A with sulfate and hinders the early strength development of the cement type HE (de Bem et al. 2018, Giraudeau et al. 2009, Pott et al. 2020, Ramachandran et al. 2002b, Yoshioka et al. 2002). The SNF superplasticizer can also interfere and delay the C₃S and C₂S hydration, even though less often and less pronounced than the reaction with C₃A. Besides, SNF’s molecule retards the dissolution between water and Ca²⁺ in the cement-solution interface, inhibiting nucleation and creating Ca²⁺ components (de Bem et al. 2018).

Another finding is that if SNF adsorption onto the surface of MWCNTs occurs first, the retarding effect of the SNF in the cement type HE does not show up. Moreover, the early development of strength in the [PM+SNF+MWCNTs] sample is enabled (Figure 7c). The SNF is a high-range water-reducing superplasticizer and an anionic compound that carry a negative charge on their hydrophilic head. Previous reports suggested that superplasticizer molecules like SNF work through weak physical adsorption on the MWCNTs surface by π-π stacking interaction. Thus, the molecules wrap themselves around the MWCNT, given them a highly negative charge, so they repel each other in an aqueous solution. Upon hydration, they bound the MWCNT to the matrix. Therefore, the [PM+SNF+MWCNTs] indirect tensile strength increased as a function of MWCNTs concentration. The high Pearson correlation coefficient between the indirect tensile strength and the MWCNT wt% (r = 0.99) supports this result. The indirect tensile strength data fit a linear model with R² = 0.98 and a slope of 0.47 (Figure 7c). Therefore, 2 wt% of MWCNTs are needed to improve the indirect tensile strength in 1 MPa. However, the relative improvement will decrease as the design unconfined compression strength increase. Mainly due to the constant bond strength between the MWCNT and the cement matrix. More studies are needed to develop this idea further.

Thus, we found that the [PM+SNF+MWCNTs] indirect tensile strength improves 35% compared to the [PM] samples at the 7 days. Similar results
were reported by Evangelista et al. (2019), who reported an improvement of 27% at 7 days on the indirect tensile strength of MWCNTs reinforced ordinary Portland cement mortar composite. These results could imply that the SNF superplasticizers wrap themselves around the MWCNTs, leading to a mild improvement of the indirect tensile strength (mechanical properties). More studies are needed to confirm this hypothesis.

The results in Figure 7b suggest that CC does not affect the indirect tensile strength of HE cement-mortar composites samples with and without MWCNTs. Although the CC accelerates the setting time and the early strength development, Ramachandran et al. (2002a) reported that this effect is marginal in the cement type HE. Thus, we do not observe any improvement or deleterious effect on the samples tested at seven days. The CC adsorption on the CNT surface involves weak physical adsorption on the CNTs’ surfaces by Coulombic attraction. The CC’s bonding effect is weak and does not improve the indirect tensile strength. This result reflects those of Skripkiunas et al. (2019). They also found that the density, flexural, and compressive strength of cement mortar modified by CC and MWCNT did not change significantly. Skripkiunas et al. (2019) did not identify new bonds and phases in the hardened cement paste modified by CC and MWCNT. Thus, Skripkiunas et al. (2019) concluded that the absence of new bonds and phases is related to the absence of improvement in mortars’ physical properties.

Similarly, this study found that the HPs do not affect the HE cement-mortar composite sample’s indirect tensile strength with and without MWCNTs (Figure 7a). The HPs is a water-reducing plasticizer and a nonionic compound that does not charge its hydrophilic head group. The HPs do not produce ions in an aqueous solution, and the steric effect prevents MWCNTs agglomeration. The HPs adsorption on the MWCNTs surface involves weak physical interaction on the CNTs’ surface by hydrophobic interaction. Thus, the bonding effect of the HPs is weak and does not lead to an improvement in the indirect tensile strength.

**Evolution of the applied load over time**

This study found evidence of the MWCNTs’ potential to control crack propagation phenomena (Figures 8 and 9). The enhancements in the indirect tensile strength of \([\text{PM+SNF+MWCNTs}]\) samples are related to the bridging effect of MWCNTs across micro-cracks and pores. MWCNTs form a network (Figure 9) capable of enhancing load transmission capacity through the matrix. Moreover, nanotubes give additional strain capacity to the matrix by dissipating energy through multiple pulls out of nanotubes from the matrix before reaching a critical state (Figure 8). Rodríguez et al. (2017) reported similar results regarding the bridging effect of CNTs. Therefore, the ability of \([\text{PM+SNF+MWCNTs}]\) samples to maintain their integrity after the crack initiation suggests an interplay between the MWCNTs and the hydrated products mediated by a strong bond (Figure 9). This bond is due to the nucleation of hydrate products (C-S-H) by the MWCNTs. Therefore, CNTs connect the hydrated products, enhancing the bridging effect between pores (Jeevanagoudar et al. 2017, Li & Lin 2019). The micrograph in Figure 9 confirms that MWCNTs are distributed and are attached to the hydrated products. More studies are needed to analyze the interface between the surface of MWCNTs and the hydrate products in cement-based composites. The strength of the bonding determines the reinforcement effect of CNTs.
Electrical conductivity

This study found that the electrical resistivity (The inverse of the conductivity) of the [PM], [PM+CAd], and [PM+CAd+MWCNTs] samples stabilized after 5 min has elapsed (Figure 10a). The time of 5 min is similar to that found by D’Alessandro et al. (2016) for [PM+CAd+MWCNTs] samples. In general, the variation in resistivity within the first few minutes after the voltage is applied is mainly due to the materials’ dielectric properties exhibiting a polarization effect. The polarization effect depends on CAd, MWCNTs, water, and ion contents in the material’s matrix and the temperature, among others (Han et al. 2011).

Another finding is that HPs, CC, and SNF decreased the electrical resistivity in the [PM+CAd] samples (Figure 10b). This result suggests that the electrical resistivity of the CAd is lower than the PM. This finding agrees with Chung’s conclusion that the electrical conductivity of CBM is governed by (1) the degree of admixture dispersion, (2) the contact electrical resistivity of the interface between the admixture and the cement matrix, and (3) the admixture itself (Chung 2004). Thus, the mortar electrical resistance is affected by CAd electrical conductivity and ions in the matrix.

In contrast, when using small amounts of CAd and MWCNTs, the tendency to reduce the resistivity depended on the CAd (Figure 10c). Thus, this study found that CC and MWCNTS cause [PM+CC+MWCNTs] samples electrical conductivity to increase significantly (507 %) compared with the conductivity of [PM] samples (Figure 11b). The electrical conductivity of the [PM+CC+MWCNTs] samples increased as a function of MWCNTs concentration. The high Pearson correlation coefficient between the electrical conductivity and the MWCNTs wt% (r = 0.91) supports this result. The electrical conductivity data fit a linear model with R² = 0.96 and a slope of 92.5 (Figure 11b). In contrast, there were mild increases in electrical conductivity of [PM+HPs+MWCNTs] and [PM+SNF+MWCNTs] samples (Figures 11a and 11c).

Many factors affect the electrical conductivity of MWCNTs cement mortar composites. Intrinsic factors like water/cement ratio, aging, pore structure, CAd and CNTs electrical conductivity, CAd and CNTs concentration, and the degree of MWCNTs dispersion. Also, the transition zone between the cement paste and the aggregate or CNTs might enhance the conductivity (Chung 2004). Extrinsic factors affect the electrical conductivity measurements, including specimen geometry, moisture content, temperature, and electrode spacing (Azarsa & Gupta 2017). This study keeps the water/cement ratio constant and measures the electrical conductivity seven days after preparation. The porous structure is one of the main parameters affecting the electrical resistivity in CBM (Dalla et al. 2019). However, Skripkiunas et al. (2019) show that in a cement mortar modified by MWCNT and CC in the amount 2% by weight, the total pore volume per unit volume decreased by 2.1 %. Additionally, the open capillary porosity increased by 1.6%. Thus, the CC did not affect the pore structure of the samples significantly. However, the results in Figure 10b show that the [PM+CC] electrical conductivity is higher than the [PM], suggesting that the conductivity of CC is higher. Figure 10b shows that the electrical conductivity of the [PM+CC+MWCNTs] is even higher than the [PM+CC] and [PM]. One explanation is that the electrical conductivity of the MWCNTs (10¹⁵ S/m) is many orders of magnitude higher than the conductivity of the [PM] (2x10⁻³ S/m) and the [PM+CC] (5x10⁻³ S/m). This contrast supports the fact that electron transport in [PM+CC+MWCNTs] samples occurs through an MWCNTs network in the matrix, consisting of dispersed MWCNTs (Jang et al. 2019).
In particular, electrical conductivity increases by the electrical tunneling between adjacent MWCNTs and, consequently, depends on the tunneling gap and the cement matrix’s conductivity (Yoo et al. 2019). Nevertheless, MWCNTs behave as non-conductive inclusions when their mutual distances are high. When the MWCNTS concentration increases above the percolation level and their mutual distances decrease, nanotubes interactions dominate the composite’s electrical conductivity (D’Alessandro et al. 2016). Figure 10b shows that [PM+CC+MWCNTs] electronic conductivity suddenly increased when the CNTs concentration went from 0.7 to 1 wt %, suggesting that the MWCNTs percolation level is around 0.7 wt %. Therefore, before the CNTs reach a concentration of 0.7 wt %, conductivity is governed by CC; when the CNTs concentration is above 0.7 wt %, MWCNTs govern the conductivity. Therefore, the conductivity of HE cement mortar can be improved by incorporating CC and MWCNTs. However, to the best of the authors’ knowledge, there is no published study regarding the electrical resistivity of HE cement-CC-MWCNTs mortar composites. A note of caution is due here since chloride diffusivity compromise durability. The chloride threshold establishes a limit for the chloride ion in the matrix (Dalla et al. 2019, Abd El Fattah et al. 2020). Further study is still needed to understand CC and MWCNTs interaction to make conductive MWCNT-HE cement-based composite practical.

CONCLUSIONS

This study used three available commercial CAds used for concrete: (1) HPs based water-reducing plasticizer, (2) SNF-based superplasticizer, and (3) CC-based accelerating agent. The three CAds dispersed MWCNTs in an aqueous solution. Ultrasonication and the optical absorption spectroscopy allowed the determination of optimal CAd/MWCNTs loading ratio and minimal energy for optimal CNTs dispersion. The MWCNTs dispersion occurs by the combined effect of ultrasonication energy and CAd adsorption on the nanotube surface.

The HPs based water-reducing plasticizer and the CC-based accelerating agent are compatible with HE cement. In contrast, the SNF-based superplasticizer is incompatible with the HE cement. The indirect tensile strength decreased when the SNF concentration increased.

HPs and CC’s adsorption on the MWCNTs surface disperse CNTs in aqueous solution via repulsive interaction among [HPs+MWCNTs] and [CC+MWCNTs] complexes. While the limited interference of [HPs+MWCNTs] and [CC+MWCNTs] complexes on the hydration of the HE cement particles do not improve or damage the indirect tensile strength of CNTs reinforced CBM.

The adsorption of SNF-based superplasticizer on the MWCNTs surface disperses CNTs in aqueous solution via repulsive interaction among [SNF+MWCNTs] complexes. While the interference of [SNF+MWCNTs] complexes on the HE cements particles’ hydration improves, the indirect tensile strength of CNTs reinforces CBM.

The addition of HPs, CC, SNF based CAd to the PM mixture decreases the electrical resistance (increase conductivity) of the [PM+CAd] mortar samples regardless of the CAd.

The adsorption of the CC-based accelerating agent on the MWCNTs surface favors a substantial improvement in the electrical conductivity of CNTs reinforced CBM composite. The results suggest that the [CC+MWCNTs] conductivity governs the electrical conductivity of [PM+CC+MWCNTs] composite. Moreover, further studies should verify if, below the
threshold concentration of \([\text{CC+MWCNTs}]\) complexes, the CAd's ionic contribution governs the composite's conductivity. While above the threshold concentration of \([\text{CC+MWCNTs}]\) complexes, the CNTs' conductivity controls the composite's conductivity through closer CNT electrical tunneling. Therefore, engineering the \([\text{CAd+MWCNTs}]\) complexes' electrical properties could improve \([\text{PM+CAd+MWCNTs}]\) composites' electrical conductivity properties. More studies are needed to know how the \([\text{CAd+MWCNTs}]\) complexes' properties control the improvement of the \([\text{PM+CAd+MWCNTs}]\) composite properties.

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