Tensile Modulus of Polymer Halloysite Nanotube Systems Containing Filler–Interphase Networks for Biomedical Requests

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Abstract: To date, there have been a limited number of studies modeling the tensile modulus in the polymer halloysite nanotube (HNT) systems before or after percolation onset. In this paper, an equation for a composite’s modulus post-percolation onset was developed for HNT-filled samples including the interphase and HNT network. The dispersed nanoparticles and adjoining interphase part were neglected, because they caused ineffective influences on the stiffness of the system after percolation onset. The developed model reflects the impacts of HNTs’ size, interphase depth, percolation onset and the volume shares and moduli of the HNT network and its adjacent interphase on the modulus of HNT-based systems. The impacts of issues on the nanocomposite modulus are defendable, confirming the effectiveness of the developed model. HNT length, interphase depth, HNT concentration, net modulus and net portion directly influenced the stiffness, while the HNT radius and percolation onset had inverse effects. Results show that there was a 142% improvement in the modulus of samples at an interphase depth of 40 nm. Moreover, the stiffness improved by 60% at a net modulus of 200 GPa, but it later exhibited a 180% enhancement at a net modulus of 1000 GPa. In addition, the experimental data for the modulus of numerous composites display fine agreement to the predictions, confirming the validity of the developed model.

Keywords: halloysite nanotube (HNT); nanocomposite; modulus; interphase zone; network

1. Introduction

The tough polymer–filler interfacial network, besides the large nanofiller superficial zone, often changes the polymer’s features due to adjacent nanoparticles leading to the formation of the interphase section [1–15]. This intermediate phase is different from both polymer media and nanoparticles. Several authors have tried to examine the interphase effects on the mechanics of nanocomposites [16–18]. The modeling approach in this field is highly challenging and interesting, since the analysis at the nanoscale in the interface region is a difficult task. Moreover, the interphase section around large and slender nanofiller such as halloysite nanotubes (HNTs) facilitates the development of networks within the sample, where the interphase eases the percolation onset and interlinks the nanoparticles before the physical contact [19–22]. However, the interphase section and its networking effectiveness for HNT-based systems were not argued in the previous articles. Due to the main reinforcing competences of the interphase section and filler network in nanocomposites, it is important to study them in HNT-filled systems. Figure 1 displays the location of nanotubes in a nanocomposite before, during and after percolation onset.
Many researchers have stated that the mixture of HNTs with polymer media enhances the thermal and mechanical properties [41–44]. Prashantha et al. [45] mixed neat HNT and quaternary ammonium salt-treated HNTs with polypropylene. They showed a fine dispersion of treated HNTs in comparison to the neat one, because of the tough interfacial network among polypropylene and treated HNTs. Krishnaiah et al. also functionalized the HNT with 3-aminopropyltriethoxysilane to improve the surface contact of the HNT with polypropylene and polylactide [46,47]. They also demonstrated that the fine dispersal of the treated HNT in the polymer matrix improved the mechanical and thermal performance of the composite.

Several authors have focused on the mechanical features of HNT-based systems through an experimental approach only [48–50]. Nevertheless, the theoretical models for this type of material and system are restricted due to insufficient scientific data and reports. Very few papers have considered the mechanical aspects of HNT nanocomposites, such as tensile strength and moduli [51,52]. Modified Halpin–Tsai and Pukanszky equations were utilized for calculating the tensile powers of a polylactide HNT system [51]. It was stated that an altered Halpin–Tsai equation overestimates the modulus, but the Pukanszky equation estimates acceptable results for strength. Additionally, it was suggested that a 3D finite element technique for polylactide HNT samples be included, hypothesizing an arbitrary HNT and assessing its predictive results [52]. Clearly, the modeling values for polymer HNT nanocomposites are highly insufficient, restricting the development of these materials and their applications. Therefore, novel theoretical models are an obligatory requirement for carrying out detailed scientific evaluation of the simulated interphase zone and networked HNTs.

Recently, our group published some papers on the modulus of polymer nanocomposites containing CNTs and HNTs [53,54]. We developed the Ouali model by considering the interphases around the dispersed and networked nanoparticles for the modulus of CNT-reinforced samples [53]. Moreover, we enhanced the advanced Takayanagi equation for calculating the modulus of HNT-based nanocomposites by considering the interphase around the dispersed and networked HNTs after percolation [54]. However, an Ouali model for the modulus of HNT-filled samples after percolation onset has not been developed yet. In this work, an Ouali model was developed for the calculation of the tensile
modulus of polymer HNT systems after the percolation threshold by considering the function of networked HNTs and the surrounding interphase. The dispersed nanoparticles and surrounding interphase part were neglected, since they led to ineffective results for the system’s modulus. The formulated model includes the HNT size, interphase depth, percolation onset and HNT concentration as well as the volume portions and moduli of the networked HNTs and adjacent interphase. The components of the whole system’s modulus are explained to validate the formulated equation. Moreover, the tentative results of real samples are utilized to evaluate the model’s predictions. We believe that by employing this simple methodology in the current paper, we can develop a novel modeling method for the approximation of moduli in HNT-based systems.

2. Advanced Equations

The percolation onset of CNTs in the nanocomposites was connected to CNT size and interphase depth [55] by:

$$\phi_p = \frac{\pi R^2 l}{\frac{32}{3} \pi (R + t)^3 \left[1 + \frac{3}{4} \left(\frac{l}{R + t}\right)^2 + \frac{3}{32} \left(\frac{l}{R + t}\right)^2\right]}$$

where “R” and “l” demonstrate the radius and length of the HNT, individually. Additionally, “t” is interphase depth. This equation is usable for the HNT system, since both CNTs and HNTs have a similar tubular shape, and the percolation onset mainly correlates to filler shape and the surrounding interphase section. The interfacial adhesion by hydrogen, van der Waals or covalent bonds is considered by the interphase depth and modulus. Additionally, a thicker and stronger interphase presents stronger interfacial bonds within the samples.

The whole volume fraction of the interphase in the system comprising a tube-like filler such as an HNT is given [55] by:

$$\phi_i = \phi_f \left(1 + \frac{t}{R}\right)^2 - 1$$

where “$$\phi_f$$” is HNT volume fraction. This equation speculates the extent of the HNT and interphase besides the HNT amount, as expected.

The interphase area acts as an imperative, reinforcing efficiency in nanocomposites. Hence, both the HNT and nearby interphase toughen the specimens. The actual HNT amount equates the entire quantity of the reinforcing elements as:

$$\phi_{eff} = \phi_f + \phi_i = \phi_f \left(1 + \frac{t}{R}\right)^2$$

which regulates the reinforcing methods of the nanofiller and interphase by one factor.

Additionally, the proportion of a tubular nanofiller such as the HNT producing the network in the sample is expressed [56] by:

$$f = \frac{\phi_f^{1/3} - \phi_p^{1/3}}{1 - \phi_p^{1/3}}$$

The effective filler portion is further modified and given by the latter equation as:

$$f = \frac{\phi_{eff}^{1/3} - \phi_p^{1/3}}{1 - \phi_p^{1/3}}$$

Further, the original Ouali model and its developed form are expressed by the mentioned factors.
Ouali et al. [57] designed a model for the composite’s modulus by using the following percolation concept:

$$E = \frac{(1 - 2\psi + \psi \phi) E_m E_f + (1 - \phi_f) \psi E_f^2}{(1 - \phi_f) E_f + (\phi_f - \psi) E_m}$$

(6)

where “$E_m$” is the medium’s modulus, and “$E_f$” denotes the modulus of the filler. “$\psi$” correlates to percolation onset ($\phi_p$), as given by:

$$\psi = 0$$

$$\phi_f \leq \phi_p$$

(7)

$$\psi = \phi_f \left(\frac{\phi_f - \phi_p}{1 - \phi_p}\right)^b$$

$$\phi_f > \phi_p$$

(8)

where “$b$” is an exponent, which obtains a value of 0.4 for a 3D structure [57].

When $\psi = 0$, this model diminishes the converse roles of mixtures as:

$$E = \frac{E_m E_f}{(1 - \phi_f) E_f + \phi_f E_m}$$

(9)

In our previous article [53], we developed the Ouali model using the isolated and networked nanofiller and neighboring interphase pieces after percolation onset. It was observed that the volume portions and moduli of the detached filler and adjoining interphase were ineffective on the modulus of nanocomposites [53]. Indeed, only the networks of particles and interphase controlled the stiffness of systems after percolation onset. Therefore, only the networked filler and surrounding interphase have the reinforcing components in the nanocomposites.

These terms extend the Ouali model to:

$$E = \frac{(1 - 2\phi_N + \phi_N \phi_i) E_m E_N + (1 - \phi_N) \phi_N E_N^2 + (1 - 2\phi_iN + \phi_iN \phi_i) E_m E_iN + (1 - \phi_iN) \phi_iN E_iN^2}{(1 - \phi_N) E_N + (\phi_N - \psi) E_m + (1 - \phi_iN) E_iN + (\phi_N - \psi) E_m}$$

(10)

where “$\phi_N$” and “$E_N$” are the volume portion and modulus of networked HNTs, in that order. Additionally, “$\phi_iN$” and “$E_iN$” show the volume share and modulus of the interphase nearby the HNT network, correspondingly. Equation (10) considers the imperative components of the network and interphase.

Based on Equation (8), “$\psi_N$” and “$\psi_iN$”, after percolation starts, progress to:

$$\psi_N = \phi_f \left(\frac{\phi_f - \phi_p}{1 - \phi_p}\right)^{0.4}$$

(11)

$$\psi_iN = \phi_i \left(\frac{\phi_i - \phi_p}{1 - \phi_p}\right)^{0.4}$$

(12)

considering the roles of percolation onset and the amounts of both the HNT and the interphase.

Additionally, “$\phi_N$” and “$\phi_iN$” are calculated by:

$$\phi_N = f \phi_f$$

(13)

$$\phi_iN = f \phi_i$$

(14)
When “f” from Equation (5) is used as a replacement in the latter equations, “φ_N” and “φ_{IN}” are given by:

\[
φ_N = \frac{φ_{eff}^{1/3} - φ_f^{1/3}}{1 - φ_f^{1/3}}
\]
(15)

\[
φ_{IN} = \frac{φ_{eff}^{1/3} - φ_f^{1/3}}{1 - φ_f^{1/3}}
\]
(16)

assuming the roles of HNT size, interphase depth, HNT concentration and percolation in the amounts of HNTs and interphase nets.

The relative modulus, \( E/E_m \), modifies the developed model (Equation (10)) to:

\[
E_R = \frac{(1 - 2φ_N + ψ_Nφ_N)E_N + (1 - φ_N)ψ_NE_m^2/E_m + (1 - 2φ_{IN} + ψ_{IN}φ_{IN})E_{IN} + (1 - φ_{IN})ψ_{IN}E_{IN}^2/E_m}{(1 - φ_N)E_N + (φ_N - ψ_N)E_m + (1 - φ_{IN})E_{IN} + (φ_{IN} - ψ_{IN})E_m}
\]
(17)

which undertakes the impacts of the networked HNT and neighboring interphase section on the improved stiffness of the HNT-based system. The above equations were developed for tubular nanofillers. In fact, the developed equations are only valid for the nanocomposites containing tubular nanoparticles.

3. Results and Discussion

The effects of all of the factors on the modulus of HNT-reinforced composites were elucidated to certify the formulated model.

Figure 2 displays the “\( E_R \)” estimated by the novel model at unlike values of HNT content and constant ranges of \( E_m = 2 \) GPa, \( R = 20 \) nm, \( l = 2 \) μm, \( l = 20 \) nm, \( E_N = 500 \) GPa and \( E_{IN} = 100 \) GPa. After the addition of HNT content, the stiffness increased. “\( E_R \)” reached 2.32 at the HNT volume share of 0.025. This level indicates that the HNT volume fraction of 0.025 increased the stiffness by 132%. However, the relative modulus had a low level at the minor ranges of HNT content.

![Figure 2. Predictions of “\( E_R \)” at various HNT volume portions.](image)

This observation is reasonable, since the HNT was exceptionally firmer than the polymer matrix, and the adding of the HNT to the matrix enhanced the stiffness of samples. In fact, the extremely high difference between the moduli of the HNT and polymer caused the upgrading of the modulus in the samples, due to the better shifting of the load from polymer media to the stiff nanoparticles via the interphase section. All models for mechanical performance of the system demonstrated the same trend between the stiffness/strength...
of the system and filler concentration, since stiff particles commonly reinforce the system [58–60]. Accordingly, the advanced model logically estimated the effect of the HNT amount on the stiffness.

Figure 3 expresses the effect of interphase depth on the \( E_R \) at \( R = 20 \text{ nm}, \phi_f = 0.02, \ E_m = 2 \text{ GPa}, \ l = 2 \text{ \( \mu \text{m} \)}, \ E_{IN} = 100 \text{ GPa} \) and \( E_N = 500 \text{ GPa} \). In the absence of the interphase area (\( t = 0 \)), the relative modulus of 1.68 was calculated. Conversely, the relative modulus increased to 2.42 at \( t = 40 \text{ nm} \). As a result, the nanocomposite’s modulus significantly grew by thickening the interphase zone. This evidence indicates that the interphase depth is a vital factor in improving the stiffness of HNT-based systems. In fact, it is crucial to provide a dense interphase to enhance the modulus of HNT systems.

The interphase depth had positive effects on the effectiveness of HNT concentrations, percolation onset and the networking of the interphase (see Section 2). In fact, a thick interphase decreased the percolation onset and boosted the volume of the interphase zone in the system. During this stage, a larger interphase produced a more prominent network, which undoubtedly increased the stiffness of the system. In other words, a thicker interphase yielded an improved network in the system, thus increasing the resistance of the system against the load bearing. Instead, a narrow interphase created a small network, which undoubtedly diminished the rigidity of sample, owing to little stress bearing. This explanation validates the advanced equation for the modulus of this system.

Figure 4 plots the estimations of \( E_R \) at various orders of the HNT radius and \( E_m = 2 \text{ GPa}, \phi_f = 0.02, \ t = 20 \text{ nm}, \ l = 2 \text{ \( \mu \text{m} \)}, \ E_N = 500 \text{ GPa} \) and \( E_{IN} = 100 \text{ GPa} \) according to the advanced model. An HNT radius of 10 nm caused \( E_R = 2.45 \), while the \( E_R \) was reduced to 1.68 at an HNT radius of 40 nm. Consequently, the HNT radius undesirably managed the stiffness, and the slender HNTs produced the stiffer samples. This observation is also crucial, since nanoparticles such as HNTs are inclined to aggregation/agglomeration during the fabrication of nanocomposites [61,62]. Indeed, it is vital to control the size of HNTs to obtain significant reinforcement within the sample.

The desirable role of thin HNTs in the modulus of samples is logical, because thin HNTs decrease the percolation onset and widen the interphase. According to the developed equations, thin HNTs produce a lower percolation onset and a highly effective filler fraction, which increases the network portion in the composites. Therefore, narrow HNTs certainly increase the modulus of samples, since they expand the interphase and the network stiffening within the samples. Conversely, thick HNTs cause slender interphase sections and few network interactions, which loosely reinforce the system. According to this explanation, the advanced model produced reasonable outputs at dissimilar levels of HNT radii.
HNT length influenced the percolation onset in the system. Large HNT shifted the percolation onset to a low filler concentration, which positively influenced the fraction of networked HNTs. So, it can be said that large HNTs produced a huge network within the system, whereas the short HNTs caused the small one. Since the extent of the network mainly controls the load bearing in the system, big HNTs advantageously influenced the stiffness of the system. Instead, short HNTs slightly manipulated the size of the network, yielding a minor reinforcing capacity. Generally, a higher aspect ratio of nanoparticles (length per diameter) obtains stiffer samples, due to the positive effects of large and narrow nanoparticles on the extents of the interphase section and networks [63,64] (see Section 2). Hence, the novel equation suitably predicted the role of HNT length in the rigidity of the system.

**Figure 4.** Influence of filler radius on the “$E_R^{m}$”.

Figure 5 displays the discrepancy of “$E_R^{m}$” at various orders of the HNT radius and $\phi_f = 0.02$, $E_m = 2$ GPa, $t = 20$ nm, $R = 20$ nm, $E_{IN} = 100$ GPa and $E_N = 500$ GPa using the current model. $E_R = 1.843$ was observed at $l = 0.8$ μm, but “$E_R^{m}$” increased to 1.972 at $l = 3$ μm. So, HNT size directly managed the stiffness, and large HNTs were needed to reinforce the samples. The large variation in HNT length from 0.8 to 3 μm only varied the relative modulus by about 0.13, which is a small amount.

**Figure 5.** Dependency of relative modulus on the HNT length.
Figure 6 plots the effect of the net modulus on the “$E_R$” at $E_m = 2$ GPa, $R = 20$ nm, $\phi_f = 0.02$, $l = 2$ μm, $t = 20$ nm and $E_{iN} = 100$ GPa using the innovative model. The “$E_R$” of 1.6 was developed at a net modulus of 200 GPa, but “$E_R$” significantly grew to 2.8 at $E_N = 1000$ GPa. These outputs display the direct correlation between the moduli of system and its network. A stronger network produced a stiffer sample, whereas poor networks ineffectively reinforced the sample. The big range of the relative modulus from 1.6 to 2.8 verifies that the net modulus is important for the reinforcement of the system. Consequently, researchers should try to extend the HNT network in the system to achieve a better stiffening efficiency.

![Figure 6](image_url)

Figure 6. Relation between the moduli of HNT network and whole system according to Equation (17).

The stiffer network in the nanocomposite bore higher amounts of stress. However, the poorer network could not stand against the stress and breaks during loading. Therefore, the stiffer network induced the higher modulus in the system, but the poorer network contrarily weakened the sample. In fact, there was a direct link between the moduli of the network and the whole system; meanwhile, the stiffness of the network mainly manipulated the stiffness of the whole system. This explanation confirms the approximations of the model at various ranks of net modulus. This relation was also reported in the earlier research on the stiffness/strength of samples [53,65].

Figure 7 expresses the model’s estimations correlating to the interphase modulus at $R = 20$ nm, $\phi_f = 0.02$, $E_m = 2$ GPa, $t = 20$ nm, $l = 2$ μm and $E_N = 500$ GPa. The relative modulus was 1.9 at $E_{iN} = 50$ GPa; nonetheless, the “$E_R$” increased to 2.33 at $E_{iN} = 250$ GPa. These calculations establish that the stiffness grew by 43% when the modulus of the interphase net varied from 50 to 250 GPa. As a result, the stiffness of the networked interphase directly influenced the overall stiffness. This output indicates that the modulus of the interphase network is a main factor influencing the modulus of the whole sample.

The modulus of the networked interphase is also imperative, since the interphase section around the network is a bridge transferring the stress from poor polymer media to the HNT network. A firmer interphase zone neighboring the network undoubtedly transferred a higher stress amount, but a defective interphase declined during stress charging. Hence, a tough interphase network positively grew the modulus of the whole system, since it can tolerate further stress. In contrast, a poorer interphase network accepted a lower volume of stress, which is inefficient for the stiffness of the whole system. Therefore, the innovative model suitably predicted the effects of the interphase net modulus on the stiffness of the system.
Figure 7. Outputs of Equation (17) at numerous values of interphase net modulus.

Figure 8 projects the results of the present model at several values of percolation onset at constant $R = 20$ nm, $E_m = 2$ GPa, $\phi_f = 0.02$, $t = 20$ nm, $l = 2 \mu$m, $E_N = 500$ GPa and $E_{IN} = 100$ GPa. “$E_R$” was 2.01 at the percolation onset of 0.001, but the relative modulus declined to 1.81 when the percolation onset increased to 0.01. Therefore, a higher modulus was observed at a lower percolation onset, and it is essential to decrease the percolation onset, leading to stiffness. Indeed, the percolation onset adversely controlled the modulus of samples (see Section 2). This occurrence is expected, since a low percolation beginning boosts the networked HNT component (Equation (5)). A lower percolation onset increases the network size, which increases the modulus. Contrariwise, a high percolation induces the smaller networks of HNT and interphase, due to the low value of “$f$” (see Equation (5)) in this condition. Consequently, the percolation onset unfavorably governs the stiffness, due to its effect on the extent of network. This description validates the results of Equation (17) at several ranges of percolation onset.

Figure 9 establishes the role of “$f$” as the portion of the filler–interphase network in the modulus of the system. $f = 0.1$ produced $E_R = 1.94$; nonetheless, the “$E_R$” rose to 1.958 at $f = 0.8$. These results signify the direct connection between the stiffness and the share of
networked HNTs or interphase. However, numerous points of “f” from 0.1 to 0.8 slightly changed the relative modulus.

“f” determines the size of networks (both interphase and HNT) in the system. A large “f” depicts the contribution of numerous HNTs and a large interphase to the network (Equations (15) and (16)). So, a larger “f” shows the creation of larger networks of HNT and interphase within the system, which certainly raises the modulus. Conversely, a lower “f” reveals the involvement of few HNTs or decreased interphase section in the network, which is undesirable for the stiffness of system. In summary, “f” reveals the size of networked interphase and HNTs, and thus “f” directly controls the stiffness of samples, due to the effective role of network size in the load bearing. Consequently, the novel model appropriately foresaw the impact of “f” on the stiffness of HNT-based samples.

Now, the outputs of Equation (17) are correlated to the tentative numbers of the modulus for real examples. Table 1 depicts HNT systems and their details (\(E_m, R\) and \(l\)) from valid published articles. Further details about this system are accessible in the original references. These data are substituted into the developed equations and the final model (Equation (17)) to analyze the “\(E_R\)” for the required study specimens. Figure 10 displays the measured and hypothetical extents of “\(E_R\)” for the given examples. It is depicted that all theoretical values matched the measured numbers at entire HNT weight fractions. Hence, the innovative model successfully predicted the modulus of real samples, and thus the developed model offers a valid equation for the modulus of HNT-filled systems when compared to experimental values.

**Table 1.** Properties for HNT examples and the estimations of developed equations.

| No. | Samples [Ref.] | \(E_m\) (GPa) | \(R\) (nm) | \(l\) (μm) | \(t\) (nm) | \(\phi_p\) | \(E_N\) (GPa) | \(E_N\) (GPa) |
|-----|----------------|--------------|-----------|----------|--------|----------|-------------|-------------|
| 1   | PA12/HNT [66]  | 1.55         | 35        | 1.0      | 3.0    | 0.024    | 20          | 150         |
| 2   | Cellulose/HNT [67] | 1.80        | 25        | 2.0      | 6.0    | 0.009    | 40          | 250         |
| 3   | Starch/HNT [68] | 0.08         | 35        | 1.2      | 9.0    | 0.018    | 25          | 150         |
| 4   | PLA/HNT [51]   | 2.90         | 40        | 1.2      | 15     | 0.018    | 45          | 220         |
The modulus of the interphase network was estimated by the predictions of the current model. When the experimented moduli of nanocomposites were compared to the developed model, numerous values for \( E_{iN} \) were obtained, and the average interphase modulus was reported for each sample. The values of the interphase net modulus were also shown to range from 20 to 45 GPa. The strongest and the weakest interphase networks were found in PLA and PA12 samples, respectively. Furthermore, the modulus of the HNT network in the samples was obtained and found to range from 150 to 250 GPa. The strongest network was observed in the cellulose system, but PA12 and starch examples show the same network modulus for 150 GPa. The firm and vast networks of the interphase section and HNT led to more significant improvements in the modulus of the samples, thus equally validating the existing data obtained with the newly modified equations.

Figure 10. Experimental “\( E_R \)” and the calculations of established technique for (a) PA12 [66], (b) cellulose [67], (c) starch [68] and (d) PLA [51] systems containing HNT.

The values of the interphase and network properties were also calculated, as listed in Table 1. “\( t \)”, as interphase depth, changed from 3 to 15 nm for the current samples. These results are significant, since they were obtained at the nanoscale. The thickest and the thinnest interphases were detected for polylactide (PLA) and polyamide 12 (PA12) samples, respectively. When HNT size and interphase depth were replaced in Equation (1), the percolation onset was estimated. The lowest and the highest percolation were found for cellulose and PA12 samples, correspondingly. These outputs indicate that the network in cellulose samples is established at an extremely low HNT content. However, the network in PA12 needs more HNTs (high HNT concentration) to be constructed, due to the low level of interphase depth.

The modulus of the interphase network was estimated by the predictions of the current model. When the experimented moduli of nanocomposites were compared to the developed model, numerous values for “\( E_{iN} \)” were obtained, and the average interphase modulus was reported for each sample. The values of the interphase net modulus were also shown to range from 20 to 45 GPa. The strongest and the weakest interphase networks were found in PLA and PA12 samples, respectively. Furthermore, the modulus of the HNT network in the samples was obtained and found to range from 150 to 250 GPa. The strongest network was observed in the cellulose system, but PA12 and starch examples show the same network modulus for 150 GPa. The firm and vast networks of the interphase section and HNT led to more significant improvements in the modulus of the samples, thus equally validating the existing data obtained with the newly modified equations.
4. Conclusions

The reported and earlier published literature studies have ignored the interphase–HNT networks in the HNT-reinforced systems. Moreover, they used complex and undefined equations, which baffled readers. In this work, the Ouali model was developed for a polymer HNT system assuming the networks of the interphase and nearby HNT after percolation. The developed model in the current study considers the properties of interphase–HNT networks after percolation onset through simple, meaningful equations and parameters, which is the prime novelty and advantage of this study. It is highly evident from the data that the HNT content directly influenced the system modulus, and $E_R$ reached 2.32 at an HNT volume share of 0.025. $t = 0$ led to the “$E_R$” of 1.68, although the “$E_R$” increased to 2.42 at $t = 40$ nm. So, a stronger interphase is vital to obtaining a better modulus. An HNT radius of 10 nm produced $E_R = 2.45$, while the “$E_R$” was reduced to 1.68 at an HNT radius of 40 nm. Thus, narrow HNTs are desirable for the system. HNT length directly controlled the stiffness, but it had a non-significant effect. The “$E_R$” was 1.6 at a net modulus of 200 GPa; however, the “$E_R$” considerably improved to 2.8 at $E_N = 1000$ GPa. Additionally, $E_R = 1.9$ at $E_{IN} = 50$ GPa, but the “$E_R$” grew to 2.33 at $E_{IN} = 250$ GPa. Consequently, the stronger networks of HNTs or surrounding interphase section produced a stiffer sample. A higher “$E_R$” was observed at a lower percolation onset, and it is important to decrease the percolation onset to reinforce the system. There was a direct connection between the stiffness and the portion of networks. The predicted calculations of Equation (17) acceptably match the experimental data of many recorded systems. This is another indication for the significant validation of Equation (17). The highest HNT content, the strongest interface between the polymer and HNT (the thickest interphase), the thinnest and the longest HNT and the strongest networks of HNT and interphase produced the lowest percolation onset and the biggest/strongest networks providing the toughest samples.

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