A theoretical modelling framework was proposed to predict tensile moduli and tensile strengths of bioepoxy/clay nanocomposites in terms of clay content and epoxidised soybean oil (ESO) content, which could be influenced by properties of blended matrices in nanocomposites, clay filler type, orientation and dispersion status, clay morphological structures, and filler-matrix interfacial bonding. The random orientation of dispersed clay fillers played a significant role in predicting elastic moduli of bioepoxy/clay nanocomposites at clay contents of 1-8 wt% (ESO content: 20 wt%) according to Hui-Shia (H-S) laminate model and Halpin-Tsai (H-T) laminate model. In addition, when clay content was fixed at 5 wt%, H-S laminate model coincided well with the experimental data of bioepoxy/clay nanocomposites at the ESO contents of 0-40 wt%. Whereas, Hirsch model showed closer estimated values with experimental data at the ESO content of 60 wt%. Finally, Turcsányi-Pukánky-Tüdős (T-P-T) model predicted better tensile strengths of bioepoxy/clay nanocomposites at clay contents of 1-5 wt% (ESO content: 20 wt%) and at an ESO content of 20-60 wt% (clay content: 5 wt%).

1. Introduction

The depletion of fossil fuel stock and plastic waste is a critical issue in recent years with urgent actions required to utilise other alternative resources from natural polymers that can substitute for derived petroleum-based polymers [1, 2]. Researchers have shown enormous interests in bioepoxies because of their similar triglyceride epoxy side groups equivalent to conventional epoxy resins [3–6]. Bioepoxy polymers, such as epoxidised plant oils, are produced from derived plant oils and usually employed as lubricants and plasticisers for polyvinyl chloride (PVC) [3, 5, 7–13]. Epoxidised plant oils based on soybean and linseed are commercially available in large volumes with a reasonable price. Among these, epoxidised soybean oil (ESO) has been widely investigated in the manufacture of polymer composites due to its abundance, inexpensiveness, and high reactivity as comonomers when combined with other monomer-based plant oils to form crosslinking networks [11, 14, 15]. With respect to mechanical properties, the substitution of petroleum-based epoxy resin with ESO exhibits an improvement of fracture toughness and impact strength, not mentioning the decreases in tensile modulus and tensile strength of bioepoxy blends. Several studies have investigated the effect of additional ESO to undermine mechanical properties of epoxy resins such as their tensile strengths and tensile moduli [16, 17], flexural strengths and flexural moduli [17], and hardness [8] for ESO/diglycidyl ether of bisphenol A (DGEBA) blends, glass transition temperature ($T_g$) and storage modulus of epoxidised canola oil (ECO)/DGEBA blends [5], and thermal properties of epoxidised vegetable oil (EVO)/DGEBA blends [18]. To overcome those drawbacks mentioned earlier, the incorporation of rigid nanofillers such as nanoclay can enhance mechanical and thermal properties of bioepoxy resins and further extend the widespread applications of these nanocomposites [19–21]. High surface areas of well-dispersed and platelet-like clay nanostructures are believed to improve mechanical
properties of bioepoxy nanocomposites, as evidenced by their higher tensile strengths and tensile moduli to a certain extent when compared with those of neat biopolymers [22–25].

Mechanical properties of bionanocomposites reinforced with nanoclays are well-understood to be affected by the compatibility and reactivity of coupling agents including biopolymers, their catalysts, and nanoclay fillers, as well as associated manufacturing methods and processing parameters [24–30]. The good compatibility between polymer matrices and their catalysts enables to improve the crosslinking density of resulting nanocomposites. Moreover, the compatibility between matrices and nanoclays can also influence their adhesion or interfacial bonding [31]. Obviously, the modification of clay interlayers also further improves clay compatibility by means of increasing interfacial interactions between matrices and clay interlayers. Indeed, the tensile strengths of nanocomposites strongly depend on the good interfacial interaction between nanoclays and polymer matrices in order to achieve effective stress transfer from nanofillers to matrices accordingly [32–39].

On the other hand, clay wettability within epoxy matrices is also critical to achieve desirable mechanical properties of epoxy/clay nanocomposites, which are mainly attributed to homogeneous clay dispersion and favourable clay orientation [40, 41]. Other factors to influence mechanical properties of nanocomposites consist of clay content, clay aspect ratio, clay stiffness, and the dispersion and orientation of clay nanofillers [42–44]. Miyagawa et al. [45] investigated the effect of dispersion structures of organo-montmorillonite (MMT) nanoclays on elastic modulus, impact strength, and fracture properties of epoxidised linseed oil (ELO)/epoxy nanocomposites and ESO/epoxy nanocomposites. The addition of 5 wt% organoclays with exfoliated structures was found to improve the storage modulus by approximately 30%. On the contrary, intercalated nanocomposites reinforced with 5 wt% organoclays only yielded 8 and 29% increases in fracture toughness when compared with those exfoliated nanocomposites and neat epoxy, respectively.

In this study, the correlation between theoretical models based on composite theory and experimental data obtained was holistically evaluated in order to successfully predict tensile moduli and tensile strengths of bioepoxy/clay nanocomposites based on ESO by considering several factors such as clay content, clay orientation, clay aspect ratio, and ESO content in bioepoxy matrices. It is anticipated to eventually offer an appropriate guidance to the manufacture of bioepoxy/clay nanocomposites in order to replace the “trial and error” method used for the valid estimation of nanocomposite properties.

2. Theoretical Models of Polymer/Clay Nanocomposites

Modelling approaches employed for polymer nanocomposite systems generally consist of three categories of molecular-scaled, microscaled, and meso-/macroscaled methods according to different size effects [46]. Molecular-scaled methods focus on molecular dynamics, Monte Carlo method, and molecular mechanics for atoms, molecules, or clusters of the units. On the other hand, microscaled methods tend to link up the gap between the molecular-scaled method and meso-/macroscaled methods. It is well known that composite constituents are not only affected on the molecular-scaled level, but also at the microscopic scale to investigate the evolution of structures, bulk flow of materials, and the bonding interaction between matrices and fillers. Various methods such as Brownian dynamics, dissipative particle dynamics, lattice Boltzmann method, the time-dependent Ginsburg-Landau theory, and dynamic density functional theories have been developed to study microscopic structures and the interaction of composite constituents [46]. In particular, the continuum method has been considered as the most popular one among the meso-/macroscaled methods under the combination condition of both molecular structures and material nature with the homogenisation at different scaled levels. Such a method concentrates on the deformation of composite materials influenced by externally applied loads and their resulting stresses and strains. Macroscaled methods follow essential laws for the continuity of mass; the equilibrium of force, energy, and momentum; and conservation of energy and conservation of entropy [46]. Modelling and numerical simulations have been used to evaluate continuum methods including micromechanical models, equivalent-continuum models, self-consistent models, and finite element analysis. In this study, we only discussed the application of micromechanical models in predicting tensile properties of bioepoxy/clay nanocomposites in terms of their tensile moduli and tensile strengths based on the effect of individual nanocomposite constituents. Six theoretical models, namely, Voigt and Reuss models, Hirsch model, Halpin-Tsai (H-T) model, modified H-T models, Hui-Shia (H-S) model, and laminate model, were applied to predict tensile moduli of bionanocomposites. On the other hand, the estimation of their tensile strengths was based on Danusso-Tieghi (D-T) model, Nicolais-Nicodemo (N-N) model, Lu model, and Turcsányi-Pukánszky-Tődős (T-P-T) model. All modelling results were compared with corresponding experimental data to validate the feasibility of relevant theoretical models used as an essential guidance for nanocomposite manufacturing.

2.1. Modulus of Polymer Particulate Composites

2.1.1. Rule of Mixture (ROM). Rule of mixture (ROM) comprises a fundamental concept for micromechanical modelling in polymer particulate composite systems. ROM is based on the assumption that both fillers and polymer matrices are linearly elastic with the symmetry of fillers including filler shape, size and aspect ratio, and good interfacial bonding existing between fillers and matrices [46]. Voigt and Reuss models are deemed as the upper and lower bonds of ROM, respectively, in the theoretical prediction of composite systems. Voigt model involves the combination of overall composite constituents influenced by the average elastic modulus and volume fraction of each constituent, namely, parallel model [47]. Furthermore, it is also assumed that an isostrain condition takes place for both fillers and matrices.
in this upper bound, resulting in the modulus of composites given by

\[ E_c = E_p\phi_p + E_m(1 - \phi_p). \]  

(1)

In contrast, Reuss model or series model is regarded as the inverse ROM or lower bound of ROM. It is based on the assumption that uniform stress occurs in both matrices and fillers [48]. Reuss model can be written in the following equations:

\[ E_c = \frac{E_pE_m}{E_m\phi_p + E_p(1 - \phi_p)}, \]  

(2)

with

\[ \phi_p = \frac{W_p}{W_p + \left(\frac{\rho_f}{\rho_m}\right)(1 - W_p)}, \]  

(3)

\[ \rho_{blend} = \rho_1\phi_1 + \rho_2\phi_2, \]  

(4)

where \( \phi_p \) is the volume fraction of fillers in composites, \( W_p \) is the weight fraction of fillers, and \( \rho_m \) and \( \rho_p \) are densities of matrices and fillers, respectively, which can be obtained from the technical data sheet provided by the material supplier (Table 1). For the density of matrix blends \( (\rho_{blend}) \), it is calculated according to Equation (4) that is influenced by the densities \( (\rho_1 \) and \( \rho_2 \)) and corresponding volume fractions \( (\phi_1 \) and \( \phi_2 \)) of two constituents in blend matrices. In addition, \( E_c, E_m, \) and \( E_p \) are the moduli of composites, matrices, and fillers, respectively. In this study, \( E_m \) of bioepoxy blends was determined based on the experimental results obtained from tensile tests of bioepoxy blends at different ESO contents.

On the other hand, elastic modulus of fillers \( (E_p) \) can vary according to different clay dispersion statuses within bioepoxy matrices in nanocomposite systems. For exfoliated clay structures, \( E_p \) is given by 178 GPa, which is the modulus of the single clay platelet layer [54]. In addition, with respect to intercalated clay structures, the estimation of effective modulus of clay fillers would be different and can be calculated by applying ROM [54, 55] as follows:

\[ E_{\text{intercalation}} = E_{\text{MMT}}\phi_{\text{MMT}} + E_{\text{gallery}}(1 - \phi_{\text{MMT}}), \]  

if \( E_{\text{gallery}} \ll E_{\text{MMT}}, \)

\[ E_{\text{intercalation}} \approx E_{\text{MMT}}\phi_{\text{MMT}} \approx \left(\frac{d_{\text{MMT}}}{d_{002}}\right)E_{\text{MMT}}, \]  

(6)

where \( \phi_{\text{MMT}} \) and \( E_{\text{MMT}} \) are the volume fraction in MMT stacks and elastic modulus of MMT, respectively, while \( E_{\text{gallery}} \) is the gallery (interlayer) modulus. When the gallery modulus is much smaller than MMT modulus \( (E_{\text{gallery}} \ll E_{\text{MMT}}) \), \( E_{\text{gallery}} \) would make an insignificant contribution to the modulus of intercalated clay structures. Accordingly, the modulus of intercalated clay structures can be approximated to be in the expression of Equation (6) in terms of \( d_{002} \)-spacing obtained from XRD results. Based on XRD and TEM results, clay structures have been formed with the combination of dominantly intercalated, exfoliated, and aggregated clay structures in our prepared bioepoxy/clay nanocomposites [56]. The interlayer spacing \( d_{002} \) of clay fillers was selected demonstrating a shifting interlayer spacing of clay platelet layers. As such, effective moduli of intercalated clay fillers at different filler contents and ESO contents were determined, as listed in Table 2.

2.1.2. Modified Rule of Mixture (MROM). In general, ROM is limited to the effect of elastic moduli and volume fractions of composite constituents while filler shape, filler orientation, and 3D spatial position of fillers are usually not considered. In fact, the latter more or less undermines elastic moduli of composite materials, which can be compensated for with the incorporation of modulus reduction factor (MRF) in MROM [57]. This is particularly the case in using the MRF along with filler aspect ratio \((\alpha)\) [58] when imperfect filler orientation in composite materials is taken into account. As such, MROM can be given by

\[ E_c = E_p\phi_p(MRF) + E_m(1 - \phi_p). \]  

(7)

The determination of MRF can be based on two different forms for flake-like fillers in composites, namely, Riley’s rule [59] and Padawer and Beecher’s rule [60] as follows:

Riley’s rule:

\[ (\text{MRF}) = 1 - \frac{\ln (u + 1)}{u}, \]  

(8)

Padawer and Beecher’s rule:

\[ (\text{MRF}) = \tanh \frac{u}{\alpha}, \]  

(9)

with

\[ u = \frac{1}{2} \sqrt{\frac{\phi_p G_m}{E_p(1 - \phi_p)}}, \]  

(10)

in which \( G_m \) is the shear modulus of the matrix and \( \alpha \) is the aspect ratio of fillers in composite systems. Wu et al. [58] investigated the MRF of rubber/clay nanocomposites by fitting corresponding experimental data. It was found that MRF = 0.66 could better fit experimental data for the modulus prediction when the filler volume fraction was less than 6 vol%.

2.1.3. Hirsch Model. Hirsch [61] proposed a model for estimating the elastic modulus of composites with the combination of parallel and series models. The general form of such
Table 1: Material properties of epoxy resin, ESO, and Cloisite 15 clays in bionanocomposites.

<table>
<thead>
<tr>
<th>Material</th>
<th>Properties</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>DGEBA</td>
<td>Epoxy equivalent weight (EEW): 190-195 g/eq</td>
<td>[49, 50]</td>
</tr>
<tr>
<td></td>
<td>Molecular weight: 340.42 g/mol</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Density: 1.17 g/cm³</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Viscosity: ~11.000 mPa-s</td>
<td></td>
</tr>
<tr>
<td>ESO</td>
<td>Oxirane content: 6.2%</td>
<td>[51, 52]</td>
</tr>
<tr>
<td></td>
<td>Molecular weight: 975.40 g/mol</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Density: 0.982 g/cm³</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Viscosity: 325 mPa-s</td>
<td></td>
</tr>
<tr>
<td>Cloisite 15</td>
<td>Organo-modified MMT clay</td>
<td>[53]</td>
</tr>
<tr>
<td></td>
<td>Bis(hydrogenated tallow alkyl) dimethyl, salt</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$d_{001} = 3.63$ nm</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cation exchange capacity (CEC) = 125 meq/100 g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Density $\rho = 1.66$ g/cm³</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$E_{MMT} = 178$ GPa for single MMT platelet</td>
<td>[54]</td>
</tr>
</tbody>
</table>

Table 2: Effective modulus of intercalated clay particles in bionanocomposites (ESO content: 20 wt%).

<table>
<thead>
<tr>
<th>Clay filler content (wt%)</th>
<th>Bionanocomposites with 20 wt% ESO</th>
<th>$d_{001}$ (nm) [56]</th>
<th>$E_{\text{intercal}}$ (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>1.97</td>
<td>85.06</td>
</tr>
<tr>
<td>3</td>
<td></td>
<td>1.96</td>
<td>85.59</td>
</tr>
<tr>
<td>5</td>
<td></td>
<td>1.96</td>
<td>85.59</td>
</tr>
<tr>
<td>8</td>
<td></td>
<td>1.96</td>
<td>85.59</td>
</tr>
</tbody>
</table>

where $E_c$, $E_m$, and $E_p$ represent the elastic moduli of composites, matrices, and fillers, respectively, and $\eta_L$ is given by

$$\eta_L = \frac{E_p/E_m}{1 - \eta_L}$$

An equation is the effect of matrix modulus, the modulus of fillers, and their volume fractions along with an empirical constant $x$. Hirsch model equation is given by

$$E_c = x(E_m(1 - \varnothing_p) + E_p\varnothing_p) + (1 - x)\frac{E_pE_m}{E_m\varnothing_p + E_p(1 - \varnothing_p)},$$

(11)

where $x$ is an empirical constant to control the stress transfer between fillers and matrices in composites, which is based on the curve fitting with experimental data ($0 < x < 1$). “$x$” is regarded as a significant parameter in predicting elastic moduli of nanocomposites, which can also determine the filler orientation of nanocomposites.

2.1.4. Halpin-Tsai Model. Halpin-Tsai (H-T) model [62, 63] offers a reasonable prediction for elastic moduli of unidirectional composites with respect to volume fraction and geometry of fillers. H-T model is generally used for continuous or discontinuous fillers such as fibre-like or flake-like fillers with different alignment directions. The elastic modulus of composites can be expressed in the following form:

$$E_c = E_m \frac{1 + \xi \eta_L \varnothing_p}{1 - \eta_L \varnothing_p},$$

(12)

where $\xi$ is a constant depending on the geometry and aspect ratios of fillers in composites, $\xi$ can be given in the following forms:

$$\xi = 2a = 2\left(\frac{t}{l}\right)$$

for longitudinal modulus ($E_{11}$),

(14)

$$\xi = 2$$

for transverse modulus ($E_{22}$),

(15)

in which $l$ and $t$ are the length and thickness/depth of dispersed fillers in composites, respectively. In addition, when $\xi$ is very large ($\xi \to \infty$), H-T model is equal to ROM (i.e., a parallel model), as given in Equation (1). Contrarily, when $\xi$ is very small ($\xi \to 0$), H-T model becomes an inverse model of ROM (i.e., series model) presented in Equation (2). Thus, H-T model can be used for most prediction situations between lower and upper limits in ROM.

On the other hand, in polymer/clay nanocomposite system, H-T model can be used to predict elastic moduli of nanocomposites influenced by volume fraction and aspect ratio of clay fillers as well as clay dispersion status in nanocomposites [54, 55, 64]. In completely exfoliated clay structures, the prediction of elastic modulus of nanocomposites would be calculated using Equation (11) with $E_p/E_m$, being equal to the ratio of the modulus of single clay platelet layer to matrix modulus. In addition, for an intercalated clay structure system, elastic modulus prediction is also determined according to Equation (11), but $E_p/E_m$ can change and become the ratio of elastic modulus of platelet stacks to that of matrices. The elastic modulus of intercalated clay
structures can be calculated based on Equation (6). On the contrary, for randomly oriented clay platelets, the elastic modulus of nanocomposites can be calculated using the combination of longitudinal modulus \((E_{11})\) and transverse modulus \((E_{22})\) of nanocomposites with the same aspect ratio and volume fraction of clay fillers, as given below [64, 65]:

\[
E_c = E_m \left[ \frac{3}{8} E_{11} + \frac{5}{8} E_{22} \right]
\]

(16)

or

\[
E_c = E_m \left[ \frac{3 \left( 1 + \xi \eta_L \phi_p \right)}{8 \left( 1 - \eta_L \phi_p \right)} + \frac{5 \left( 1 + 2 \eta_T \phi_p \right)}{8 \left( 1 - \eta_T \phi_p \right)} \right]
\]

(17)

with

\[
\eta_T = \left( \frac{E_p}{E_m} \right) - 1 \left( \frac{E_p}{E_m} \right) + 2,
\]

(18)

\[
\xi = \frac{2}{3} \alpha = \left( \frac{2\ell_p}{l} \right).
\]

(19)

Additionally, 2D structure shape of clay platelets can possibly make less contribution to the modulus of nanocomposites as compared with those filled by 1D fibre-like counterparts. Thus, the MRF for clay platelet fillers is incorporated in modified H-T model according to Equations (12) and (16), which are presented below accordingly:

\[
E_c = E_m \frac{1 + \xi (\text{MRF}) \eta_L \phi_p}{1 - \eta_L \phi_p},
\]

(20)

\[
E_c = E_m \left[ \frac{3 \left( 1 + \xi (\text{MRF}) \eta_L \phi_p \right)}{8 \left( 1 - \eta_L \phi_p \right)} + \frac{5 \left( 1 + 2 \text{MRF} \eta_T \phi_p \right)}{8 \left( 1 - \eta_T \phi_p \right)} \right].
\]

(21)

2.1.5. Hui-Shia (H-S) Model. Hui-Shia (H-S) model develops the prediction of elastic modulus of unidirectionally aligned two-phase composites with fibre-like or flake-like filler inclusions [57, 66]. H-S model simplifies the orientation of fillers based on the assumption of perfect interfacial bonding between matrices and fillers with similar Poisson’s ratios. The associated elastic modulus equations are given below:

Longitudinal elastic modulus \((E_{11})\):

\[
\frac{E_c}{E_m} = \frac{E_{11}}{E_m} = \frac{1}{1 - \left( \phi_p / 4 \right) \left[ \left( 1 / \xi' \right) + 3 \left( \xi' + \Lambda \right) \right]},
\]

(22)

Transverse elastic modulus \((E_{22})\):

\[
\frac{E_c}{E_m} = \frac{E_{22}}{E_m} = \frac{1}{1 - \left( \phi_p / \xi' \right)},
\]

(23)

with

\[
\xi' = \phi_p + \frac{E_m}{E_p - E_m} \left( 1 - \phi_p \right) \left( 1 - g \right) \alpha'^2 - \left( \frac{g}{2} \right) \alpha'^2 - 1,
\]

(24)

\[
\Lambda = \left( 1 - \phi_p \right) \left[ 3 \left( \alpha'^2 + 0.25 \right) g - 2 \alpha'^2 \right],
\]

(25)

where \(\alpha'\) is the inverse of aspect ratio \(\alpha\) or the ratio of thickness to the length of fillers (i.e., \(\alpha' \approx t / l\)), and \(g\) is a geometric parameter of composites defined in Equations (23), (24), and (25). In case of \(\alpha' = 1\), \(g = 2 / 3\), \(\alpha' \geq 1\) belongs to the inclusion of fibre-like fillers while \(\alpha' \leq 1\) is used for the inclusion of flake-like or disk-like fillers according to Equation (26). In addition, for a perfect interface, \(g\) would be given in Equation (27).

\[
g = \frac{\pi}{2} \alpha'.
\]

(27)

2.1.6. Laminate Model. Laminate model is developed with respect to geometric properties of filler inclusions in matrices. From the experimental point of view, typical fillers in composites are generally subjected to random orientation or misalignment, as compared with unidirectionally aligned reinforcements. The aspect ratio and modulus of fillers, as well as the orientation of dispersed fillers have a significant effect on elastic moduli of composites [54]. In case of fully random orientation in the 3D orthogonal direction, platelet-like fillers would induce more effective planar reinforcements as opposed to fibre-like fillers. The related equations based on fibre-like or platelet-like fillers are proposed as follows [67, 68]:

\[
E_{\text{fiber}}^{\text{ran-3D}} = 0.184E_{11} + 0.816E_{22},
\]

(28)

\[
E_{\text{platelet}}^{\text{ran-3D}} = 0.49E_{11} + 0.51E_{22},
\]

(29)

where \(E_{11}\) and \(E_{22}\) are longitudinal and transverse moduli of composites, respectively.
2.2. Strength of Polymer Particulate Composites

2.2.1. Danusso-Tieghi (D-T) Model. Danusso and Tieghi [69] proposed a relationship of mechanical strength and volume fraction of rigid matrix-based composites. It is assumed in Danusso and Tieghi (D-T) model that there is no adhesion between matrices and fillers, resulting from no load transfer from matrices to fillers. Thus, the total load is equal to that carried by matrices alone. In addition, D-T model can be applied from regular or irregular filler distribution to random filler dispersion. Nonetheless, this model can also be used with the existing adhesion. A simple equation for the strength of composites is given by:

\[
\sigma_c = \sigma_m(1 - \psi),
\]

where \(\sigma_c\) and \(\sigma_m\) are tensile strengths of composites and matrices, respectively, and \(\psi\) is the area fraction in the cross section. With those aforementioned model assumptions and random orientation of filler structures, the area fraction of the matrix must be the same for any cross section through the matrix, for which the area fraction of the matrix’s cross section is equal to the volume fraction of fillers (\(\psi \approx \phi_p\)) [69, 70].

2.2.2. Nicolais and Narkis (N-N) Model. This model is based on previous D-T model by replacing the volume fraction with a power law function in terms of volume fraction as follows [71]:

\[
\sigma_c = \sigma_m\left(1 - a\phi_p^b\right),
\]

where \(a\) and \(b\) are constants influenced by particle shape and arrangement in composites. Based on the hypothesis that there is no adhesion between matrices and fillers, Nicolais and Narkis (N-N) model is considered for cubic filler shape with uniformly dispersed spherical particles. However, it is assumed that a minimum cross section of continuous phase should be perpendicular to the applied load. Thus, the strength of composites is calculated according to the equation below:

\[
\sigma_c = \sigma_m\left(1 - 1.21\phi_p^{2/3}\right).
\]

It is used to predict the lower-bound strength of composites. In addition, for their upper bound, the strength is calculated with the assumption of perfect adhesion between matrices and fillers. As such, the strength of composites is the same as that of the matrix or unfilled polymer (\(\sigma_c = \sigma_m\)).

2.2.3. Lu Model. Lu model [72] has been developed by modifying N-N model when there is some adhesion existing between matrices and fillers, which is given by:

\[
\sigma_c = \sigma_m\left(1 - 1.07\phi_p^{2/3}\right).
\]

This equation is established by using the combined properties from micromechanical measurement, microdamage monitoring, and micromechanical analysis in order to obtain the interfacial bonding strength between matrices and fillers.

2.2.4. Turcsányi-Pukánszky-Tüdös (T-P-T) Model. Turcsányi et al. [73] proposed a semiempirical equation for very strong particle-matrix interfacial bonding with a simple hyperbolic function to describe the change of filler cross section with the filler content, which is known as the Turcsányi-Pukánszky-Tüdös (T-P-T) model [74–76]. Nonetheless, the functionalisation of exponential function is subjected to the unexpected dependence of the matrix strength and filler volume fraction. Additionally, T-P-T model can be applied to composites with the inclusion of spherical particles and anisotropic particles. The equation is generally formulated as follows:

\[
\sigma_c = \left[1 - \phi_p - \sigma_m \frac{1}{1 + 2.5\phi_p - \sigma_m} \exp\left(B\phi_p\right)\right],
\]

where \(\sigma_m\) and \(\phi_p\) are the strength of matrices and volume fraction of fillers, respectively. \(B\) is a constant that depends on the surface area of particles, their density, and interfacial bonding with matrices. If \(B = 0\), the fillers act as voids having poor interfacial bonding without adhesion and load transfer at the matrix-filler interface. Nonetheless, if \(B \leq 3\), the filler-matrix interface is weak without reinforcing effect. The constant \(B\) was calculated by fitting experimental data with theoretical results and derived from the minimum sum of squares of variance from experimental data of composite strengths.

3. Result and Discussion

3.1. Tensile Modulus of Bionanocomposites

3.1.1. Effect of Clay Content. Theoretical models according to Equations (1) to (29) were used to predict the effect of clay content on tensile moduli of bio epoxy/clay nanocomposites cured by isophorone diamine (IPDA) determined in our previous study [56]. Material properties of bionanocomposite constituents including the moduli of single platelet MMT (\(E_{MMT}\)), DGEBA as a conventional epoxy resin, and ESO as well as the molecular weight and density of matrices (i.e., DGEBA resin and ESO) and Cloisite 15 clay nanofillers are listed in Table 1. In addition, Equation (6) was used to calculate the effective elastic modulus of intercalated clay structures detected in bionanocomposites. The \(d\)-spacing values of intercalated clay structures were obtained according to previous XRD results [56]. The calculation results for effective elastic moduli of intercalated clay structures in bionanocomposites are listed in Table 2, which was used to estimate theoretical elastic moduli of bionanocomposites at different clay contents. Additionally, with respect to the random orientation of clay structures, 3D platelet-like filler laminate model was employed to incorporate actual filler geometries. Nonetheless, MRF = 0.66 [58] was also implemented based on a mix of partially intercalated/exfoliated and aggregated clay structures in bionanocomposites by assumption. The abovementioned geometric parameters including clay
orientation and dispersion status were determined according to previous results [56] obtained from TEM micrographs of bionanocomposites cured by IPDA at a fixed ESO content of 20 wt%, as shown in Figure 1.

Theoretical prediction of elastic moduli of bioepoxy/clay nanocomposites as a function of clay content was compared with experimental data, as illustrated in Figures 2–4. In Figure 2, it can be seen that Voigt model demonstrated an overestimated result as opposed to experimental data despite the inclusion of MRF. On the contrary, the theoretical prediction of elastic modulus by Reuss model, which is known as inverse Voigt model, presented an underestimation for experimental data instead. Voigt and Reuss models only give a rough estimation for elastic moduli of bionanocomposites, thus demonstrating their upper and lower bounds, respectively. This phenomenon indicated that those two models neglected the geometry, dispersion status, and orientation of fillers, which further made the theoretical prediction become questionable. In order to enhance the prediction accuracy, filler geometric parameters, filler dispersion status, and relevant orientation were taken into account by utilising Hirsch model and H-T model. It is assumed in Hirsch model that the random orientation of clay fillers takes place with the combination of parallel and series filler orientations in bionanocomposites. The curve fitting with experimental data by Hirsch model revealed better agreement with experimental data when compared with those predicted by Voigt and Reuss models. Nonetheless, it has been found that $x = 0.029$ for parallel filler orientation to the stress direction. This can be indicated that the series orientation of clay platelet fillers is significantly affected by the behaviour of bionanocomposites in real experiments.

On the other hand, in order to enhance the accuracy of geometric parameters, H-T model and modified H-T model as well as modified equations with the incorporation of MRF were employed in predicting elastic moduli of bionanocomposites at the ESO content of 20 wt% by varying aspect ratios of fillers according to Equations (13) to (21) and (29), respectively. Figure 3 presented the theoretical prediction and experimental data for elastic moduli of bionanocomposites according to H-T model and modified H-T laminate model. Figure 3(a) showed that increasing the aspect ratio, from 3 to 60, improved the estimated elastic moduli in line with increasing the clay content owing to the enhanced wettability and contact surface areas between clay fillers and matrices. The lower clay content yields higher filler aspect ratios as expected. Nonetheless, the use of MRF in H-T laminate model with random orientation exhibited a reduction in predicted elastic moduli, which could eventually affect the prediction of aspect ratios for clay fillers. The percentage difference between experimental data and theoretical results was ranging from 2 to 8.7% and 0.7–2.7% for H-T laminate model and H-T laminate model with MRF, respectively. Similar approaches were also revealed for the theoretical prediction of elastic moduli of nanocomposites based on modified H-T model with the random filler orientation according to Equation (16), as shown in Figure 3(b). The predicted elastic moduli of bionanocomposites were in fairly close agreement with experimental data. The percentage errors for modified H-T model with the random filler orientation were in the range of 0.7-7.6%. On the other hand, for those models with MRF, the percentage errors were found to be between 0.9 and 2.5%. Detailed percentage errors of each theoretical model are summarised in Table 3. Consequently, the prediction of elastic modulus by H-T laminate model and modified H-T random model with MRF appeared to be much closer to experimental data when compared with those without MRF.

Additionally, as seen in Figure 4, elastic moduli of bionanocomposites were also predicted by H-S laminate model in terms of aspect ratio of clay fillers and clay content. Similarly, with the implementation of H-T model and modified H-T model, theoretically predicted elastic moduli of bionanocomposites exhibited different aspect ratios with increasing clay content. The increase in clay content revealed the reduction of aspect ratios for clay fillers in bionanocomposites. Nonetheless, the percentage errors between theoretical results and experimental data were ranging between 0.07 and 3.5%, indicating excellent consistency with experimental data. It is implied that H-S laminate model offers better theoretical prediction for elastic moduli of bionanocomposites as opposed to other theoretical models used such as ROM model, Hirsch model, H-T laminate model and modified H-T random model.

Clay aspect ratios presented special geometric characteristics of clay fillers in bionanocomposites. When clay platelet layers are dispersed well in matrices with the existence of single platelet layers, it leads to exfoliated structures with an aspect ratio as high as 1000 [77]. However, with respect to undispersed clay fillers in matrices, their aspect ratios can be significantly reduced to approximately 10 or even less. The theoretical prediction of elastic modulus by H-T model and H-S model could obtain estimated aspect ratios of clay fillers in bionanocomposite systems at different clay contents, as shown in Table 4. It was found that at a lower clay content of 1 wt%, the effective aspect ratio reached the highest value in the range of 30 to 60 despite being still lower than that of single clay platelet at 100-1000. When the clay content increased up to 3 and 5 wt%, effective aspect ratios decreased to a range of 15-30 and 10-15, respectively. In addition, experiment data at the highest clay content of 8 wt% revealed the lowest aspect ratio ($\alpha \leq 5$), possibly implying poor clay dispersion in nanocomposites. In relation to effective aspect ratios of clay particles, it was revealed that a closer aspect ratio prediction at clay contents of 3-8 wt% was manifested in H-T laminate model and modified H-T random model with MRF. Similarly, a better aspect ratio prediction was also detected in both H-T models without MRF at the same clay contents. However, at the lower clay content of 1 wt%, various aspect ratios were estimated for different H-T models, which could be attributed to different assumptions used for determining geometric parameters to predict elastic moduli of nanocomposites using H-T laminate model and modified H-T random model.

On the other hand, H-S laminated model exhibited a higher aspect ratio prediction with more sensitivity when compared with various H-T models according to Table 4. Despite various aspect ratios, the prediction of effective
aspect ratios is consistent with morphological structures of bionanocomposites depicted in Figure 1, which is evident in the presence of the random orientation of clay fillers in matrices and the increasing level of clay aggregation with increasing the clay content. More interestingly, elastic modulus predictions obtained using H-S laminate model were closer to experimental data as opposed to those obtained from H-T models, especially at clay contents of 5-8 wt%.
shown in Table 3. The percentage errors of elastic modulus prediction for bionanocomposites at clay contents of 5 and 8 wt% by using H-S laminate model were found to be approximately 1.84 and 0.07%, respectively. Nonetheless, those predicted by modified H-T random model were about 2.46 and 0.85% accordingly. In contrast, at lower clay contents of 1 and 3 wt%, the percentage errors in elastic modulus prediction by H-S laminate model were higher than those using H-T models (i.e., approximately 1.6 and 3.48%, respectively). As such, it is implied that H-S laminate random model is more applicable to predict elastic moduli of bionanocomposites beyond the clay content of 5 wt%. On the other hand, H-T laminate model and modified H-T random model with/without MRF were confirmed to be more feasible for nanocomposites at the clay content below 3 wt%.

3.1.2. Effect of ESO Content. Similar to the effect of clay content, theoretical predictions to elastic moduli of bionanocomposites were compared with experimental data using Equations (1) to (29). The estimated results of those predictions were depicted in Figures 5–8. At different ESO contents from 0 to 60 wt%, elastic moduli of bioepoxy blends varied according to the ESO content, as shown in Table 5. In a similar manner, the densities of blend matrices were also altered based on different ESO contents. Additionally, the dispersion

![Figure 3: Theoretical prediction curves for elastic moduli of bionanocomposites with intercalated and randomly oriented clay fillers: (a) H-T laminate model and (b) modified H-T model with random filler orientation.](image-url)
The status of clay fillers in nanocomposites affected elastic moduli of bionanocomposites. Hence, elastic moduli of clay particles were calculated based on Equation (6) along with the results listed in Table 5. Such elastic moduli of clay particles were calculated with the assumption that intercalated clay structures were dominantly formed as compared to other dispersion statuses (i.e., exfoliated and aggregated clay structures), which were based on XRD analysis results in the previous work [56] along with other parameters used for theoretical models in Table 5. The clay content of 5 wt% in nanocomposites was fixed when using all different ESO contents in bionanocomposites.

In Figure 5, elastic moduli of nanocomposites show a declining trend with increasing the ESO content. As previously mentioned, the decrease in elastic modulus could result from the ESO plasticisation effect [56]. Increasing the ESO content yielded bionanocomposites with more rubber-like behaviour and less brittle nature in material properties. Voigt model and the Reuss model could only provide rough predictions for the elastic moduli of bionanocomposites at different ESO contents, as depicted in Figure 5. It was seen that theoretical prediction of Reuss model being the lower bound gave better estimation than Voigt model as the upper bound. This result suggested that series model for fibre-matrix interaction became more dominant in good accordance with Reuss model mentioned earlier.

On the other hand, calculated percentage errors for each theoretical model were listed in Table 6, which were obtained from the percentage of the difference between experimental data and theoretical modelling results. As clearly seen, theoretical predictions using Reuss model and Hirsch model

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**Table 3: Percentage errors in theoretical models in comparison with experimental data at different clay contents in bionanocomposites.**

<table>
<thead>
<tr>
<th>Theoretical model</th>
<th>Clay content (wt%)</th>
<th>1</th>
<th>3</th>
<th>5</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voigt</td>
<td>79.54</td>
<td>259.82</td>
<td>428.63</td>
<td>809.72</td>
<td></td>
</tr>
<tr>
<td>Voigt (MRF)</td>
<td>46.25</td>
<td>163.54</td>
<td>273.55</td>
<td>527.57</td>
<td></td>
</tr>
<tr>
<td>Reuss</td>
<td>-17.25</td>
<td>-20.11</td>
<td>-22.19</td>
<td>-10.41</td>
<td></td>
</tr>
<tr>
<td>Hirsch</td>
<td>-14.47</td>
<td>-12.09</td>
<td>-9.27</td>
<td>13.10</td>
<td></td>
</tr>
<tr>
<td>H-T laminate random</td>
<td>2.75</td>
<td>1.95</td>
<td>3.27</td>
<td>8.66</td>
<td></td>
</tr>
<tr>
<td>H-T laminate random with MRF</td>
<td>-1.98</td>
<td>-0.74</td>
<td>2.73</td>
<td>2.28</td>
<td></td>
</tr>
<tr>
<td>Modified H-T random</td>
<td>0.71</td>
<td>-2.40</td>
<td>6.59</td>
<td>5.85</td>
<td></td>
</tr>
<tr>
<td>Modified H-T random with MRF</td>
<td>-0.91</td>
<td>-2.03</td>
<td>-2.46</td>
<td>0.85</td>
<td></td>
</tr>
<tr>
<td>H-S laminate</td>
<td>-1.60</td>
<td>-3.48</td>
<td>-1.84</td>
<td>-0.07</td>
<td></td>
</tr>
</tbody>
</table>

**Table 4: Aspect ratios of clay fillers based on curve fitting between experimental data and theoretical modelling results in bionanocomposites.**

<table>
<thead>
<tr>
<th>Theoretical model</th>
<th>Aspect ratio</th>
<th>Clay content (wt%)</th>
<th>1</th>
<th>3</th>
<th>5</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>H-T laminate</td>
<td>~40</td>
<td>10-15</td>
<td>5-10</td>
<td>&lt;3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H-T laminate with MRF</td>
<td>~60</td>
<td>~20</td>
<td>10-15</td>
<td>&lt;3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Modified H-T random</td>
<td>~30</td>
<td>10-15</td>
<td>5-10</td>
<td>&lt;3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Modified H-T random with MRF</td>
<td>~40</td>
<td>15-20</td>
<td>10-15</td>
<td>~3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H-S laminate</td>
<td>~60</td>
<td>20-30</td>
<td>~15</td>
<td>~5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: Negative values in percentage errors represent lower prediction values than experimental data.
became much closer to experimental data as compared with Voigt model and Voigt (MRF) model. For instance, without ESO (i.e., 0 wt% ESO), comparable percentage errors of approximately 5.30 and 5.89% in Reuss model and Hirsch model were reported, respectively, as opposed to Voigt model with about 100% in percentage error. More interestingly, theoretical predictions of elastic moduli of bionanocomposites with the addition of 60 wt% ESO were shown to be in good agreement with experimental data using Hirsch model. The percentage error of proposed Hirsch model at the ESO content of 60 wt% was approximately 7%, which was the closest prediction when compared with other proposed theoretical models listed in Table 6.

With respect to the random orientation of clay fillers in matrices, 3D laminate model was employed to predict elastic moduli of bionanocomposites at different ESO contents, along with H-T model and H-S model according to Equations (14) to (29). Nonetheless, since it was found that clay structures were the combination of partial intercalation/exfoliation and particle aggregation, the MRF of 0.66 was used in H-T model equations. The associated curves showing the elastic modulus prediction by H-T laminate model and modified H-T random model with MRF were depicted in Figures 6 and 7. It was seen that the modification of H-T model affected the prediction of filler aspect ratios, especially at the ESO contents of 20 and 40 wt%, even though the percentage errors in modified H-T random model with MRF were comparable to those in H-T laminate model with MRF shown in Table 6. Additionally, it was found that filler aspect ratios of bionanocomposites varied according to different ESO contents exhibited in Figure 7. Nonetheless, the use of MRF showed an increasing aspect ratio as compared with that without MRF. The aspect ratios of conventional nanocomposites (i.e., 0 wt% ESO) exhibited comparable values when predicted using H-T laminate model and modified H-T random model with MRF resulting in aspect ratios of fillers less than 3. This could suggest that clay fillers were undispersed in conventional epoxy, which was likely to form aggregated clay fillers in matrices. However, a further increase in ESO content revealed increasing aspect ratio values. At the ESO content of 20 wt%, the aspect ratio predicted by H-T laminate model with MRF was improved in range of 5-10. Whereas, the additional ESO at approximately 40 wt% yielded the aspect ratios of 10-15. In addition, with respect to modified H-T random model with MRF, its predicted aspect ratios of bionanocomposites at 20 and 40 wt% were slightly higher than those based on H-T laminate model with MRF, leading to aspect ratios of 10-15 and approximately 20, respectively. Finally, bionanocomposites with 60 wt% ESO exhibited an aspect ratio over 20 for both H-T model and modified H-T model. Nonetheless, the percentage errors in each H-T model were observed to be similar.

Figure 5: Comparison between theoretical modelling results and experimental data for elastic moduli of bionanocomposites as a function of ESO content.
which were found to be ~99% less than those obtained from corresponding experimental data. This unpredicted elastic modulus could be attributed to particle-size effect in relation to filler aspect ratios, as well as particle content [54, 78]. Since aspect ratio values represented particle sizes of clays dispersed in bioepoxy matrices, the reduction in clay particle sizes would increase the aspect ratios of clay fillers. Similarly, with the effect of filler inclusion, the increase in the filler aspect ratio would decrease the size of dispersed clay fillers in matrices as expected, which might further improve the rigidity of nanocomposites. More interestingly, the high elastic modulus of bionanocomposites at 85.08 GPa in experimental data with 60 wt% ESO was likely to be associated with a far higher modulus of clay particles as opposed to that of corresponding bioepoxy (i.e., 10^-4 GPa). The high elastic modulus of clay fillers could improve the stiffness of bionanocomposites. A direct effect arising from the clay inclusion into rubber-like bioepoxy matrices would significantly influence the deformation reduction of bionanocomposites as opposed to that of neat bioepoxy at high ESO contents.

In addition to the effect of aspect ratio and modulus of fillers, the orientation of dispersed clay fillers also significantly affected elastic modulus, which was indicative of the

![Diagram](image-url)
percentage of the difference between theoretical prediction calculated according to Equation (29) for H-T laminate random model and Equation (16) for modified H-T random model and experimental data. From an experimental point of view, modified H-T random model with lower percentage errors is more applicable to predict elastic moduli of bionanocomposites when compared with H-T laminate random model. This can be associated with different interpretations in determining the constant values of \([^E_{11}\text{]}\), which are 0.375 and 0.49 GPa for modified H-T random model and H-T laminate model, respectively. Furthermore, it is very critical to predict filler aspect ratios in matrices based on the theoretical prediction of elastic moduli of nanocomposites with the combination of random orientation of fillers in the matrices.

On the other hand, with respect to H-S laminate model, the theoretical prediction of bionanocomposites in comparison with experimental data was shown in Figure 8. Clearly, the curve fitting by H-S laminate model demonstrated good prediction for filler aspect ratios in experiment data. The aspect ratios predicted by H-S laminate model were summarised in Table 7. Increasing the ESO content dramatically enhanced the aspect ratios of clay fillers in bionanocomposites. Conventional nanocomposites (i.e., 0 wt% ESO) presented the lowest aspect ratios below 3. The aspect ratios of

Figure 7: Comparison between theoretical modelling results and experimental data for elastic moduli of bionanocomposites at different aspect ratios calculated based on (a) modified H-T random model and (b) modified H-T random model with MRF.
bionanocomposites with 20 wt% ESO were in the range of 15-20, as opposed to nearly 30 for bionanocomposites at the ESO content of 40 wt%. When the ESO content was 60 wt%, the predicted filler aspect ratio reached a level higher than 30, as illustrated in Figure 8. Additionally, the effect of clay content also revealed that the theoretical prediction of elastic moduli of bionanocomposites determined by H-S laminate model led to closer prediction values given in Table 6. The percentage errors for bionanocomposites up to 40 wt% ESO were below 7%, which appeared to be greater up to almost 100% at the ESO content of 60 wt%. In contrast to other theoretical models, the application of H-S laminate model yielded much better elastic modulus prediction with lower percentage errors, especially for bionanocomposites at the ESO contents of 20 and 40 wt% (i.e., ~1.84 and 1.16%, respectively). Moreover, filler aspect ratios predicted by H-S laminate model remained at a high level as opposed to those estimated by other theoretical models. As a result, when the effect of ESO content on bionanocomposites reinforced by 5 wt% clays was investigated, it was evident that H-S laminate model appeared to be more reliable in predicting their elastic moduli against experimental data. This was especially the case for bionanocomposites at the ESO contents below 40 wt%. On the other hand, when the ESO content was as high as 60 wt%, Hirsch model with the combination of parallel and series orientations became more applicable in estimating elastic moduli of bionanocomposites.

### 3.2. Tensile Strength of Nanocomposites

#### 3.2.1. Effect of Clay Fillers

The theoretical tensile strength values predicted by Equations (30)–(34) were compared with experimental data of bionanocomposites cured by IPDA at a fixed ESO content of 20 wt%. Figure 9 showed the theoretical prediction of nanocomposites by using different theoretical models to fit experimental data. Tensile strengths of bionanocomposites demonstrated an initially enhancing trend when increasing the clay content from 1 to 5 wt% along with the further reduction at the higher clay content of 8 wt%, as illustrated in Figure 9. Four different theoretical models were also employed to predict tensile strength values for bionanocomposites. N-N model and T-P-T model \((B=0)\) underestimated the experimental data in relation to tensile strengths of bionanocomposites, implying that bionanocomposites possessed better interfacial adhesion between clay fillers and matrices, thus resulting in an increase in tensile strength with the inclusion of clay fillers. The further analysis for experimental data of bionanocomposites was based on the adoption of D-T model and Lu model in comparison with experimental data. As shown in Figure 9, it was observed that a comparable correlation between theoretical modelling results and experimental data predicted by D-T model was

<table>
<thead>
<tr>
<th>ESO content (wt%)</th>
<th>(E_{\text{blend}}) (GPa)</th>
<th>(\sigma_{\text{blend}}) (MPa)</th>
<th>(\rho_{\text{blend}}) (g·cm(^{-3}))</th>
<th>Imaginary modulus (d_{\text{Ip}}) [56] (nm)</th>
<th>Interlaminar elastic modulus (E_{\text{intercalation}}) (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>3.30</td>
<td>59.56</td>
<td>1.17</td>
<td>1.80</td>
<td>92.72</td>
</tr>
<tr>
<td>20</td>
<td>0.49</td>
<td>28.53</td>
<td>1.13</td>
<td>1.96</td>
<td>85.59</td>
</tr>
<tr>
<td>40</td>
<td>0.13</td>
<td>10.26</td>
<td>1.09</td>
<td>2.07</td>
<td>80.65</td>
</tr>
<tr>
<td>60</td>
<td>1x10(^{-4})</td>
<td>0.29</td>
<td>1.06</td>
<td>1.97</td>
<td>85.08</td>
</tr>
</tbody>
</table>

![Figure 8](image_url): Elastic modulus predictions of bionanocomposites reinforced with 5 wt% clays as a function of ESO content at different filler aspect ratios using H-S laminate model.

![Table 5](image_url): Tensile properties of blend matrices and effective moduli of intercalated clay particles in bionanocomposites (fixed clay content: 5 wt%).
achieved as opposed to Lu model at the high clay content of 8 wt%.

On the other hand, T-P-T model was also employed to be compared with experimental data for bionanocomposites, and the curve fitting based on T-P-T model yielded $B = 4.718$. It was suggested that bionanocomposites could benefit from the reinforcing effect by increasing the clay filler content along with better matrix-filler interfacial bonding when $B$ value is over 3. Additionally, due to the high difference between theoretical prediction and experimental data in bionanocomposites at the clay content of 8 wt%, the theoretical prediction obtained from T-P-T model was in better accordance with experimental data as a function of ESO content, Figure 9. It was determined that the empirical constant of $B = 8.693$, which indicated that bionanocomposites at clay contents from 1 to 5 wt%, exhibited better interfacial bonding resulting in more effective filler-matrix load transfer than that at 8 wt%. In addition, bionanocomposites at the clay content of 8 wt% presented a lower tensile strength value as compared with that predicted by T-P-T model with $B = 3$. Furthermore, those results suggested that such bionanocomposites at the clay content of 8 wt% might arise from the weak interfacial bonding between clay fillers and matrices without reinforcing effect [73–76]. Nonetheless, such a prediction had good agreement with SEM analysis results, as indicated in our previous study [56] where microcracks existed close to interfacial areas between clay fillers and matrices.

### Table 6: Percentage errors in theoretical predictions in comparison with experimental data based on different theoretical models at various ESO contents in bionanocomposites.

<table>
<thead>
<tr>
<th>Theoretical model</th>
<th>ESO content (wt%)</th>
<th>0</th>
<th>20</th>
<th>40</th>
<th>60</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voigt</td>
<td>100.08</td>
<td>428.63</td>
<td>1249.97</td>
<td>17121.94</td>
<td></td>
</tr>
<tr>
<td>Voigt (MRF)</td>
<td>65.38</td>
<td>273.55</td>
<td>811.35</td>
<td>11266.69</td>
<td></td>
</tr>
<tr>
<td>Reuss</td>
<td>5.30</td>
<td>-22.19</td>
<td>-35.87</td>
<td>-99.35</td>
<td></td>
</tr>
<tr>
<td>Hirsch</td>
<td>5.89</td>
<td>-9.27</td>
<td>-27.91</td>
<td>7.20</td>
<td></td>
</tr>
<tr>
<td>H-T laminate</td>
<td>22.48</td>
<td>3.27</td>
<td>5.64</td>
<td>-98.52</td>
<td></td>
</tr>
<tr>
<td>H-T laminate with MRF</td>
<td>17.39</td>
<td>2.73</td>
<td>5.24</td>
<td>-98.80</td>
<td></td>
</tr>
<tr>
<td>Modified H-T random</td>
<td>16.04</td>
<td>6.59</td>
<td>-1.20</td>
<td>-98.90</td>
<td></td>
</tr>
<tr>
<td>Modified H-T random with MRF</td>
<td>12.38</td>
<td>-2.46</td>
<td>-5.35</td>
<td>-99.06</td>
<td></td>
</tr>
<tr>
<td>H-S laminate random</td>
<td>7.79</td>
<td>-1.84</td>
<td>-1.16</td>
<td>-99.01</td>
<td></td>
</tr>
</tbody>
</table>

Note: Negative values in percentage errors represent lower prediction values than experimental data.

### Table 7: Aspect ratios of clay fillers based on curve fitting of experimental data with theoretical models at different ESO contents in bionanocomposites.

<table>
<thead>
<tr>
<th>Theoretical model</th>
<th>ESO content (wt%)</th>
<th>Aspect ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>H-T laminate</td>
<td>&lt;3 3-5 10 20</td>
<td></td>
</tr>
<tr>
<td>H-T laminate with MRF</td>
<td>&lt;3 3-5 10 15 20</td>
<td></td>
</tr>
<tr>
<td>Modified H-T random</td>
<td>&lt;3 3-5 10 15 20</td>
<td></td>
</tr>
<tr>
<td>Modified H-T random with MRF</td>
<td>&lt;3 10-15 20 30</td>
<td></td>
</tr>
<tr>
<td>H-S laminate random</td>
<td>&lt;3 15-20 30 30</td>
<td></td>
</tr>
</tbody>
</table>

3.2.2. Effect of ESO Content. Similar to the effect of clay content, the prediction for tensile strengths of bionanocomposites at a fixed clay content of 5 wt% was compared with experimental data as a function of ESO content, Figure 10. It was clearly seen that experimental data for tensile strengths of bionanocomposites were reduced when increasing the ESO content from 20 to 60 wt% despite the strength improvement as opposed to that of neat matrices. In addition, tensile strengths of conventional nanocomposites (i.e., 0 wt% ESO) exhibited a declining tendency with the inclusion of 5 wt% clay fillers.

With respect to the theoretical prediction, as can be seen in Figure 10, all proposed models failed to present good predictions to fit all experimental data of tensile strengths. The percentage errors of proposed model prediction for tensile strength were enhanced with increasing the ESO content, as displayed in Table 8. The proposed models could thus only fit one or two experimental data. From the hypothesis of particle-matrix interfacial interaction, N-N model and T-P-T (at $B = 0$) model were in better accordance with experimental data for conventional nanocomposites (0 wt% ESO). Percentage errors in N-N model and T-P-T model ($B = 0$) exhibited comparable values of approximately 0.27 and 2.17%, respectively, as opposed to experimental data (i.e., 0 wt% ESO). Since N-N model and T-P-T model ($B = 0$) were categorised as lower bounds for modelling tensile strength, it could be suggested that conventional nanocomposites (i.e., 0 wt% ESO) had poor filler-matrix interfacial bonding. However, other theoretical models such as D-T model and Lu model presented an Overestimation of tensile strengths in contrast with experimental data. The percentage error was found to be over 10% for those two models according to Table 8.

On the other hand, with respect to T-P-T model, the curve fitting with experimental data offered an empirical constant $B = 1.273$. However, the prediction curve failed to fit well with the experimental data at the ESO contents from 0 to 60 wt%. Furthermore, in the further analysis, experimental data were only used from 20 to 60 wt% in the subsequent model prediction as bionanocomposites with 0 wt% ESO presented a comparable prediction using T-P-T model with an empirical constant $B = 0$. Further prediction calculations...
based on three proposed experimental data were shown in Figure 10. The curve fitting with experimental data (20 to 60 wt% ESO) revealed better agreement with T-P-T model results along with the resulting value of $B = 7.960$. From this prediction using T-P-T model ($B = 7.960$), the percentage differences between experimental data and theoretical prediction were 0.67 and 4.63% for tensile strengths of bionanocomposites with the inclusions of 20 and 40 wt% ESO, respectively. Moreover, T-P-T model ($B = 7.960$) revealed a closer correlation with experimental data as opposed to other proposed strength prediction models according to the percentage errors listed in Table 8.

On the other hand, prediction models for tensile strengths of bionanocomposites with 60 wt% ESO presented a high percentage of the difference when compared with experimental data (Table 8). As such, it was suggested that the high improvement in tensile strength of bionanocomposites (i.e., 60 wt% ESO) could be attributed to structural factors due to different morphological structures and interfacial bonding interaction between fillers and matrices. Turcsányi et al. [73] has noted that structural factors, which could influence stress concentration, are the shape and the size of dispersed fillers, spatial distribution of matrices, and interface thickness. In practice, as previously mentioned in

![Figure 9: Comparison between theoretical modelling results and experimental data of tensile strengths of bionanocomposites at different clay contents.](image1)

![Figure 10: Tensile strength predictions of bionanocomposites reinforced with fixed 5 wt% clays as a function of ESO content using different theoretical models.](image2)
the theoretical predictions of elastic moduli of bionanocomposites, increasing the ESO content yielded higher aspect ratios of clay fillers in matrices. Furthermore, the improvement of aspect ratios can also be associated with the reduction of filler size in matrices, resulting in a much higher increase in clay surface areas. As such, high clay surface areas could cause better matrix-filler interaction, which offered positive impact on the strength improvement with more efficient stress transfer from fillers to matrices.

4. Conclusions

A comparison between experimental data and theoretical predictions with respect to tensile properties of bioepoxy/clay nanocomposites has been demonstrated successfully. Most proposed theoretical models for tensile modulus predictions revealed good agreement with experimental data except Voigt model and Reuss model. Hirsch model with the combination of series and parallel filler orientations revealed reasonable prediction for constant parameter \( x \) where \( x = 0.029 \) in the parallel direction. Among various models, H-T laminate model and modified H-T random model with MRF, as well as H-S laminate model, revealed very good correlation with experimental data when the percentage errors appeared to be below 3.5%. Nonetheless, it was also clearly noted that aspect ratios of clay fillers in bionanocomposites decreased with increasing the clay filler content.

Theoretical predictions of tensile modulus as a function of ESO content demonstrated partial correlation with experimental data for most proposed models except Voigt model. Similar to the effect of clay content, Voigt model overestimated experimental data owing to the assumption of filler alignment in the longitudinal direction. In case of conventional nanocomposites (i.e., 0 wt% ESO), lower-bound model predictions (i.e., Reuss model and Hirsch model with \( x = 0.006 \)) and H-S laminate model fitted well with experimental data (error percentage < 10%). However, at the ESO contents of 20–40 wt%, H-T laminate model, modified H-T random model, and H-S laminate model demonstrated very good agreement with experimental data resulting in the percentage errors being less than 5%. In addition, as for bionanocomposites with 60 wt% ESO, most proposed theoretical models underestimated experimental data except Hirsch model. A good agreement between experimental data and Hirsch model was observed when \( x = 0.006 \) in the parallel filler orientation. This suggested that the series orientation of fillers was more dominant in predicting elastic moduli of bionanocomposites at the ESO content of 60 wt% (error percentage: approximately 7%). Additionally, according to H-T model and H-S model, aspect ratios of clay fillers in nanocomposites were found to increase with increasing the ESO content.

In addition, with respect to tensile strength as a function of clay content, D-T model, Lu model, and T-P-T model at \( B = 3 \) had good partial agreement with experimental data of bionanocomposites with 5 wt% clay inclusion. In particular, the curve fitting of experimental data at clay contents from 0 to 5 wt% was reasonably good when T-P-T model was employed with empirical constant \( B = 8.693 \) due to strong filler-matrix interfacial bonding.

When the effect of ESO content on tensile strengths of bionanocomposites was considered, selected models except D-T model and Lu model showed partially good fitting with experimental data. N-N model and T-P-T model (\( B = 0 \)) fully fitted only experimental data of conventional nanocomposites (i.e., 0 wt% ESO). In contrast, T-P-T model matched better experimental data at the ESO contents of 20 and 40 wt% on account of strong filler-matrix interfacial bonding (\( B = 7.960 \)). In addition, experimental data with 60 wt% ESO did not fit all proposed models with a clear underestimation instead of the comparable fitted model because aspect ratios of fillers to enhance stress transfer between fillers and matrices were not involved in proposed theoretical models.

Data Availability

All data generated or analysed during this study are included in this manuscript.

Conflicts of Interest

The authors declare that they have no conflicts of interest and no financial interest related to this study.

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