Mathematical modelling of microtubule-tau protein transients: insights into the superior mechanical behavior of axon

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ABSTRACT
Axons with staggered microtubules cross-linked by tau protein possess a remarkable mechanical balance of high specific stiffness and toughness. Owing to their viscoelastic nature, axons exhibit stress rate-dependent mechanical behavior, which is relevant to their selective vulnerability to damage in traumatic brain injury. A Kelvin-Voigt viscoelastic shear lag model is developed to elucidate the mechanical responses of axons under transient tensile force. Analytical closed-form expressions are derived to characterize the relative sliding, stress transfer and failure mechanism between microtubule and tau protein while the axon is stretched transiently. The results from the theoretical solutions elucidate how the MT-tau interface length and stress rate affect the mechanical responses of axon. It is found that axonal failure mechanism may be different under different loading conditions. Long microtubules are more vulnerable to rupture at high stress rate, yet short microtubules are likely to detach from microtubule bundles under large deformations. In the view of multi-level failure of axon, it is illustrated how the vulnerable axons protect themselves from overall damage, and how the axon can simultaneously achieve an outstanding mechanical balance of high specific stiffness and toughness.

KEYWORDS: Staggered composite; Biocomposite; Shear lag model; Kelvin-Voigt viscoelastic model; Traumatic brain injury

1. Introduction
Staggered biocomposites are found to exhibit a remarkable mechanical balance of fracture toughness versus stiffness and strength [1, 2]. In consideration of the structure of staggered biocomposite and the vulnerability of axon, it may possess the innate ability of mechanical balance of high stiffness and high toughness, and self-protecting over the course of evolution; hence the axon is taken as an example for investigation in this study.

Traumatic brain injury (TBI) is primarily resulted from an impact to the brain tissue from an external mechanical force including contact force, penetration of a projectile, or inertia forces induced by rapid
acceleration and deceleration [3]. In recent years, TBI has become a major public health issue, killing 15 per 100,000 individuals in European countries [4] and affecting over 2.5 million people in the USA [5] per year.

Plenty of *in vitro* researches indicated that the mechanical rupture of microtubules (MTs) results in traumatic brain injury, commonly referred to as “concussion” [6, 7]. As depicted in Fig. 1a, a single long slender axon and multiple shorter dendrites are responsible for transmitting and receiving signals, respectively [8]. Histologically, axonal cytoskeleton consists of MT bundles connected by microtubule-associated protein (MAP) tau, as illustrated in Fig. 1b [8, 9]. Parallel array of MTs, acting as reinforcing components with a main function of transporting electrochemical cargo, are the stiffest structural element within the axon [10-13].

Owing to their viscoelastic nature, axons exhibit strain or stress rate-dependent rupture [7]. In order to characterize the micromechanical breaking of axonal ultrastructure, axons were uniaxially stretched transiently by using transmission electron microscopy (TEM) under different strain rates [6, 7]. Experimental techniques are tricky to capture the immediate mechanical behavior of axonal MT bundles in tension. It is reported that the axon behaved like an elastic element while it is subjected to sudden changes in force [14]. Indeed, viscoelasticity may be more appropriate to capture the major biophysical behavior of axon and associated MTs in such conditions.

In recent years, there is an increasing interest in modelling the mechanical response of axon. However, little is known about which component of the axonal cytoskeleton might break under transient mechanical loading and large deformation. Tolomeo et al. [15] reported that cross-linking proteins provided shear resistance between MTs. Peter et al. [16] used a discrete bead-spring model to simulate the biomechanical behavior of axonal microtubule bundle under uniaxial tension, and they assumed that both MT and tau protein are linear elastic materials. Shamloo et al. [17] considered MTs as a large number of discrete masses and modelled tau protein as Kelvin-Voigt element so that viscoelastic model can be employed to simulate the transient response of axonal MTs under sudden forces. de Rooij et al. modelled the time-dependent behaviour of brain tissue by adopting an analytical hyperelastic approach [18], and developed a computational mechanics model to elucidate cellular-level characteristics of MT bundles [19]. Combined with experimental results and finite element analysis (FEA), viscosity has been identified as the most critical factor causing the mechanical vulnerability of axon under transient loading [10].

Within the axonal cytoskeleton, tau protein plays a vital role in the assembly of individual MTs [20], and the breaking of tau protein may be relevant to some of neurodegenerative diseases, such as Alzheimer’s disease [21, 22] and Parkinsonism [23]. Different viscoelastic models consisting of linear elastic MT and viscoelastic tau protein, a combination of elastic springs and viscous dashpots, were postulated to elucidate the macroscopic viscoelastic behavior of the axon by some researchers [14, 24, 25]. Ahmadzadeh et al. [10] verified that Kelvin viscoelastic model can efficiently characterize the viscoelastic mechanical behavior of axon, but only MT failure is investigated in their model without consideration of tau protein breaking. However, once tau protein axial strain exceeds its ultimate strain, MTs will detach from the MT bundle [26, 27]. What’s more, tau protein breaking is found to be associated with some of neurodegenerative diseases. Therefore, it’s significant to develop a
computational model comprehensively characterizing the axonal failure evidence of MT rupture and tau protein breaking.

Though many researches revealed the mechanical behavior of tau protein and MTs [10, 14, 24, 25], the molecular mechanism of axonal failure remains poorly understood. **In Ref. [28] we have investigated** MT-tau interface failure mechanism under transient torsional loading. Our goals of this study is to start a dialog between experimentalists and modelers, so that experimentalists are more informed about what type of data is needed for model development and validation. At the microscopic length scale of axon, randomly staggered alignment of discontinuous MTs are held together by connective tau proteins. The objective of this work is to investigate the mechanical response of axons under transient tensile force and the fundamental design principles of biocomposites for engineering fiber reinforce composites. To explore the mechanical responses of axons, shear lag model (SLM) including viscoelastic tau proteins is developed to predict the relative sliding and stress transfer in the axon under transient loading. In order to accomplish these research objectives, three research questions are to be answered:

1. The structure of vulnerable axon is mainly composed of parallel staggered reinforcing MTs cross-linked by tau protein. What kind of superior mechanical properties does the axon have?
2. How does the vulnerable axon protect itself from overall damage over the course of evolution?
3. How does the axon simultaneously achieve an outstanding mechanical balance of high specific stiffness and toughness?

**Fig. 1.** (a) Neuron and cross-section diagram of the axon, reprinted from [19]; (b) Electron microscopy image of axonal MTs cross-linked by tau proteins (Arrows); (c) Cross-sectional electron micrograph of MT bundle, adapted from [29].

2. Materials and methods

2.1. Fundamental formulas and theoretical derivation
In 1952, SLM was firstly proposed by Cox [30], which has now been modified and developed for biomechanical applications [31-38]. SLM can be employed to characterize stress transfer between MTs and tau protein under transient tensile conditions. As shown in Fig. 2a, the cytoskeleton of MT is hollow cylindrical structures with outer radius \( R_o \) and inner radius \( R_i \). As the elastic modulus of MTs is much larger than that of tau protein, MTs are assumed to follow linear elastic constitutive relation, and its Young’s modulus is denoted by \( E_M \).

Before theoretical derivations, the assumptions are adopted in the analysis presented in this study as follows:

(i) The stiffest discontinuous MTs are connected by tau protein, leading to deformation and deformation rate primarily occur in tau protein, thus, all the MTs are assumed to be linear elastic materials, and tau protein follows viscoelastic constitutive law.

(ii) The MT-tau interface length \( L \) is over two orders of magnitude larger than the distance between adjacent MTs so that the deformation in axon is essentially along x-direction.

(iii) All the MTs are parallel array along x-direction, and the MT-tau interface length \( L \) is nearly half of their entire length (see Fig. 2a).

\[
\eta \dot{\delta} + \delta = \frac{F}{K}
\]

where \( \eta=\mu/K \) is tau protein dashpot timescale. A Cartesian coordinate system is established at the midpoint of an entire length of MT so that the longitudinal direction of MTs is along x-axis (see Fig.
2c). In the existing study of axon [10], the axon is simplified as a staggered arrangement of discontinuous MTs connected by tau proteins, which is represented by a periodic unit cell that is half the entire length of MT. When the unit cell is stretched by a tensile force, stress will be transferred from one MT to the adjacent MTs through tau protein, which leads to relative sliding between MTs. Tau protein elongation $\delta$ and the relative sliding between adjacent MTs $[u_1(x,t) - u_2(x,t)]$ follow the geometrical relationship as follows

$$\delta(x,t) \approx [u_1(x,t) - u_2(x,t)] \cos \beta$$

where $\beta$ is the angle between the tau protein and the MT; $u_1$ and $u_2$ are displacements of upper and lower MTs, respectively.

Tensile forces $F(x,t)$ resulting from the elongation of tau protein can be converted to shear stress $\tau(x,t)$ acting over the entire MT circumference, which can be written as

$$\tau(x,t) = \frac{\alpha F(x,t) \cos \beta}{2\pi R_d d_t}$$

where $d_t$ is the center-to-center distance between neighboring tau protein. The number of adjacent MTs is $\alpha=6$ based on the cross-sectional electron micrograph of MT bundle (Fig. 1c and Fig. 2b).

According to the unit cell (see Fig. 2c) showing the two adjacent MTs, shear stress $\tau(x,t)$ acts over the MT circumference leading to the following equilibrium equations

$$\frac{\partial \sigma_1(x,t)}{\partial x} = \frac{2R_0}{R_0^2 - R_i^2} \tau(x,t)$$

$$-\frac{\partial \sigma_2(x,t)}{\partial x} = \frac{2R_0}{R_0^2 - R_i^2} \tau(x,t)$$

where $R_0$ and $R_i$ are the outer radius and inner radius of the MTs, respectively. $\sigma_1$ and $\sigma_2$ are the normal stresses of the linear elastic upper and lower MTs, which can be expressed as

$$\sigma_1(x,t) = E_M \varepsilon_1 = E_M \frac{\delta u_1(x,t)}{\delta x}, \quad \sigma_2(x,t) = E_M \varepsilon_2 = E_M \frac{\delta u_2(x,t)}{\delta x}$$

where $E_M$ is the Young’s modulus of MTs. $\varepsilon_1$ and $\varepsilon_2$ are the axial strain of the upper and lower MTs, respectively.

While MT has a negligible mass, at any point $x$ along MTs longitude direction, the tensile force $P(t)$ should maintain mechanical balance with the normal stresses acting on the MT cross-section, thus we can obtain

$$\sigma_1(x,t)S + \sigma_2(x,t)S = P(t)$$

where $S=\pi(R_0^2-R_i^2)$ is the cross-sectional area of MT.

Letting

$$\varepsilon_1^2 = \eta, \quad L_e^2 = \frac{E_M d_t S}{2\alpha K \cos^2 \beta}$$

and combining Eqs. (1)-(8), we can get the governing equation for tau protein elongation

$$\frac{\partial \delta}{\partial t} = L_e^2 \frac{\partial^2 \delta}{\partial x^2} - \frac{1}{\varepsilon_1^2} \frac{\partial \delta}{\partial x}$$

The upper MT normal stress $\sigma_1$ along the $x$-direction can be written as
\[ \sigma_i = \frac{E_m}{2\cos \beta} \frac{\partial \delta}{\partial x} + \frac{P}{2S} \] (10)

with boundary conditions \( \sigma_i(0,t) = 0, \quad \sigma_i(L,t) = P(t)/\pi(R_o^2 - R_i^2) \). Based on Eq. (10), boundary and initial conditions are alternatively expressed as

\[ \frac{\partial \delta(0,t)}{\partial x} = -\frac{P(t)\cos \beta}{E_m S}, \quad \frac{\partial \delta(L,t)}{\partial x} = -\frac{P(t)\cos \beta}{E_m S} \] (11)

\[ \delta(x,0) = \varphi(x) \] (12)

where \( L_C \) is the characteristic length over which the stress is transferred between MT and tau protein.

Tau protein elongation can be derived from solving the nonhomogeneous partial differential equation (Eq. (9)) with nonhomogeneous boundary conditions (Eq. (11)) and initial condition (Eq. (12))

\[ \delta(x,t) = [C_o + D_o(t)] e^{\frac{1}{2}t} + \sum_{n=1}^\infty [C_n + D_n(t)] e^{\left(\frac{1}{2} - \frac{\xi_n^2}{L^2}\right)t} \cos \frac{n\pi x}{L} \] (13)

where

\[ C_o = \frac{1}{L} \int_0^L \left[ \varphi(x) + \frac{E_m S}{E_m SL} \frac{P(0)\cos \beta}{x^2} + \frac{P(0)\cos \beta}{E_m S} x \right] dx \]

\[ C_n = \frac{2}{L} \int_0^L \left[ \varphi(x) - \frac{E_m S}{E_m SL} \frac{P(0)\cos \beta}{x^2} + \frac{P(0)\cos \beta}{E_m S} x \right] \cos \frac{n\pi x}{L} dx \]

\[ D_o(t) = \int_0^t e^{\frac{1}{2}(t-s)} h_0(s) ds \]

\[ D_n(t) = \int_0^t e^{\frac{1}{2}(t-s)} h_n(s) ds \]

\[ h_0(t) = \frac{1}{L} \int_0^L H(x,t) dx \]

\[ h_n(t) = \frac{2}{L} \int_0^L H(x,t) \cos \frac{n\pi x}{L} dx \]

\[ H(x,t) = \frac{\cos \beta}{\zeta E_m SL} \left[ 2L_o^2P(t) - P(t)x^2 + P(t)Lx - \zeta^2 P'(t)x^2 + \zeta^2 P'(t)Lx \right] \]

It can be seen from the above equations that, the values of \( C_0 \) and \( C_n \) depend on the initial condition. If there are no elongation and no force in the beginning, we can get \( C_0 = C_n = 0 \). \( D_0 \) and \( D_n \) are determined by the force \( P(t) \) stretched at the midpoint of the entire MT.

After the expression of tau protein elongation is gained, the solutions for normal stress in the MT and shear stress acting over the MT circumference yield

\[ \sigma_i(x,t) = \sigma_i(x-L,t) = -\frac{n\pi}{2L} \sum_{n=1}^\infty \left[ C_n + D_n(t) \right] e^{\left(\frac{1}{2} - \frac{\xi_n^2}{L^2}\right)t} \sin \frac{n\pi x}{L} + \frac{2P(t)\cos \beta}{SL} x - \frac{P(t)\cos \beta}{2S} \] (14)

\[ \tau(x,t) = -\frac{n^2\pi^2}{4R_i L^2} \sum_{n=1}^\infty \left[ C_n + D_n(t) \right] e^{\left(\frac{1}{2} - \frac{\xi_n^2}{L^2}\right)t} \cos \frac{n\pi x}{L} + \frac{P(t)\cos \beta}{SL} \] (15)

Obviously, MT axial strain can be derived from Eq. (6). Tau protein axial strain can be derived from the definition of axial strain \( \varepsilon = \delta \sin \beta/\delta x \), where \( \delta x \) is the surface-to-surface distance between adjacent
2.2. Failure criteria and universal applicability of the solutions

In general, the mechanical properties of strength and toughness are mutually exclusive [42]; yet these properties are a significant requirement for most structural materials. Strength is invariably a stress representing a material’s resistance to non-recoverable deformation [43]. When the transient tensile force continuously exerts at the end of the MT, the main component of either MT or tau protein will reach the ultimate strain. It was reported that MTs are shown to be ruptured under 50% of axial strain by several researchers [44-46]. With increasing strain, tau-tau bonds begin to break in exceed of approximately 40% [41]. Once MT axial strain reaches 50%, MT ruptures and transient loading comes to an end. Detachment of MT from bundle occurs at $x=0$ and $x=L$ once tau protein axial strain reaches >40%. Thus, three typical types of axonal failure may appear when MTs are increasingly stretched by a transient tensile force, as shown in Fig. 3. Fig. 3 records the axonal failure during the time (upper: MT sliding; left: MT rupture; middle: a portion of MT detachment; right: MT rupture; bottom: all the MT detachment). Generally, breaking of the bonds between the paired tau proteins starts from both ends and propagates sequentially toward the midpoint of the MT-tau interface. $L_s$ and $L_d$ denote the MTs sliding length and detachment length, respectively (see Fig. 3). Solutions in sliding region (Eqs. (13)-(15)) remain valid in case a) the initiation of tau protein elongation at both ends is considered as the initial condition $\phi(x)$; b) the Cartesian coordinate system moves the right with the distance of $L_d$; c) $L$ is replaced by $L_s$. As MTs are assumed to be linear elastic material, solutions in MTs detachment region can be easily obtained.

![Fig. 3. Three types of axonal failure.](image)

2.3. Related experiments and parameters

It can be noted from Eq. (9) that, the micromechanical interaction behavior of axon is determined by $(L_C/\zeta)^2$ and $1/\zeta^2$. These two parameters depends on the geometrical and material parameters of MT and tau protein, excluding the MT-tau interface length $L$. Nevertheless, boundary and initial conditions are determined by external force $P(t)$ and the MT-tau interface length $L$. In brief, the mechanical response of axon bundle depends on external force and geometrical and material properties of MT and tau.
Experimental data from numerous researches are applied to assign values to the parameters in viscoelastic SLM [9, 16, 17, 41, 44-49]. The length of MT is about 1-10 μm, and its outer and inner diameters are 25 nm and 14 nm, respectively [9]. Axonal MTs are hexagonally aligned with surface-to-surface spacing of approximately 20 nm [16, 17, 29]. In general, the Young’s modulus of MT could be captured by the slope of its stress-strain curve. The ultimate tensile strain of MT can be defined as the axial strain when the MT ruptures. The Young’s modulus of MTs was found to be 1.51 ± 0.19 GPa [9], which agreed well with experimental data reported by Suresh [50]. The rupture tensile strain can be set at 50% according to experimental measurements of the MT rupture strain performed by Janmey et al. [44], consistent with Suresh’s experimental observation [50].

The geometrical and material parameters of tau protein are also of great significance. About 95% of tau proteins are formed by paired helical filaments with a radius of 4-10 nm [47]. Based on statistical analysis, Hirokawa et al. [48] reported that tau protein center-to-center spacing fell within the typical range, 20-40 nm. The Young’s modulus of tau protein is typically about 5 MPa [16, 49], which gives the tau cross-link axial spring constant of 33.3 pN/nm as calculated from $K=E_t\pi R_t^2 \sin \beta / d_{st}$. According to the experimental data [41], Ahmadzadeh et al. [10] estimated the viscoelastic parameter $\eta=0.35$ s by using Bell’s equation [51]. Additionally, interactions in hTau40 broke at maximum force 300 pN observed by Wegmann et al. [41], and therefore tau protein breaking tensile strain is estimated in exceed of 40%.

In the current work, the geometrical and material parameters are gained originally from experiment data or estimations, as listed in Table 1. The angle $\beta$ between the tau protein and the MT is assumed to be 60°. According to these parameters, we can obtain the characteristic length from Eq. (8), $L_C=0.390$ μm.

**Table 1** Variables and values for viscoelastic SLM

| Variable | Value    | References |
|----------|----------|------------|
| $R_O$    | 12.5 nm  | [9]        |
| $R_I$    | 7 nm     | [9]        |
| $R_T$    | 4-10 nm  | [47]       |
| $E_M$    | 1.5 GPa  | [49]       |
| $d_{st}$ | 20-38 nm | [20]       |
| $d_T$    | 20-40 nm | [48]       |
| $2L$     | 1-10 μm  | [9]        |
| $K$      | 33.3 pN/nm | [16, 49] |
| $\eta$   | 0.35 s   | [10, 41]   |
| -        | 50%      | [44-46]    |
| -        | 40%      | [41]       |
3. Results

3.1. Analysis of axonal failure process

3.1.1. Prediction of the axon failure mode

Axonal MTs are experimentally found to be parallel arrays, with an average length of 4 μm [3]. Fig. 4 shows the maximum MT and tau protein axial strain history of MT-tau interface length about 2 μm. It can be noted from Fig. 4 that MT and tau protein reach their ultimate strain at the same time when $\dot{\sigma} = 92.5E_M \text{ Pa/s}$, which indicates that MT failure and tau protein failure occur simultaneously at applied stress rate $\dot{\sigma} = 92.5E_M \text{ Pa/s}$. When the applied stress rate is $100E_M \text{ Pa/s}$, MT reaches its ultimate strain firstly, thus axonal failure follows the S-R regime. Otherwise, it follows either S-D regime or S-D-R regime. Additionally, axon failure process under different stress rates is illustrated in Fig. 4b.

**Fig. 4.** (a) Prediction of the time sequence of MT failure and tau protein failure. The blue and orange lines capture the maximum MT and tau protein axial strains, respectively; and different linetypes characterize different stress rates. (b) Axon failure process under different stress rates. Model parameter used in calculation are $n=1-30$, $R_o=12.5 \text{ nm}$, $R_I=7 \text{ nm}$, $E_M=1.5 \text{ GPa}$, $R_T=7 \text{ nm}$, $d_M=20 \text{ nm}$, $d_T=30 \text{ nm}$, $\beta=60^\circ$, $\alpha=6$, $K=33.3 \text{ pN/nm}$, $\eta=0.35 \text{ s}$, $L = 5L_c = 1.9479 \mu\text{m}$, stress rate $\dot{\sigma} = 100E_M, 92.5E_M, 75E_M, 50E_M, 10E_M, 0.1E_M \text{ Pa/s}$.

3.1.2. Sliding-rupture regime

Fig. 5 shows the maximum axial strain of tau protein at both ends of the MT-tau interface length ($L$), and the minimum tau protein axial strain at the center, which agrees with the analytical results [17]. The maximum and minimum axial strains of MTs are found to be at the midpoint and both ends of the entire MT length ($2L$), respectively. Under the applied stress rate $\dot{\sigma} = 92.5E_M \text{ Pa/s}$, the initiation breaking of tau-tau bond and MT rupture occur at the same time; yet MTs come to failure first at the applied stress rate $\dot{\sigma} = 100E_M \text{ Pa/s}$, and MT-tau interface remains intact at all times. While MT is continuously stretched at a high stress rate, tau protein axial strain is extremely large at both ends ($x=0$ or $L$), but is nearly zero in the region $0.1 < x/L < 0.9$ (Fig. 5a and 5c). The axial strain distribution of the upper MT and lower MT surrounding the midpoint ($0.1 < x/L < 0.9$) remain constant, which equals to half of the maximum strain (Fig. 5b and 5d). It can be noted from Fig. 5b and 5d that, evidence of MT
rupture is more sensitive to stress rate rather than tensile stress, consistent with the experimental observation in the region of axonal swelling by Tang-Schomer et al. [7]. MTs are considered as the tracks for chemical cargo transport, and they will rupture under high stress rate ($\sigma \geq 92.5E_M$ Pa/s) found in TBI, leading to the transport interruption [6, 7]. This mathematical modelling can help us to understand the mechanism of TBI, and predict the symptom appearance of TBI when the brain is subjected to sudden external mechanical forces.

Fig. 5. Distribution of (a) tau protein axial strain along the MTs and (b) MT axial strain with $L=5L_c=1.9479$ μm, stress rate $\dot{\sigma}=92.5E_M$ Pa/s, MT rupture time $t_0=0.0054$ s; (c) tau protein axial strain along the MTs and (d) MT axial strain with $L=5L_c=1.9479$ μm, stress rate $\dot{\sigma}=100E_M$ Pa/s, MT rupture time $t_0=0.005$ s (In Fig. 5b and 5d, solid line and dashed line represent the upper MT and the lower MT plotted in Fig. 2c, respectively).

3.1.3. Sliding-detachment regime
The second type of axonal failure is caused by tau protein breaking, leading to MT detachment from MT bundle. Tau protein and MT axial strain are shown in Fig. 6. When applied stress rate are much smaller than 92.5$E_M$ Pa/s, e.g. $\dot{\sigma}=10E_M$ Pa/s or 0.1$E_M$ Pa/s, axonal failure follows S-D regime. It can be seen from Fig. 6 that, as the stretched loading continuously increases, the breaking of bonds between the paired tau proteins starts from both ends and propagates sequentially toward the midpoint of the MT-tau interface, as observed in the experiment [16]. According to the MT detachment time, MT-tau interface breaks faster and faster, as the applied loading increases continuously.
3.1.4. Sliding-detachment-rupture regime

A complicated failure process of axon, containing tau protein breaking and MT rupture, is illustrated in Fig. 7. When applied stress rate is slightly smaller than $92.5E_M$ Pa/s, e.g. $\dot{\sigma} = 50E_M$ Pa/s or $75E_M$ Pa/s, shear stress will be concentrated at both ends of the MT-tau interface even though the relative sliding is small. Obviously, tau protein closed to both ends breaks first, and then MTs rupture not until all the connected tau proteins come to failure, as shown in Fig. 7.

Both MT rupture and tau protein breaking will occur in different MT-tau interface lengths under transient loading conditions [3, 16]. However, to the best of our knowledge, there is no experimentally
reported data characterizing this complicated type of axonal failure—a portion of MT detachment and MT rupture successively happen to the same MT, although this type of axonal failure may happen in the light of our theoretical prediction, which is different from the S-R axonal failure regime.

Fig. 7. Distribution of (a) tau protein axial strain along the MTs and (b) MT axial strain with \( L=5L_c=1.9479 \, \mu m \), stress rate \( \dot{\sigma}=50E_M \, Pa/s \), tau-tau bonds breaking time \( \tau_0=8.1893 \, ms \), MT rupture time (end) at 10 ms; (c) tau protein axial strain along the MTs and (d) MT axial strain with \( L=5L_c=1.9479 \, \mu m \), stress rate \( \dot{\sigma}=75E_M \, Pa/s \), tau-tau bonds breaking time \( \tau_0=6.222 \, ms \), MT rupture time (end) at 6.667 ms.

### 3.2. Effect of the stress rate on axonal failure process

In this section, we investigate the strain distribution at different stress rates by comparing Figs. 5–7. While axon is stretched under \( \dot{\sigma}>50E_M \, Pa/s \), axial strain of tau protein at both ends \( (x=0 \) and \( L) \) is extremely large and tau protein axial strain in the region \( 0.1<x/L<0.9 \) is close to zero; the MT axial strain distribution surrounding the midpoint \( (x=L/2) \) remains constant. While the applied stress rate is very small, the tau protein axial strain distribution surrounding the midpoint \( (x=L/2) \) is not close to zero, and MT axial strain surrounding the midpoint is variable, as shown in Fig. 6a. Thus, we can conclude that during the loading process MT axial strain along the interface length is nearly unchanged, and tau-tau bonds break simultaneously in the end if the applied stress rate is small enough (quasi-static).
In order to clarify the relationship between the applied stress rate and mechanical response of axons containing Kelvin-Voigt viscoelastic tau protein, the time and spatial coordinate system should be rescaled as $T = \dot{\sigma} t$, $X = x/L$, in such a way that variables in Eq. (9) are normalized as $\Delta(X, T) = \delta/L$, $\partial \Delta/\partial T = (\partial \delta/\partial t)/(\dot{\sigma} L)$, $\partial^2 \Delta/\partial X^2 = (\partial^2 \delta/\partial x^2)/L$. According to the rescale variables, Eq. (9) can be converted to

$$\frac{\delta \mu}{K} \frac{\partial \Delta}{\partial T} = \frac{L^2}{L} \frac{\partial^2 \Delta}{\partial x^2} - \Delta \tag{16}$$

Eq. (16) implies that tau protein with high viscosity has similar relative sliding with high applied stress rate when all other parameters remain the same. On the contrary, tau protein with high spring constant has an opposite tendency. Interestingly, axon bundle can withstand strains of over 100% and recover to its original configuration without evidence of damage [7]. However, axons appeared to faille when subjected to a sudden change in tensile force [52]. Yuen et al. [53] reported that MT rupture begins at axon strain 5%, when axon is stretched under strain rate 50 s$^{-1}$. This viscoelastic shear lag model can explain why the axonal injury is sensitive to the stress rate.

### 3.3. Effect of the MT-tau interface length on stress transfer

Researchers found that the interface length affects the stress transferring and fracture propagation between fiber and matrix in the composites and biocomposites [54, 55]. Recently, finite element models were established to explore how MT-tau interface length affects the mechanical behavior of axon [10]. In this study, SLM including viscoelastic tau protein is constructed in order to illustrate the effects of MT-tau interface length on relative sliding and stress transfer under transient tensile force, as shown in Fig. 8.

While MTs are stretched by a force with a low stress rate $\dot{\sigma} = 0.001E_M$ Pa/s, tau protein axial strain along short MT-tau interface length ($L = L_C$) is very large (Fig. 8c). Whereas if the applied stress rate is $10E_M$ Pa/s, tau protein axial strains in both ends are identical, which indicates that the initiation of tau protein breaking starts simultaneously, and overall tau proteins along the short interface length come to failure in a shorter period of time (Fig. 8a). Normal stress in the MT will be concentrated at the both ends when the stress rate is high and MT-tau interface is long enough, as illustrated in Fig. 8b and 8d.

Histologically, long and short MTs are parallel aligned in the axon bundle. Bass et al. [56] found that different lengths of MT have different mechanical functions. Long MT is responsible for carrying loads, which results in a higher resistance to the sliding [57], and implies that long MTs are more susceptible to rupture. Yet, tau proteins in the short MT-tau interface are responsible for sliding, which may explain the significant elongation of axon following traumatic stretch. Our model prediction is consistent with the experimental observation proposed by Peter et al. [16].

![Diagram](image-url)
3.4. Effective stiffness of axon

Micromechanical analysis can provide a basis for the mechanical performance of macrostructures. For instance, the Young’s modulus of axon bundle could be computed from the ratio of the average force over the MT cross-sectional area and the average strain:

\[ E(t) = \frac{c_M L \sigma(L,t)}{2u_i(L,t)} \]  

(17)

where \( c_M \) is MT volume fraction. For the hexagonal lattice of MTs, \( c_M \) can be expressed as:

\[ c_M = \frac{\pi (R_o^2 - R_i^2)}{(d_o + 2R_i)^2 \sin 60^\circ} \]  

(18)

A mathematical model has been established to characterize the mechanical properties of axon at a nano-scale, where MTs and tau proteins act as coupled Kelvin-Voigt viscoelastic systems. Scientists have powerful experimental equipment, such as TEM, atomic force microscopy (AFM) and microelectromechanical systems (MEMS), etc., but it’s difficult to measure the elongation of tau protein along the MTs and axial strain of MTs. A general discussion on the stress-strain or force-displacement relation of axon, obtained from the experiments, can be found in Refs. [14, 58].

A series of in vivo tensile tests on axon were recently conducted to measure the mechanical properties...
of axon [14, 59-61]. Impressively, Rajagopalan et al. [14] reported that the axons behave like elastic springs under suddenly applied force condition. The effective stiffness of the axons, represented by the slope of the force-deformation curve, takes values in the range 0.2-1.2 nN/μm. The force-deformation curve of axonal bundle under tensile stress is plotted in Fig. 9. It shows that the slope of force-elongation curve of our current model agrees well with that of experimental data, which implies that our prediction for the effective stiffness of the axonal bundles is in excellent agreement with the existing experiment.

Fig. 9. Force-deformation curves of two axons from different Drosophila neurons. The triangles and circles represent experimental data measured by Rajagopalan et al. [14]. The dashed lines are fitting by experimental data. Our models are calculated with the parameter $E_M=1.5$ GPa, $d_T=20$ nm (red solid-line) and $E_M=1.0$ GPa, $d_T=40$ nm (green solid-line).

4. Discussion

Diffuse axonal injury (DAI) is one of the most common types of TBI [53]. Unfortunately, today’s imaging methods are unable to detect the microscopic diffuse injuries to the axon [62-64]. Moreover, tension of axon bundle is such a tricky experiment, which has been prompting scientists to resort to computational modelling. Computational modelling is considered as an efficient tool to predict the mechanical response of axons for substantially improving contemporary understanding of the underlying pathology and molecular mechanisms of axonal injury.

From the perspective of material mechanics, the stiffest structural component MTs reinforce the axon, guaranteeing high stiffness of axon. Additionally, weakest-linking tau proteins ensure large deformation of the axon. The stiffest discontinuous MTs are weakest-linked by tau protein, making vulnerable axon can withstand large deformation and tensile force, achieving high stiffness and toughness. Under low stress and low stress rate conditions, axons are highly compliant and ductile, and MTs slide smoothly.

Long MTs in the axon break with all tau proteins binding at high stress rate ($\sigma \geq 92.5E_M$ Pa/s). When the applied stress rate is low ($\sigma \leq 10E_M$ Pa/s), tau proteins along short MT-tau interface are more likely to break, and short MTs will detach from the MT bundle in the end. Under slightly high stress rate ($10E_M$ Pa/s$<\sigma <92.5E_M$ Pa/s), parts of the tau proteins located at the both ends of slightly long MTs break at first, and then MTs come to failure. In other words, MAP tau will sacrifice themselves to protect MTs from overall damage when $\sigma <92.5E_M$ Pa/s, and vice versa when it is under suddenly
applied forces condition ($\sigma \geq 92.5 E_M$ Pa/s). The type of axonal mechanical failure is multi-level, as illustrated in Fig. 10. Therefore, the vulnerable axon can achieve high stiffness and toughness, protecting itself from overall damage.

![Fig. 10.](image)

**Fig. 10.** Different types of deformation and damage evolution in the axon after TBI due to various MT-tau interface lengths and stress rates. Tau protein serves as a molecular switch between MT sliding, rupture, detachment and detachment-rupture.

In our previous study [65], we have explained that the fundamental load-carrying elements of regularly staggered biocomposite structures are discontinuous fibers, instead of continuous long fibers along the entire tissue. The discontinuous fibers are more capable to protect themselves from overall damage. As for axons, MTs and tau proteins will come to failure at different locations in the axon bundle, rather than break in a single section, and therefore an overall mechanical stability and high toughness could be achieved. This may be a key reason why randomly staggered biocomposite structures are observed in the axon.

The mechanical properties of engineering fiber reinforced composites depend to a great extent on the bond length. In contrast to the axon, high-performance fiber reinforced composites typically use continuous fibers, thus achieving high stiffness and strength but presenting limited toughness and ductility. From the above analysis, we know that the use of discontinuous fibers could potentially improve the ductility and fracture toughness [66, 67]. Longer interface length is not conducive to stress transfer, but discontinuous fibers could help dissipate energy and protect biocomposites from overall damage. Additionally, crack bridging by discontinuous fibers can make brittle materials tougher by transferring stress from the crack tip to the ductile matrix.

Despite the ultrastructure of the axon has the advantage of dissipating energy, MTs are vulnerable to rupture at high stress rate ($\dot{\sigma} \geq 92.5 E_M$ Pa/s), resulting in TBI. The rupture of MTs results in accumulation of transported cargo in axonal swellings [7]. Fortunately, MT has the ability to recover to its original configuration through self-repair [13, 68]. In addition, TBI may trigger chronic traumatic encephalopathy. Progressive axonal injury and structural degradation are considered as the classical features of chronic traumatic encephalopathy. These symptoms may be related to Parkinsonism and Alzheimer's disease [23, 28, 69].

From the perspective of biomedical engineering, this viscoelastic SLM is likely to become the core of the broader multi-scale model in time and space. As for the time, the computational window from
milliseconds to years can explain axonal injuries in the gradual deterioration of the mechanism of time. This can help identify the early markers of neurodegenerative disease and promote early treatment. As for the space, bridging from MTs to MAP tau can explain the molecular mechanism within the axon, which can provide new coping strategies to slow, block or reverse neurodegenerative disease [70]. This viscoelastic SLM may be a crucial step for shedding new light on the complicated interactions in MT-tau transient.

Though the viscoelastic SLM provides a new method to predict the axonal injuries under comparatively high stress rate over a typical timescale from microseconds to decades of seconds, the current study is not without limitations. A major limitation of this work is the choice of the Kelvin-Voigt model, which cannot represent stress relaxation. Surely if the axon is deformed and held at the final deformation, stress relaxation occurs in the tau protein. In order to accurately predict the underlying pathophysiology of DAI, future work will seek to improve the accuracy of the model by relaxing some assumptions and considering more of the related biomedical phenomena.

5. Conclusions
The aim of this study is to explore how the axon can simultaneously achieve a remarkable mechanical balance of high specific stiffness and toughness, and why the randomly staggered alignment microtubules are selected in the axon. Of course, viscosity is one of the significant physical attributes for axons [6, 7], which will cause them to be sensitive to loading speed. In this study, we have extended the previous SLM by including Kelvin-Voigt viscoelastic behaviors for the tau protein of the axon subjected to transient loading. This Kelvin-Voigt viscoelastic SLM is an essential extension to the previous SLM which only considered the elastic or elasto-plastic behaviors under static or quasi-static conditions. Kelvin-Voigt viscoelastic SLM has been developed to elucidate the biomechanical behavior between microtubules and tau protein in the axon under transient loading. Theoretical closed-form solutions is presented to predict the stress transfer and failure forms while axon is stretched transiently. According to the recent experiments [14, 60] and results presented in this paper, the conclusions can be drawn as follows:

(1) The maximum axial strain of tau protein is at both ends of the MT-tau interface length, and the minimum tau protein axial strain is at the center. The maximum and minimum axial strains of MTs are at the midpoint and both ends of the entire MT length, respectively.

(2) Axonal failure mechanism may be different under different loading conditions, such as microtubule rupture, detachment and detachment-rupture. In the process of MT-tau protein interface failure, tau protein serves as a molecular switch. Long microtubules are more vulnerable to rupture at high stress rate, yet short microtubules are likely to detach from the microtubule bundle under large deformations.

(3) Discontinuous, stiffest and randomly staggered alignment microtubules weakest-linked by tau proteins enable axon to withstand large deformations and have the advantage of dissipating energy. In this case, the forms of axonal mechanical failure is multi-level resulting in protecting the axon from overall damage. Therefore, composites with randomly staggered element can simultaneously achieve an outstanding mechanical balance of high strength and high toughness.
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