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PERCOLATION STUDY OF NANO-COMPOSITE CONDUCTIVITY USING MONTE CARLO SIMULATION

by

JING BAI
B.S. Jiang Nan University, 2007

A thesis submitted in partial fulfillment of the requirements for the degree of Master of Science in the Department of Mechanical, Materials and Aerospace Engineering in the College of Engineering and Computer Science at the University of Central Florida Orlando, Florida

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Major Professor: Kuo-Chi Lin
A Monte Carlo model is developed for predicting electrical conductivity of carbon nanofiber composite materials. The conductive nanofibers are models as both 2D and 3D network of finite sites that are randomly distributed. The percolation behavior of the network is studied using the Monte Carlo method, which leads to the determination of the percolation threshold. The effect of the nanofiber aspect ratio on the critical nanofiber volume rate is investigated in the current model, each of the nanofibers needs five independent geometrical parameters (i.e., three coordinates in space and two orientation angles) for its identification. There are three controlling parameters for each nanofiber, which includes the nanofiber length, the nanofiber diameter, and the nanofiber aspect ratio. The simulation results reveal a relationship between the fiber aspect ratio and the percolation threshold: the higher the aspect ratio, the lower the threshold. With the simulation results obtained from the Monte Carlo model, the effective electrical conductivity of the composite is then determined by assuming the conductivity is proportional to the ratio of the number of nanofibers forming the largest cluster to the total number of nanofibers. The numerical results indicate that as the volume rate reaches a critical value, the conductivity starts to rise sharply. These obtained simulation results agree fairly with experimental and numerical data published earlier by others. In addition, we investigate the convergence of the current percolation model. We also find the tunneling effect does not affect the critical volume rate greatly. We propose that the percolation model is not scalable as well.
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Table 5: Volume rates under different dimensions with $L = 0.05$, $R = 0.0005$, $L/D = 50$ ............ 52
A variety of conductive particles or fibers have been added to polymer matrices to form nanocomposites, finding important applications in aerospace and other industries. Carbon nanotubes and carbon nanofibers have been an area of intense research and play a significant role in current development of nanotechnology. They own unique characteristics and properties, which includes low density, high surface area, good chemical stability, high electrical and thermal conductivity, excellent mechanical properties and fire resistance. (Krishnan et al., 1998; Yu et al., 2000; Sandler et al., 1999; Geng et al., 2002; Beguin et al., 2002; Subramony et al., 1998; Kashiwagi et al., 2002; Kashiwagi et al., 2005). For example, as the volume fraction increasing, there exists an insulator to conductor transition around a critical threshold (Bigg, 1979). That is, there is a sharp increase of the electrical conductivity at a critical fiber volume fraction. Polymer matrix composites reinforced by nickel nanostrands, which are pure nickel filaments with nanometer diameters and high aspect ratios, are very promising nanocomposites that can reduce lightning strike damage with minimum added weight (Hansen, 2005; Li et al., 2008).
Carbon nanotubes consist of graphitic cylinders with diameter of 1-100 nm and high aspect ratio of a few micro lengths, leading to high Van der Waals force between adjacent tubes. The high Van der Waals force and high aspect ratios with a combination of high flexibilities make these nanotubes easily assemble into bundles or ropes. As a result, it becomes quite difficult to disperse carbon nanotubes in polymer matrix. In addition, their manufacturing and purification processes are still too complicated and expensive for mass production. All these factors limit their application in manufacturing high performance carbon nanotubes/polymer composite (Hill et al., 2005; Thess et al., 1996; Kokai et al., 1999).

Compared to carbon nanotubes, carbon nanofibers have recently gained considerable attention in nanocomposite applications. Carbon nanofibers can be produced at lower cost by catalytic chemical vapor deposition of carbon containing gases (Toebes et al., 2004; Ros et al., 2002; Vieira et al., 2004; Dandekar et al., 1998; Durkic et al., 1997; Wang et al., 2003). They possess less order and more edge sites on the outer wall (Kim and Lee, 2004; Werner et al., 2005), which makes carbon nanofibers have better dispersion and wettability in polymer matrix. The higher proportion of edge plane defects also may lead to more facile electron transfer (Salimin et al., 2004; Banks and Compton, 2005). In fact, carbon nanofibers can be prepared having grapheme sheets stacked either parallel (ribbon structure), perpendicular (platelet structure), or canted (herring-bone structure) relative to their long axis (Rodriguez et al., 1995; Carneiro et al., 2003). The exterior surface of carbon nanofibers possessing a herring-bone structure is composed of highly reactive carbon or partially hydrogenated carbon edge-sites. This structural characteristic provides a unique opportunity for covalent binding of linking molecules containing reactive pendant functional groups to the surfaces of carbons nanofibers. The reaction of such surface-
bound pendant functional groups with polymers could make carbon nanofiber/polymer nanocomposites exhibit exceptional carbon fiber-polymer wettability due to the presence of covalent binding across the nanofiber-polymer interface (Zhong et al., 2004). Based on the as-mentioned characteristics, an increasing number of carbon nanofiber/polymer nanocomposites have been researched. (Thostenson and Chou, 2006; Yang et al., 2005; Lau et al., 2005; Shi et al., 2007; Hirai et al., 2007; Cho and Bahadur, 2007; Hasan et al., 2007; Kobayashi and Kawai, 2007; Yang et al., 2007)

Percolation theory has been widely used to describe the effective conductivity of a composite material (Kirkpatrick, 1973; Lebovka et al., 2006; Stauffer, 1979). Percolation theory, originally, was to study the problem involving liquid passing through the porous materials and answer the question that if the liquid can make it from hole to hole and reach the other side of the material. It was later developed into a pure mathematical problem that describes the behavior of connected clusters in a random graph. This model is perfect for our purpose that treats the nanofiber cluster as the paths of electronics and study conductivity due to those paths.

In the “site percolation” and “bond percolation” model, the conductive fillers in a composite are models as 2D and 3D, and they are randomly and regularly located in a composite system. In the “bond percolation” model, a known fraction of bonds, distributed randomly, are missing from the lattice and the current can not form between them. In the “site percolation” model, the absence of a site means there is no current flow through any of the bonds which join the site to its neighbors. So if the possibility of allowing the current to go through is \( p \) between each two neighboring sites, then there is no connection between them with the possibility of \( 1 - p \). We have observed
the following phenomenon in the laboratory. In an insulated substrate, we add the conductive carbon nanofiber to form a composite. When the volume ratio of the nanofiber is low, the conductivity of the composite is very low and even zero. As the increasing of the nanofibers, when the volume ratio reaches a critical value, the conductivity starts to rise sharply. Therefore, there exists a threshold value $p_c$, above which a continuous conductive path forms. The probability of carbon nanofibers forming a conductive path is very low when the volume rate is low. Accordingly, the conductivity of the composite is close to zero. After the volume rate reaches a critical value, the carbon nanofibers start to form at least one conductive path and the possibility of existing multiple conductive paths increases; therefore the conductivity of the composite increases sharply. Many researchers has studied the electrical conductivity of the composite system and investigated the percolation threshold in order to better understand the principle of the sharp change in conductivity near the critical transition point.

Monte Carlo methods are a class of computational algorithms that rely on repeated random sampling to compute their results. Monte Carlo methods are often used when simulating physical and mathematical systems. Because of their reliance on repeated computation and random or pseudo-random numbers, Monte Carlo methods are most suited to calculation by computer. Monte Carlo methods tend to be used when it is infeasible or impossible to compute an exact result with a deterministic algorithm.

Monte Carlo simulation methods are especially useful in studying systems with a large number of coupled degrees of freedom, such as fluids, disordered materials, strongly coupled solids, and
cellular structures. More broadly, Monte Carlo methods are useful for modeling phenomena with significant uncertainly inputs.

The Monte Carlo simulation is a widely used numerical technique of modeling the effects of disorder resistor networks and can be applied to a wide range of situations. By using the method, the percolation behavior is appropriately simulated with advanced computational algorithms and the topological disorder of a random microstructure is directly constructed. In 1974, Pick and Seager (1974) carried out a 2D study on composites filled by straight sticks. They assumed that all the sticks have the equal length and no width. Their work was extended by Balberg and Binenbaum (1983), who account for the macroscopic anisotropy induced by sticks with preferred orientation and unequal length. Natsuki et al. (2005) considered the width of sticks in their Monte Carlo simulation of 2D networks, which also predict the relationship between the percolation threshold and the fiber aspect ratio and the orientation angle.

The Monte Carlo simulation can also be used to study the 3D percolation behavior. Balberg et al. (1984) first conducted the 3D Monte Carlo study on percolation behavior of systems consisting of randomly oriented sticks, and they explored the effects of the stick aspect ratio and macroscopic anisotropy. By developing the study of Balberg et al and the effective medium method, Taya and Ueda (1987) used the Monte Carlo approach to study the electrical conductivity of a 3D composite reinforced by straight short fibers. Lee and Kim (1995) also performed the 3D Monte Carlo simulation for unidirectional short-fiber reinforced composites. In their research, the fiber orientation was predetermined and the fiber length distribution, fiber volume fraction and fiber aspect ratio are allowed to change.
Few attentions have been paid to the effects of the fiber waviness on the percolation behavior of fibrous composites. Ye et al. (2004) predicted that the increasing of the curliness of the fibers cause the greatly increasing of the percolation threshold. Li and Chou reported a general continuum percolation model for the composites with fibers of arbitrary shape in 2D scale. In 3D aspect, Dalmas et al (2006) developed the simulation of 3D entangled fibrous networks using spline-shaped fibers. They reported that the percolation threshold increased as the increase of the fiber tortuosity and the larger fiber aspect ratio can also bring the larger effect of the tortuosity. However, they did not study the effect of the fiber width in the 3D simulations. Therefore, there still need to develop 3D Monte Carlo models to explore the effects of the fiber aspect ratio, fiber width, fiber length and fiber aspect ratio on the behavior of the composites system.

1.2 Research Objectives

This work has following research objectives:

1. Development of a conductivity percolation model for the carbon nanofiber composite materials. The model is based on the randomly generated cylinders, which model the nanofibers, forming clusters through the tunneling effect. When a cluster spans the substrate from one side to the opposite side, percolation happens.

2. Using Monte Carlo simulation method, study the convergence of current model and by statistical analysis, we get the density distribution of the critical volume rate.
3. Simulation results demonstrate the percolation phenomenon when the volume rate of the nanofibers in the composite exceeds a critical value. Fundamentally understanding the relationship between the nanofiber aspect ratio and the fiber volume rate.

4. The research also studies the tunneling effect to the average volume rate, and proposes the percolation model is not a scalable system. The electrical conductivity also studied for both the 2D and 3D models.

5. The simulation model shows the qualitative characteristics of the nanocomposite percolation behavior. This study will lead to the better understanding of the influence of certain parameters to reduce the number trials and errors in the experiments.
CHAPTER 2
PERCOLATION CONCEPTS AND APPROACHES

2.1 Background

Percolation theory was to study the problem involving liquid passing through the porous materials and answer the question that if the liquid can make it from hole to hole and reach the other side of the material. It was later developed into a pure mathematical problem that describes the behavior of connected clusters in a random graph. This model is perfect for our purpose that treats the nanofiber clusters as the paths of electronics and study conductivity due to those paths. The percolation models are composed of sites and of bonds between sites.

2.2 Site percolation and bond percolation

Broadbent and Hammersley (1957) proposed the first mathematical model of the classical threshold. They reported a simplified “lattice percolation” model for the flow of a fluid through a porous medium from one side to the other side and showed clearly their model possessed a percolation threshold. The model requires fluid to go through the “bonds” which connect to the nearest neighbors in a regular lattice of “sites”. In the “bond percolation”, some known fraction of bonds, which distributed randomly, is missing from the lattice. In the “site percolation”, a
known fraction of the sites are missing from the lattice causes no current can flow through any of
the bonds which join that site to its neighbors, so no current can go through the sites.

The research of the Kirkpatrick (1973) pointed that in the case of the site percolation on a lattice
which includes \( N \) sites, where the number \( N \) is large. The parameters relevant to percolation are
defined in the limit \( N \to \infty \), and will depend on the concentration of connected sites \( x \) and the
geometry of the lattice. When the concentration is low, for example \( x \leq x_c \), the allowed sites are
single and in small separated clusters of adjacent allowed sites. As the increase of \( x \), larger
cluster formed and the mean size of a cluster increased accordingly. As \( x \) approaches \( x_c \) from
below, the larger clusters begin to merge and the mean cluster size diverges at \( x_c \) in the limit
\( N \to \infty \). For the finite \( N \), this indicates that there is a completed path of neighboring allowed
sites crossing the system, therefore the macroscopic flow through the system becomes possible.
If \( N \) is large enough, there will be only one large cluster remains in the system, along with many
small ones. As \( x \) increases above \( x_c \), the infinite cluster grows rapidly and absorbs small
clusters. Accordingly, the site percolation probability \( P^{(s)}(x) \) rises greatly from zero just above
threshold, and the isolated clusters become rare. \( P^{(s)}(x) \) near threshold can be demonstrated by a
simple power law,

\[
P^{(s)}(x) \propto (x - x_c)^s
\]

(1)

Where \( s \) is approximately the same for all three lattices.

\[0.3 \leq s \leq 0.4\].

Thresholds of this type can be applied to all the lattices and are not restricted to nearest neighbor
percolation on regular lattices.
2.3 Effective Medium Theory

The effective medium theory, which was originally formulated to describe the conductivity of binary mixture, has been developed to treat disordered networks. Moreover, some attempts to generalize the theory to anisotropic random network were made to verify some general aspects of conduction in anisotropic materials. During the evolution of such theories many methods have been used to obtain statistical information about the heterogeneous systems [50].

The distribution of potential in a random resistor network to which a voltage has been applied along one axis may be regarded as due to both an “external field” which increases the voltages by a constant amount per row of nodes, and a fluctuating “local field,” whose average over any sufficiently large region will be zero [51]. We use an effective medium, in which the total medium inside is equal to the external field, to represent the average effects of the random resistors by an effective medium. Such a medium should be homogenous. To simplify the model, we consider it to be made up of a set of equal conductance, $g_m$, connecting the nearest neighbors on the cubic mesh. The criterion to fix $g_m$ is the extra voltages induced the local fields, when individual conductance $g_g$ replaces $g_m$, in this medium, should average to zero.

Consider one conductance oriented along the external field, surrounded by the effective medium, and having the value $g_{AB} = g_0$. The solution of the network equation
\[ \sum_j g_j (V_i - V_j) = 0 \]  \hspace{1cm} (2)

Figure 1: Constructions used in calculating the voltage induced across one conductance, \( g_0 \), surrounded by a uniform medium [51].

In the presence of \( g_{AB} \) is constructed by superposition. To the uniform field solution, in which the voltages increase by a constant amount, \( V_m \), per row, we add the effects of a fictitious current, \( i_0 \), introduced at A and extracted at B. Since the uniform solution fails to satisfy current conservation at A and B, the magnitude of \( i_0 \) is chosen to correct for this:

\[ V_m (g_m - g_0) = i_0 \]  \hspace{1cm} (3)

The extra voltage, \( V_0 \), induced between A and B, can be calculated if we know the conductance \( G_{AB} \) of the network between points A and B when \( g_{AB} \) is absent.

\[ V_0 = i_0 / (g_0 + G_{AB}) \]  \hspace{1cm} (4)

To calculate \( G_{AB} \), we first obtain the conductance \( G_{AB} \) between A and B in the uniform effective medium, since \( G_{AB} = G_{AB}^\prime + g_m \). A symmetry argument is useful: Express the current distribution
with \( g_0 = g_m \) as the sum of two contributions, a current \( i_0 \), introduced at A and extracted at a very large distance in all directions, and an equal current, introduced at infinity and extracted at B. In each case, the current flowing through each of the \( z \) equivalent bonds at the point where the current enters is \( i_0 / z \), so that a total current of \( 2i_0 / z \) flows through the AB bond. This determines the voltage developed across AB, and from that follows the result, \( G_{AB} = (z/2)g_m \), or \( G_{AB}' = (z/2 - 1)g_m \). Using Eqs. (3) and (4), we obtain:

\[
V_0 = V_m(g_m - g_0) /[g_0 + (z/2 - 1)g_m]
\] (5)
valid in both 2D and 3D.

If the value of a bond, \( g_{ij} \), is distributed according to a probability distribution \( f(g) \) (which may be either continuous or discrete), the requirement that the average of \( V_0 \) vanish gives a condition determining \( g_m \):

\[
\int dg f(g)(g_m - g)/[g + (z/2 - 1)g_m] = 0
\] (6)

For a binary distribution, we have

\[
f(g) = p\delta(g - 1) + (1 - p)\delta(g - \alpha)
\] (7)

As is appropriate to the percolation network models, Eq. (6) reduces to a quadratic equation for \( g_m \):

\[
(z/2 - 1)g_m^2 + \{(z/2)p - 1 + \alpha[z/2(1 - p) - 1]\}g_m - \alpha = 0
\] (8)

The relevant root of Eq. (8)

\[
g_m = \{(z/2)p - 1 + \alpha[z/2(1 - p) - 1]\}/(z - 2) + \{(z/2)p - 1 + \alpha[z/2(1 - p) - 1]\}^2 + 2(z - 2)\alpha^{1/2}(z - 2)^{-1}
\] (9)
Has a simple limiting form when $\alpha \to 0$:

$$g_m(\alpha = 0) = 1 - (1 - p)/(1 - 2/z)$$  \hspace{1cm} (10)

A straight line in which $g_m$ goes to zero when $p = 2/z$.

It was shown in Kirkpatrick (1971), the effective medium theory is accurate to within a few percent at all concentrations whenever $\alpha \geq 0.1$ for the bond percolation model. However, for the site percolation model, it gives the wrong slope at high concentration (small concentrations of missing sites).

### 2.4 Excluded Volume Method

There are several methods capable of predicting the critical concentration in percolating systems. The excluded volume approach is effective in treating the problem of percolating systems in which the objectives possess a large aspect ratio [53]. The excluded volume and excluded area in two dimensions of an object is defined as the volume or area around an object into which the center of another similar object is not allowed to enter if overlapping of the two objects is to be avoided. The excluded volume for an elongated volume can be obtained by multiplying this volume by $N_c$, which is the critical number density of objects in the system. If $\langle V \rangle$ defines an average excluded volume, the total excluded volume $\langle V_{ex} \rangle$ is given by

$$\langle V_{ex} \rangle = \langle V \rangle N_c$$  \hspace{1cm} (11)

Similarly, the total excluded area is
\[ \langle A_{\alpha} \rangle = \langle A \rangle N_c \]  

(12)

Remember that \( \langle V \rangle \) and \( \langle A \rangle \) of an object is very different, and its value can be calculated according to the shapes such as cylindrical rods in three-dimensions (3D) and line segments and narrow strips in two-dimensions (2D).

Figure 2: Two sticks of length L and width D, the angle between which is \( \theta \). The excluded area is obtained by following the center o as stick \( j \) travels around stick \( i \) while touching it at least at one point [54].

Two sticks (rectangles), the angle between which is \( \theta = \theta_i - \theta_j \) [54]. The excluded area can be obtained simply by moving one stick around the other and registering the center of the moving stick. Fig.2 shows a result of such a procedure. The shaded area represents the stationary stick and the curve is the path of the center of the other sick as it is moved around the first stick. The area within the curve is the excluded area.
This excluded area is given by
\[ (L \sin \theta + D + D \cos \theta)(L + D \sin \theta + L \cos \theta) - (L^2 + D^2) \sin \theta \cos \theta \] (13)

For the uniform distribution of angles we must consider all possible angles \( \theta_i \) and \( \theta_j \) and their corresponding uniform probability
\[ P(\theta) = 1/2\theta \] (14)
in the interval \( 2\theta \). Hence the averaged excluded area is
\[ \langle A \rangle = L^2 \int_{-\pi/2}^{\pi/2} \int_{-\pi/2}^{\pi/2} \sin |\theta_i - \theta_j| P(\theta_i)P(\theta_j) \times d\theta_i d\theta_j \] (15)

Substituting the distribution (14) and (15) yields the average excluded area
\[ \langle A \rangle = 2DL[1 + \left( 1/2\theta \right)^2 (1 - \cos 2\theta)] + (L^2 + W^2)(4\theta - 2\sin 2\theta)/(4\theta^2) \] (16)

The average excluded volume \( \langle V \rangle \) for a CNT, modeled as a cylinder of length \( L \) and diameter \( D \), is given by [54]:
\[ \langle V \rangle = \frac{4}{3} \pi D^3 + 2\pi D^2 L + 2DL^2 \langle \sin \theta \rangle \] (17)

Where \( \langle \sin \theta \rangle \) is the average value of \( \sin \theta \), and \( \theta \) is an angle between two CNTs.

The term \( \langle \sin \theta \rangle \) describes the degree of CNT alignment. The calculation of \( \langle \sin \theta \rangle \) is complex. However, it is easy to obtain \( \langle \sin \theta \rangle = 0 \) when the CNTs are perfectly aligned. In the case of random distribution, the value of \( \langle \sin \theta \rangle \) is calculated to be \( \pi / 4 \).
Another two-dimensional finite-width stick is the “capped” rectangular stick. The derivation of the excluded area of this object indicates how to handle the three-dimensional problem. We assume now a rectangle of length $L$, width $D$, and the caps of radius $D/2$ at its ends. Fig. 3. shows the capped rectangle and the excluded area which is formed around it. One can readily find that the excluded area for these two sticks, which have an angle $\theta$ between them, is

$$4DL + \pi D^2 + L^2 \sin \theta$$

(18)

The average excluded area in the uniform and randomly orientated system is given as

$$\langle A \rangle = 4DL + \pi D^2 + \left(\frac{L}{2\theta_{\mu}}\right)^2 [4\theta_{\mu} - 2\sin(2\theta_{\mu})]$$

(19)

Where $W$ is the width of fibers and $\theta_{\mu}$ is the orientation angle.

Figure 3: The sticks are capped rectangles. The length of the sticks is $L$, their width is $D$ and the radius of the cap is $D/2$ [54].
The critical volume fraction of CNT is associated with total excluded volume \( \langle V_{ex} \rangle \). In the 3D system, the critical volume fraction can be estimated by

\[
\phi_c = 1 - \exp\left(-\frac{V_{ex} \nu}{V}\right)
\]  

(20)

Where \( \nu \) is the volume of CNTs capped at each end with a hemisphere.

Substituting Equation (13) into Equation (15), we have:

\[
\phi_c = 1 - \exp\left(-\frac{\langle V_{ex} \rangle \left[\frac{\pi}{4}D^2 L + \frac{\pi}{6}D^3 \right]}{(4\pi/3)D^3 + 2\pi D^2 L + 2DL(\sin \theta_{\mu})}\right)
\]  

(21)

Similarly, we obtain the critical area fraction in 2D system:

\[
\phi_c = 1 - \exp\left(-\frac{\langle A_{ex} \rangle \left[WL + \frac{\pi}{4}W^2\right]}{4WL + \pi W^2 + \left(L/2\theta_{\mu}\right)^2 \left[4\theta_{\mu} - 2\sin(2\theta_{\mu})\right]}\right)
\]  

(22)

The analytical percolation model, the excluded volume theory, also assumes that the percolation threshold is inversely proportional to the object excluded volume \( V_{ex} \) \([54, 55]\)

\[
\phi_c \propto \frac{1}{V_{ex}} \quad (23)
\]

Bug et al.\([42]\) have shown that for very thin rods \( R/L \rightarrow 0 \), the proportionality in Eq. (23) becomes an equality. Nevertheless, when the fibers are curved, the percolation threshold is underestimated by this excluded volume theory. A tortuous high aspect ratio fiber can be considered as a filler with an effective aspect ratio lower than that of the same straight fiber, leading to a lower excluded volume and a higher percolation threshold.
CHAPTER 3
NUMERICAL MODEL

In this chapter, we present two steps involving in developing the Monte Carlo model. First of all, the fibers with randomly location and random orientation are generated. Since the main goal of this research is to determine the conductive filler volume rate at the onset of high electrical conductivity, a non-dimensionalized unit cube will be used. In the second step, a percolation criterion is applied to check the connectivity between each pair of fibers in the composite system. The details of these two steps are described as follows.

3.1 Model generation

The following phenomena have been observed in the laboratory. Conductive carbon nanofiber is added into an insulated substrate to form a composite. When the volume ratio of the nanofiber is low, the conductivity of the composite is zero or very low. If we continue to add more nanofibers into the composite, as the volume ratio reaches a critical value, the conductivity starts to rise sharply. This phenomenon is theorized as follows. When the volume ratio is low, the probability of carbon nanofibers forming a conductive path is very low. Therefore, the conductivity of the composite is close to zero. As the volume ratio reaches a critical value, the carbon nanofibers start to form at least one conductive path. As the increase of volume ratio, the
probability of forming multiple conductive paths increases; therefore the conductivity of the composite increases sharply [56].

A numerical model of a nanofiber composite is built as the follows. A set of nanofibers, modeled as cylinders, are generated within the domain of the substrate with their positions, orientations, lengths, and radius following certain types of random distributions. All the fibers in the system are identical, for example, they have the same length, diameter and orientation angles. But they are randomly distributed and oriented. To generate numbers with sufficient randomness, we adopt the most common computer technique for producing random sequences, the multiplicative congruential generator.

3.2 Connection criterion

Percolation problem is to find the critical length $L_c$ by which an infinite cluster of connected sites are formed. Let us consider a sample which is a unit-size square. In the square, we plant randomly distributed sites $N_s$, defined as a stick density. The next stage is to attach a stick of length $L$ to each site according to a certain fiber orientation. A critical length of the percolation threshold depends on the stick density in the sample. We can give the average lattice constant normalized by [39]

$$r_s = 1/\sqrt{\pi N_s}$$

(24)
Percolation threshold is reached if and only if a continuous pathway of overlapping sticks exists between connecting opposing edges. The critical length $L_c$ can be normalized by $2r_s$ given by Eq. (24). The value $L_c / 2r_s$ is dependent on the orientation angle and aspect ratio.

Bonding criterions for two sticks with a finite-width can be analyzed by the schematic diagram shown in Fig. (4) and (5). Sites are planted by generating random coordinates $(x_k, y_k)$, $k = 1, 2, ..., N_s$. Each stick of length $L_k$ is centered on a site and has an assigned orientation angle $\theta_k$. Fig. 4 illustrates the relationship between two different coordinate systems that are fixed on fibers. The axes $y'$ and $y''$ indicate the orientation angles of $\theta_i$ and $\theta_j$, respectively. The relationship between the Cartesian coordinates shown in Fig.4 can be expressed as

$$\begin{align*}
\begin{bmatrix} x \\ y \end{bmatrix} &= \begin{bmatrix} x_i^0 \\ y_i^0 \end{bmatrix} + \begin{bmatrix} \cos \theta_i & \sin \theta_i \\ -\sin \theta_i & \cos \theta_i \end{bmatrix} \begin{bmatrix} x' \\ y' \end{bmatrix}, \\
\begin{bmatrix} x \\ y \end{bmatrix} &= \begin{bmatrix} x_j^0 \\ y_j^0 \end{bmatrix} + \begin{bmatrix} \cos \theta_j & \sin \theta_j \\ -\sin \theta_j & \cos \theta_j \end{bmatrix} \begin{bmatrix} x'' \\ y'' \end{bmatrix},
\end{align*}$$

(25)

(26)

Eliminating $x$ and $y$ from the above equations, we obtain

$$\begin{align*}
y' &= \frac{(\Delta x \cos \theta_j - \Delta y \sin \theta_j) + x' \cos(\theta_i - \theta_j) - x'}{\sin(\theta_j - \theta_i)} \\
y'' &= \frac{(\Delta x \cos \theta_i - \Delta y \sin \theta_i) + x'' \cos(\theta_j - \theta_i)}{\sin(\theta_j - \theta_i)}
\end{align*}$$

(27)

(28)

Where

$$\begin{align*}
\Delta x &= \begin{bmatrix} x_i^0 - x_j^0 \\ y_i^0 - y_j^0 \end{bmatrix}
\end{align*}$$

(29)
Let us consider the case of a finite-width stick, as shown in Fig.4. Since bonding occurs when two sticks overlap, the bonding criterions are satisfied by the following both conditions:

\[
\begin{align*}
|\bar{x}| & \leq D/2, \quad |\bar{x}'| \leq D/2, \\
|\bar{y}| & \leq L/2, \quad |\bar{y}'| \leq L/2,
\end{align*}
\]

(30)

Where \(D\) and \(L\) are the stick width and length, respectively. When two sticks have the same orientation angle (Fig. 5), the bonding criterions are given by

\[
A_{ij} = d_{ij} \left| \sin(\theta_i + \gamma_{ij}) \right| \leq \frac{L_i + L_j}{2},
\]

(31)

and

\[
B_{ij} = d_{ij} \left| \cos(\theta_i + \gamma_{ij}) \right| \leq D,
\]

(32)

Where

\[
d_{ij} = \sqrt{(x_i - x_j)^2 + (y_i - y_j)^2}
\]

(33)

\[
\gamma_{ij} = \tan^{-1} \left( \frac{y_i - y_j}{x_i - x_j} \right)
\]

(34)

For widthless sticks \((x' = x = 0)\), we can obtain the bonding criterions from Eqs.(27), (28) and (30) as follows:

\[
\left| \frac{\Delta x \cos \theta_j - \Delta y \sin \theta_j}{\sin(\theta_j - \theta_i)} \right| \leq \frac{L}{2},
\]

(35)

\[
\left| \frac{\Delta x \cos \theta_i - \Delta y \sin \theta_i}{\sin(\theta_j - \theta_i)} \right| \leq \frac{L}{2}.
\]

(36)

If the points of stick intersection are outside the unit square sample, the two sticks are not considered intersection.
Figure 4: Diagram of the relationship between orientated fibers in different coordinate systems [41].

Figure 5: Diagram of determining bonding criterion of fibers with the identical orientation angle [41].

For small aspect ratio, the geometry of fiber ends strongly affects the percolation threshold because of the intersection between boundaries. Fig.6 shows there are three patterns of overlapping between fibers: body-to-body, end-to-body, and body-to-end.
For the end-to-end and end-to-body connection patterns, two fibers do not satisfy the above bonding criterions but are still connected. The end-to-end and end–to- body connecting patterns are more time consuming to model than the body-to-body connection in the Monte-Carlo simulations. However, after examined the fraction of the number of these two types of connections to the number of total connections among all fibers in the system, we found that this fraction drastically decrease with the increase of the fiber aspect ratio. The difference in the critical fiber volume fraction between these two sets of simulation results is found to be negligibly small when the fibers have an aspect ratio larger than 24. Therefore, some previous research has neglected the end-to-end and end-to-body connections for the sake of computational efficiency.

3.3 Calculate the shortest distance between two fibers in space
Percolation occurs in the composite system if a continuous conducting pathway along any one or all of the three coordinate axes can be identified. To evaluate the status of the system generated above, each fiber in the system is checked against another to see whether they intersect. In the current study, the connectivity between the \( i \)th and \( j \)th fibers is determined by comparing the shortest distance between the critical threshold.

Fig. 7 is the simplified diagram which is used to explain the calculation procedure with vector.

![Diagram](image)

Figure 7: Two-dimension reduction of the shortest distance between two fibers

We assume the following conditions:

The plane \( H \) is constructed parallel to \( \overrightarrow{PQ} \).

\( \overline{PQDC} \) is a rectangle.

\[
\|\overrightarrow{XZ}\|^2 = \|\overrightarrow{XY}\|^2 + \|\overrightarrow{YZ}\|^2, \text{ with } \|\overrightarrow{YZ}\| = \text{dist}(\overrightarrow{PQ}, \text{plane } H) = \text{dist}(\overrightarrow{PQ}, \overline{CD})
\]  

(37)
Therefore, with

\[ d_3 = \min \left\{ \| XZ \| : X \in \overline{AB}, Z \in \overline{PQ} \right\} \]

\[ d_2 = \min \left\{ \| XY \| : X \in \overline{AB}, Y \in \overline{CD} \right\} \]

\[ d_1 = \text{dist}(\overline{PQ}, \text{planeH}), \quad \text{a const tan} \ t \]

it follows

\[ d_3^2 = d_2^2 + d_1^2 \]  

(38)

And there is a one-to-one correspondence between solutions of the two minimization problems.

**Proof**

\[ \min \left\{ \| XZ \|^2 : X \in \overline{AB}, Z \in \overline{PQ} \right\} \]

\[ = \min \left\{ \min \left\{ \| XZ \|^2 : Z \in \overline{PQ} \right\} : X \in \overline{AB} \right\} \]

\[ = \min \left\{ \min \left\{ \| XY \|^2 + \| YZ \|^2 : Y \in \overline{CD} \right\} : X \in \overline{AB} \right\} \]

\[ = \min \left\{ \min \left\{ \| XY \|^2 + d_1^2 : Y \in \overline{CD} \right\} : X \in \overline{AB} \right\} \]

\[ = d_1^2 + \min \left\{ \min \left\{ \| XY \|^2 : Y \in \overline{CD} \right\} : X \in \overline{AB} \right\} \]  

(39)

### 3.4 Computational implementation

The shortest distance between each pair of nanofibers is calculated. If this distance is below a certain threshold, the two nanofibers are assigned to the same cluster. After all nanofibers are
assigned, each cluster is checked to see if it spans the substrate, i.e., providing a connection path from one side to the opposite side. Once percolation happens, the conductivity is calculated based on the cluster or clusters that constitute the connection paths. The process is repeated with different distributions of the nanofiber sets to obtain statistical data.

Each fiber is assigned a fiber number and a cluster number, when the generation procedure is completed. The fiber number and the cluster number are equal and range from 1 through $N$, where $N$ is the total number of fibers in the system. Then, each fiber is checked for connection with other fibers whose fiber numbers are larger than its fiber number. For example, the $i$th fiber will be checked against the $i+1$th through the $N$th fiber. If two fibers satisfy the connection criterion, they will be assigned a common cluster number which is the smaller one of the two fiber number. As a result, all fibers within the same cluster have the same cluster number, and two clusters are given the same cluster number if they have a common fiber.

If any two fibers in opposite boundary regions have the same fiber cluster number, then it can be concluded that the system is percolated in the direction perpendicular to the two opposing bounding surfaces. When the first percolating cluster is found, the system is said to be in the critical state where the fiber critical volume fraction has been reached.
CHAPTER 4
RESULTS AND DISCUSSION

In the percolation model developed in the preceding chapter, there are three controlling parameters for a given system: the fiber length $L$, the fiber diameter $D$ and the fiber aspect ratio $\alpha$, which are all predetermined. The percolation threshold depends on these specified parameters.

For a given set of the values of the three controlling parameters, the number of fibers in the system $N$ will be increased in small increments (with the random number generator reset for each increment) until the first cluster connecting the two opposite boundaries of the system is identified, which corresponds to one critical value of the fiber volume fraction.

4.1 Simulation Results

The following are examples of the two dimensional cases for easy demonstration. There are no major differences when the model is extended to the three-dimensional cases. The substrate is a 1 by 1 square (normalized dimension). The lengths of the nanofibers are between 0.10 and 0.12, evenly distributed. The aspect ratio, $L/D$, of the nanofibers is 200; therefore, the radius of the
nanofiber is between 0.00025 and 0.00030, also evenly distributed. The threshold of the tunneling effect is 0.0005.

Figure 8 shows the result of a set of nanofibers with 2.42% volume rate. The nanofibers in black (darker) color are the largest cluster, which is of the size of about 25% of the total nanofibers. The rest of the nanofibers, in the lighter color, are grouped into a number of smaller size clusters. There is apparently no percolation and the conductivity is zero.

Figure 8: A set of nanofibers with 2.42% volume rate

Figure 9 shows the result of the set of nanofibers increased to a volume rate of 2.87%. The additional nanofibers have made the size of the largest cluster grow to 47% of the total nanofibers. From the figure, we can see there is still no percolation and the conductivity is zero. However, with a few additional nanofibers, the percolation may happen.
Figure 9: A set of nanofibers with 2.87% volume rate

Figure 10: shows the result of the same set of nanofibers with the volume rate increased to 2.90%. Percolation starts to happen. The largest cluster grows to about 72% of the total nanofibers
Figures 11, 12, and 13 show the results of volume rates increased to 3.03%, 3.63%, and 4.84%, respectively. The largest clusters are of 76.6%, 98.2%, and 99.8%, respectively, of the total nanofibers.

Figure 11: A set of nanofibers with 3.03% volume rate
Figure 12: A set of nanofibers with 3.63% volume rate

Figure 13: A set of nanofibers with 4.84% volume rate
Monte Carlo simulation capable of predicting the percolation threshold of cylindrical nanotubes embedded in a polymer matrix has been developed and applied to microscale particles. The applicability of this approach to nanoscale systems is described in this section. The nanotubes are modeled as capped cylinders of diameter $D$ and length $L$ with the randomly distributed orientation angles. Numerical simulations have also been performed to determine the critical volume ratio under the different aspect ratio and the convergence of the system. As above, the simulation volume size and nanotube dimensions were rescaled to give a unit volume. Simulations were carried out by starting with an empty polymer matrix and adding nanotubes until a percolating cluster was formed. Percolation was defined as the point which two opposite walls of the cubic simulation volume were connected by a continuous cluster of nanotubes. Statistics were collected by performing 1000 independent runs.

Figs 14-17 are the results of the three dimensional cases with the aspect ratio is 10, 20, 50 and 100, respectively. We assume the same volume size and the tunneling distance equals to zero. From the results we can find the average critical volume rate converges when running the simulation no more than 400 times.
Figure 14: Average critical volume rate VS number of runs when L/D=10, Te=0

Figure 15: Average critical volume rate VS number of runs when L/D=20, Te=0
Figure 16: Average critical volume rate VS number of runs when L/D=50 and Te=0
Next, let us further discuss the convergence of Monte Carlo simulation. We use the same Monte Carlo method as mentioned above and take the following case as an example. In this case, the size of the cubic nanocomposite is \(X = Y = Z = 0.25\), the aspect ratio \(L/D = 50\), and the volume rate is 1.3%. The result shows the percolation percentage changes as the 1000 times independent runs.

Figure 17: critical volume rate VS number of runs when L/D=100 and Te=0
From Fig. 18, we found that after 350 runs, the Monte Carlo simulation converges. However, for 200 runs, the percolation percentage is 61.5% while the percentage is 57.7% for 1000 runs. The error is about 6%.

The picture below compares the volume rates under 200 runs and 1000 runs. Under the same condition, when the percolation percentage arrives to 100%, the volume rate is 1.46% for 200 runs and 1.5% for 100 runs. The error is 2.7%, which is acceptable. Therefore, in order to save simulation time, 200 runs can be used in the simulations.
4.3 Fiber aspect ratio effect

The aspect ratio has a very important effect on the percolation threshold. It was predicted in the 2D Monte Carlo study of Natsuki et al.[41] that the percolation threshold has a linear dependence on the fiber aspect ratio in a log-log plot when the aspect ratio is above 40. Bigg[8] showed in the experimental study for the 3D composites reinforced by straight short fibers the critical fiber volume rate is strongly dependent on the aspect ratio.

In the current study, the effects of the fiber aspect ratio are investigated both in 2D and 3D cases. In 2D case, the result for straight fibers with aspect ratio increasing from 50 to 200 is illustrated in Fig.20.
Figure 20: Critical volume rate versus aspect ratio in 2D case

Figure 20 shows the higher the aspect ratios, the lower the critical volume rates. From analyzing the data, if the length remains the same, lower aspect ratios represent larger diameters. It takes fewer nanofibers to reach percolation. However, the volume rate increase is proportional to the square of the increase of the nanofiber diameter. Therefore, the critical volume rates increase with the decrease of the aspect ratios.

Table 1 and Figs. 21 and 22 show the results of the volume rate under the different aspect ratios in 3D cases. The average volume rate and the standard deviation of the critical value for aspect ratio with 10, 20, 50, 100, respectively are obtained. From the following results, we found that the critical volume rate decreases as the increase of the aspect ratio.
Table 1: Aspect ratio vs critical value of fiber volume rate (3D)

<table>
<thead>
<tr>
<th>L/D</th>
<th>Average volume rate</th>
<th>Stdev</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>6.6445</td>
<td>0.4126</td>
</tr>
<tr>
<td>20</td>
<td>3.3694</td>
<td>0.1293</td>
</tr>
<tr>
<td>50</td>
<td>1.3157</td>
<td>0.0443</td>
</tr>
<tr>
<td>100</td>
<td>0.6171</td>
<td>0.0352</td>
</tr>
</tbody>
</table>

Figure 21: Aspect ratio vs critical value of fiber volume rate
In the following case, we run the Monte Carlo simulation by adding the nanotubes into the polymer matrix gradually, each time we calculate the volume rate and percolation probability percentage based on the current amount of the nanotubes. We assume all the 3D cases with the same volume size and tunneling distance is zero. From Fig.22, we can compare the percolation percentage changing with the different volume rate under the four aspect ratios, 10, 20, 50 and 100, respectively. Fig.23 shows at the threshold, where the percolation percentage just arrives to 100%, the volume rates change as the different aspect ratio, which is also called the critical volume rate. We found the same conclusion, the higher the aspect ratio, the lower the critical volume rate and the percolation networks are more easily to form.

![Percolation probability % vs Volume Rate %](image)

Figure 22: Percolation percentage vs volume rate with the aspect ratio 10, 20, 50 and 100.
4.4 Electrical conductivity

According to the percolation theory [36,44], the effective electrical conductivity of a composite, \( \sigma_e = \sigma_f (\phi - \phi_c)^t \), where \( \sigma_f \) is the electrical conductivity of the conductive phase, \( \phi \) is the volume fraction of the conductive phase, \( \phi_c \) is the critical volume fraction, and \( t \) is the conductivity exponent. It is proposed that in this theory \( t \) depends only on the space dimensionality. A universal value of \( t = 1.3 \) was showed for 2D materials, and \( t = 1.7 \sim 2.0 \) was suggested for 3D materials.

The simplest model to predict the conductivity of the composite after percolation happens is to assume the conductivity is proportional to the ratio of the number of nanofibers forming the

Figure 23: Volume rate under the different aspect ratio at 100% percolation probability
largest cluster to the total number of nanofibers. Figure 24 shows the results of a single simulation run, in which nanofibers are added to the substrate to increase the volume rate gradually. Before the volume rate reaches the critical volume rate of 2.63\%, the conductivity is zero. After that the conductivity jumps to a certain value, which increases with the increasing volume rates. After the volume rate reaches 4.84\%, almost all nanofibers are connected into one large cluster. The conductivity is a normalized value of 1.

![Figure 24: Conductivity vs volume rate in 2D case](image)

The effective electrical conductivity of the composite varying with the fiber volume fraction is shown in Fig. 25 and table 2. The results show that there is a sharp increase of the conductivity near the threshold.
Figure 25: Nanofibers in the largest cluster/total nanofibers versus volume rate

Table 2: Nanofibers in the largest cluster/total nanofibers versus volume rate

<table>
<thead>
<tr>
<th>Volume Rate</th>
<th>Nanofibers in the largest cluster/total nanofibers</th>
<th>Volume Rate</th>
<th>Nanofibers in the largest cluster/total nanofibers</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1</td>
<td>0.379</td>
<td>9.111</td>
<td>74.700</td>
</tr>
<tr>
<td>2.119</td>
<td>0.393</td>
<td>10.053</td>
<td>82.577</td>
</tr>
<tr>
<td>3.142</td>
<td>0.540</td>
<td>11.153</td>
<td>88.002</td>
</tr>
<tr>
<td>4.084</td>
<td>0.727</td>
<td>12.095</td>
<td>91.123</td>
</tr>
<tr>
<td>5.027</td>
<td>1.325</td>
<td>13.038</td>
<td>93.284</td>
</tr>
<tr>
<td>6.126</td>
<td>3.555</td>
<td>14.137</td>
<td>95.107</td>
</tr>
<tr>
<td>7.069</td>
<td>18.958</td>
<td>15.08</td>
<td>96.268</td>
</tr>
<tr>
<td>8.011</td>
<td>57.685</td>
<td>15.865</td>
<td>96.985</td>
</tr>
</tbody>
</table>
4.5 Density distribution with various aspect ratios

We run the Monte Carlo simulation under 1000 different random seeds and the following figure shows the critical percolation volume rate of each case.

Figure 26: Critical volume rates under 1000 random seeds

Fig. 27 shows the histogram of the random data in 100 bins.
Figure 27: Histogram of the random data in 100 bins

And then we use the statistics method and get the probability density estimate of the random data, which evaluates the density estimate at 100 points covering the range of the data.
By using the same method, we got the probability density of critical percolation volume rate with different aspect ratio of the nanotubes.
4.6 Tunneling Effect

In quantum mechanics, quantum tunneling (or the *tunnel effect*) is a nanoscope phenomenon in which a particle violates the principles of classical mechanics by penetrating a potential barrier or impedance higher than the kinetic energy of the particle [57]. Balberg [58] and later on Rubin et al.[59] proposed a model based on interparticle tunneling conduction. This model implies a diverging resistance distribution between spherical particles governed by a Hertz law depending on the mean interparticle distance (which decreases with the particle content). Scarisbrickz [60] has argued that in composites containing a large fraction of inclusions, the \( V-I \) relationship is
linear due to the actual touching of inclusions. Electron tunneling was proposed to be the probable mechanism of electrical conduction in such composites. However, Reboul [61] observed a linear \( V-I \) relationship in composites containing a low volume fraction of fibers, which exhibited high resistivities and where fiber-fiber contacts were improbable. In general, the tunneling distance between fibers is a parameter relative to a matrix and has a quite small value (a few nanometers) [62]. The diameter of traditional carbon fibers is very larger in comparison with the tunneling distance. The limit of tunneling distances depends on matrix used in composite system but its determination for an actually given system is quite difficult.

Therefore, in our research, we used the Monte Carlo simulation to investigate the tunneling effect on the critical volume rate of the nanofibers. Using the same simulation model and approach as we described before, we set the tunneling distance to be zero, 0.1R, 0.3R and 0.5R, where R is the radius of the capped cylinders of the nanotube model. We plot the critical volume rate distribution with the different tunneling distance under the 1000 times independent runs. As Fig. 30 shows, the tunneling effect does not affect the critical volume rate too much. As the tunneling distance changing from zero to 0.5R, the critical volume rates are almost the same.
4.7 Non-scalability of the percolation model

In this section, we intend to apply the percolation model to analyze and address percolation is not a scalable mechanism. First, we define a typical percolation model. Imagine a two dimensional lattice of dots, which are usually think as sites. The bonds, are the lines drawn between neighboring sites. Each bond can be open with the probability $p$, or closed with the probability $(1 - p)$. A cluster is formed when a group of sites connected by open bonds. We say a cluster percolates the lattice if it extends from one side of the lattice to the opposite side. From
In this example, we can easily find the average volume rate depends on the possibility $p$ and the size of the lattice. Even $p$ remains the same, the bigger the size of the lattice is, the larger the average volume rate will be. For example, for each unit grid, since there is a site occupied, we can say it percolates. However, for the whole lattice, it does not percolate due to some bonds are missing between the neighboring grid. Our Monte Carlo simulation results also show the non-scalability of the percolation model.

![Percolation and lattice](image)

**Figure 31:** Percolation and lattice [64].

From the following results, we can find as we increase the dimensions of the matrix, under the same $L$, $R$ and $L/D$, the average critical volume rate will change accordingly. The conclusion is the bigger the size of the matrix, the harder the percolation paths can form.

<table>
<thead>
<tr>
<th>$x = y = z$</th>
<th>$x/L$</th>
<th>Vol. rate for average critical percolation</th>
<th>100% percolation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.125</td>
<td>12.5</td>
<td>6.98%</td>
<td>7.55%</td>
</tr>
<tr>
<td>0.100</td>
<td>10</td>
<td>6.96%</td>
<td>7.78%</td>
</tr>
<tr>
<td>0.075</td>
<td>7.5</td>
<td>6.91%</td>
<td>7.96%</td>
</tr>
</tbody>
</table>
Figure 32: Volume rates under different dimensions with $L = 0.01$, $R = 0.0005$, $L/D = 10$

Table 4: Volume rates under different dimensions with $L = 0.02$, $R = 0.0005$, $L/D = 20$

<table>
<thead>
<tr>
<th>$x = y = z$</th>
<th>$x/L$</th>
<th>Vol. rate for average critical percolation</th>
<th>100% percolation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.200</td>
<td>10</td>
<td>3.48%</td>
<td>3.80%</td>
</tr>
<tr>
<td>0.125</td>
<td>6.25</td>
<td>3.43%</td>
<td>3.92%</td>
</tr>
<tr>
<td>0.100</td>
<td>5</td>
<td>3.42%</td>
<td>4.05%</td>
</tr>
<tr>
<td>0.075</td>
<td>3.75</td>
<td>3.36%</td>
<td>4.27%</td>
</tr>
</tbody>
</table>
Figure 33: Volume rates under different dimensions with $L = 0.02$, $R = 0.0005$, $L/D = 20$

Table 5: Volume rates under different dimensions with $L = 0.05$, $R = 0.0005$, $L/D = 50$

<table>
<thead>
<tr>
<th>$x = y = z$</th>
<th>$x/L$</th>
<th>Vol. rate for average critical percolation</th>
<th>100% percolation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.250</td>
<td>5</td>
<td>1.27%</td>
<td>1.46%</td>
</tr>
<tr>
<td>0.125</td>
<td>2.5</td>
<td>1.18%</td>
<td>1.71%</td>
</tr>
<tr>
<td>0.100</td>
<td>2</td>
<td>1.105%</td>
<td>1.78%</td>
</tr>
</tbody>
</table>
Figure 34: Volume rates under different dimensions with $L = 0.05$, $R = 0.0005$, $L/D = 50$
CHAPTER 5

CONCLUSIONS AND FUTURE WORK

This research work has performed the computational study of the qualitative characteristics of the nanocomposite percolation behavior. The results of the research have enhanced our understanding of the percolation theory and showed the model has the potential to be used in the prediction of the influence of certain parameters. It can help in reducing the number trials and errors in the experiments.

5.1 Conclusions

A Monte Carlo model for predicting the percolation threshold and electrical conductivity of the nanocomposite material is presented. The model accounts for the nanofiber aspect ratio effect by using the randomly distributed and oriented nanofibers, which have three adjustable parameters (i.e., the nanofiber length, the nanofiber diameter, and the nanofiber aspect ratio.)

The use of the Monte Carlo method leads to the determination of the percolation threshold, and by assuming the conductivity is proportional to the ratio of the number of nanofibers forming the largest cluster to the total number of nanofibers results in the prediction of the effective electrical
conductivity of the nanocomposite. We also explore the tunneling effect on the percolation volume rate and propose the percolation model is not a scalable system.

The numerical results obtained using the developed model show a relationship between the percolation threshold and the aspect ratio: the higher the nanofiber aspect ratio, the lower the threshold. These predicted trends of the percolation threshold and composite conductivity are in good agreement with existing experimental and simulation results.

5.2 Future Work

In our research, we assume the conductivity is proportional to the ratio of the number of nanofibers forming the largest cluster to the total number of nanofibers. Based on that, we get the estimate results to predict the conductivity of the percolation system. As we put more nanofibers in the system, all the nanofibers will form one biggest cluster which occupied the whole system, so the trend of the conductivity closes and normalizes to be one eventually. However, from the physics, we know, we can not add the nanofibers into the composite without any limit. On the other hand, even we say the ratio of the number of nanofibers forming the largest cluster to the total number of nanofibers is 100%, every time when we add nanofibers, the conductivity will change due to form the different electrical paths.
Therefore, in the future research, we will further study the area when the percolation percentage reaches 100% and try to find out how the conductivity changes with the increase of the number of nanofibers.
APPENDIX: MONTE CARLO SIMULATION CODE
function
[percolation_flag,percolation_cluster_size]=main_all(n_tube,random_seed,xyz,length,radius,Te_ratio)

%clear ;
%clc ;

global max_cluster idebug ;
global P5 P6 PO N U V ; % for routine dist_3D_2_segments()

%time_start=cputime;
%n_tube      = 100 ;
%max_pair    = n_tube*(n_tube-1)/2 ;
max_cluster = n_tube ;
% --------------------------------------- nano tubes -------------------
%  int    i, j, k, k1, i1, j1, i2, j2, k2, itmp, record_per_line, data_dim ;
%  double xlo, xhi, ylo, yhi, zlo, zhi, hlo, hhi, rlo, rhi,
%  int    n_pair, n_cluster, chaining ;
%  double dist, cluster_threshold ;
%  double **sc ;
%  int    *pair_lst_i, *pair_lst_j, *cluster_ID ;
%  double *pair_dist ;
%
%  int    *x_sort_lst ; --- pre-processing : sort by x-coordinates ---
%  FILE   *fp_out_pair_dist, *fp_out_tube_cluster, *fp_out_pair_cluster ;

% ----------------------- first executible statement ----------------
% ----------------------- first executible statement ----------------

xlo      = 0.0     ;
ylo      = 0.0     ;
zlo      = 0.0     ;
xhi      = xyz ;
yhi      = xyz ;
zhl      = xyz ;
hlo      = length ; % nano tubes are cylinders with random heights
hhi      = length ; % nano tubes are cylinders with random heights
rlo      = radius ; % nano tubes are cylinders with random radii
rhi      = radius ; % nano tubes are cylinders with random radii
filo     = -pi/2   ; % -pi/2 the low value in range of latitude angle
fihi     =  pi/2   ; % pi/2 the high value in range of latitude angle
theta_lo =  0.0    ; % 0.0 the low value in range of longitude angle
theta_hi = 2*pi    ; % pi*2 the high value in range of longitude angle
Te       = Te_ratio*radius; % Tunnel Effect gap

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Average_Volume_ratio = n_tube * pi * [(rlo + rhi) / 2]^2 * (hlo + hhi) / 2 / ((xhi - xlo) * (yhi - ylo) * (zhi - zlo))

%cluster_threshold = 2 * (rlo + rhi) / 2 + Te; % --- for cluster algorithm ---
wall_threshold = (rlo + rhi) / 2 + Te; % --- for peecolation ---

If the "distance_sq_between_centers" > "distance_sq_between_centers_threshold"

dist = inf and no need to calculate the distance

distance_sq_between_centers_threshold = ((hlo + hhi) / 2 + (rlo + rhi) + Te)^2;

data_dim = 3;
idebug = 0;

% /* ---------------- data structure of (line) segment centers ---------- */
% /* ------------------------------------------------------------------------------------------
%    sc(i, 1) : x-coordinate of center of tube i
%    sc(i, 2) : y-coordinate of center of tube i
%    sc(i, 3) : z-coordinate of center of tube i
%    sc(i, 4) : height of tube i
%    sc(i, 5) : radius of tube i
%    sc(i, 6) : phi, latitude angle of tube i  new version
%    sc(i, 7) : theta, the θ angle of tube i
% /* ------------------------------------------------------------------------------------------ */
pair_dist = zeros (1, max_pair);
pair_lst_i = zeros (1, max_pair);
pair_lst_j = zeros (1, max_pair);
cluster_ID = zeros (1, max_cluster);

sc = zeros (n_tube, 7);
P1 = zeros (3, 1); % column vector, 3D
P2 = zeros (3, 1); % column vector, 3D
P3 = zeros (3, 1); % column vector, 3D
P4 = zeros (3, 1); % column vector, 3D
A = zeros (3, 1); % column vector, 3D
B = zeros (3, 1); % column vector, 3D

% /* ------------------------ global for dist_3D_2_segments --------------------- */
% /* ------------------------ global for dist_3D_2_segments --------------------- */
PO = zeros (3, 1); %*--------- mid-point of P1, P2 --------- */
U = zeros (3, 1); %* orthogonal basis { U, V, N } for plane through P0 */
V = zeros (3, 1); %* orthogonal basis { U, V, N } for plane through P0 */
N = zeros (3, 1); %* orthogonal basis { U, V, N } for plane through P0 */
P5 = zeros (3, 1); %* orthogonal projection of P3, P4 onto plane thru P0 */
P6 = zeros (3, 1); %* orthogonal projection of P3, P4 onto plane thru P0 */
sc = set_tube ( idebug, data_dim, n_tube, xlo,xhi,ylo,yhi,zlo,zhi,hlo,hhi,rlo,rhi, filo,fihi, theta_lo, theta_hi, random_seed ) ;

x_sort_lst = zeros ( 1, n_tube ) ;

x_sort_lst = linear_sort ( idebug, n_tube, sc ) ;

n_pair = 0 ;

for i1 = 1 : n_tube-1
    i = x_sort_lst(i1) ;
    for j1 = i1+1 : n_tube
        j = x_sort_lst(j1) ;
        distance_sq_between_centers = (sc(i,1)-sc(j,1))^2+(sc(i,2)-sc(j,2))^2+(sc(i,3)-sc(j,3))^2;
        if distance_sq_between_centers > distance_sq_between_centers_threshold
            dist = inf;
            cluster_threshold = 0;
        else
            % Calculate the distance of the center of i1 and j1
            % If the distance is greater than the threshold, set the distance to infinity.
            % Otherwise, set the cluster threshold to 0.
        end
    end
end
% /* --- always true that angle phi is from -pi/2 to pi/2 --- */
% /* --------------------------------------------------------------- line segment 1 --- */
% /* --- P1, P2 : bottom and top disc centers on tube i --- */
cosi6=cos(sc(i,6));
cosi7=cos(sc(i,7));
sini6=sin(sc(i,6));
sini7=sin(sc(i,7));
nix=cosi6*cosi7; % x-component of unit vector of i tube
niy=cosi6*sini7; % y-component of unit vector of i tube
niz=sini6; % z-component of unit vector of i tube
half_l_i=sc(i,4)/2; % Half length of tube i
P2(1) = sc(i,1) + half_l_i*nix ;
P2(2) = sc(i,2) + half_l_i*niy ;
P2(3) = sc(i,3) + half_l_i*niz ;
P1(1) = sc(i,1)*2 - P2(1) ;
P1(2) = sc(i,2)*2 - P2(2) ;
P1(3) = sc(i,3)*2 - P2(3) ;
% /* --------------------------------------------------------------- line segment 2 --- */
% /* --- P3, P4 : bottom and top disc centers on tube j --- */
cosj6=cos(sc(j,6));
cosj7=cos(sc(j,7));
sijn6=sin(sc(j,6));
sijn7=sin(sc(j,7));
njx=cosj6*cosj7; % x-component of unit vector of i tube
njy=cosj6*sinj7; % y-component of unit vector of i tube
njz=sijn6; % z-component of unit vector of i tube
half_l_j=sc(i,4)/2; % Half length of tube i
P4(1) = sc(j,1) + half_l_j*njx ;
P4(2) = sc(j,2) + half_l_j*njy ;
P4(3) = sc(j,3) + half_l_j*njz ;
P3(1) = sc(j,1)*2 - P4(1) ;
P3(2) = sc(j,2)*2 - P4(2) ;
P3(3) = sc(j,3)*2 - P4(3) ;
% /* --------------------------------------------------------------- */
dist = dist_3D_2_segments ( idebug, P1,P2,P3,P4,A,B );
% dist_matrix(i1,j1)=dist;
cos_theta_ij=nix*njx+niy*njy+niz*njz;
cluster_threshold=sc(i,5) + sc(j,5) + Te;
%
end
if dist < cluster_threshold
    connect_matrix=[connect_matrix;[i1,j1]];
end
%        /* ---------------------- done for tube pair (i,j) ----------- */
end %/* --- inner loop over tube j1 --- */
end % /* --- outer loop over tube i1 --- */
% Kurt Lin's code starts here

[connect_1,connect_2]=find(dist_matrix < cluster_threshold);

connect_matrix(79,107)
dist_matrix(107,134)
connect_1=connect_matrix(:,1);
connect_2=connect_matrix(:,2);
n_connect_pairs=size(connect_1,1);
tube_cluster=(-1)*ones(n_tube,1);
cluster=[];
cluster_label=0;
while ~isempty(connect_matrix)
    cluster_set=connect_matrix(1,:);
    connect_matrix(1,:)=[];
    continue_flag=1;
    while continue_flag ~= 0
        size(cluster_set,2);
        %cluster_set=cluster_set
        %pause
        n_cluster_set=size(cluster_set,2);
        cluster_set_new=[];
        for i=1:n_cluster_set
            index_col_1=find(connect_matrix(:,1)==cluster_set(i));
            if ~isempty(index_col_1)
                for j=1:size(index_col_1,1)
                    if isempty(find(cluster_set==connect_matrix(index_col_1(j),2)))
                        cluster_set_new=[cluster_set_new,connect_matrix(index_col_1(j),2)];
                    end
                    connect_matrix(index_col_1,:)=[];
                    continue_1(i)=1;
                end
                else
                    continue_1(i)=0;
                end
            end
        end
        %Leftover=connect_matrix(1:3,:)
        index_col_1=find(connect_matrix(:,1)==cluster_set(i));
        if ~isempty(index_col_1)
            for j=1:size(index_col_1,1)
                if isempty(find(cluster_set==connect_matrix(index_col_1(j),2)))
                    cluster_set_new=[cluster_set_new,connect_matrix(index_col_1(j),2)];
                end
            end
        end
        connect_matrix(index_col_1,:)=[];
        continue_1(i)=1;
        else
            continue_1(i)=0;
        end
    end
    n_leftover=connect_matrix(1:3,:)
    index_col_2=find(connect_matrix(:,2)==cluster_set(i));
    if ~isempty(index_col_2)
        continue_2(i)=1;
    else
        continue_2(i)=0;
    end
%Leftover=connect_matrix(1:3,:)
index_col_2=find(connect_matrix(:,2)==cluster_set(i));
%if ~isempty(index_col_2)

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% whatifound=connect_matrix(index_col_2,2)
%end
if ~isempty(index_col_2)
    for j=1:size(index_col_2,1)
        %Leftover=connect_matrix(1:index_col_2(j),:);
        if isempty(find(cluster_set==connect_matrix(index_col_2(j),1)))
            cluster_set_new=[cluster_set_new,connect_matrix(index_col_2(j),1)];
        end
    end
end
connect_matrix(index_col_2,:)=[];
continue_2(i)=1;
else
    continue_2(i)=0;
end
end
cluster_set=[cluster_set,cluster_set_new];
%Leftover=connect_matrix(1:20,:);
continue_flag_1=sum(continue_1);
continue_flag_2=sum(continue_2);
continue_flag=continue_flag_1+continue_flag_2;
%pause
if continue_flag==0
    %n_cluster_set=size(cluster_set,2);
    cluster_label=cluster_label+1;
    %cluster=[cluster;n_cluster_set];
    %if n_cluster_set==69
    %cluster_set=cluster_set
    %cluster_label=cluster_label
    %end
    for k=1:n_cluster_set
        tube_cluster(cluster_set(k))=cluster_label;
    end
end
end
%pause
end

% Put the size of each cluster in an array
number_of_cluster=cluster_label;
for i=1:number_of_cluster
    cluster=[cluster;size(find(tube_cluster==i),1)];
end
% Find largest cluster
[largest_cluster, n_max_cluster] = max(cluster);

% Check percolation (3-D)

x_largest_cluster_min = xhi;
x_largest_cluster_max = xlo;
y_largest_cluster_min = yhi;
y_largest_cluster_max = ylo;
z_largest_cluster_min = zhi;
z_largest_cluster_max = zlo;

for i = 1:n_tube
    i_sorted = x_sort_lst(i);
    if tube_cluster(i) == n_max_cluster
        x1 = sc(i_sorted,1) - 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*cos(sc(i_sorted,7));
        if x1 < x_largest_cluster_min
            x_largest_cluster_min = x1;
        end
        if x1 > x_largest_cluster_max
            x_largest_cluster_max = x1;
        end
        x2 = sc(i_sorted,1) + 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*cos(sc(i_sorted,7));
        if x2 < x_largest_cluster_min
            x_largest_cluster_min = x2;
        end
        if x2 > x_largest_cluster_max
            x_largest_cluster_max = x2;
        end
        %output = [i_sorted, x1, x2, x_largest_cluster_max]
    end
    y1 = sc(i_sorted,2) - 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*sin(sc(i_sorted,7));
    if y1 < y_largest_cluster_min
        y_largest_cluster_min = y1;
    end
    if y1 > y_largest_cluster_max
        y_largest_cluster_max = y1;
    end
    y2 = sc(i_sorted,2) + 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*sin(sc(i_sorted,7));
    if y2 < y_largest_cluster_min
        y_largest_cluster_min = y2;
    end
    if y2 > y_largest_cluster_max
        y_largest_cluster_max = y2;
    end
    z1 = sc(i_sorted,3) - 0.5*sc(i_sorted,4)*sin(sc(i_sorted,6));
    if z1 < z_largest_cluster_min
        z_largest_cluster_min = z1;
    end
    if z1 > z_largest_cluster_max
        z_largest_cluster_max = z1;
    end
if \( z_1 > z_{\text{largest\_cluster\_max}} \)
\[
z_{\text{largest\_cluster\_max}} = z_1;
\]
end
\( z_2 = sc(i_{\text{sorted}},3) + 0.5 \times sc(i_{\text{sorted}},4) \times \sin(sc(i_{\text{sorted}},6)) \);
if \( z_2 < z_{\text{largest\_cluster\_min}} \)
\[
z_{\text{largest\_cluster\_min}} = z_2;
\]
end
if \( z_2 > z_{\text{largest\_cluster\_max}} \)
\[
z_{\text{largest\_cluster\_max}} = z_2;
\]
end
end

end

\( x_{\text{percolation}} = 0; \)
if \( (x_{\text{largest\_cluster\_max}} > (xhi-\text{wall\_threshold})) \& (x_{\text{largest\_cluster\_min}} < (xlo+\text{wall\_threshold})) \)
\[
x_{\text{percolation}} = 1;
\]
end
\( y_{\text{percolation}} = 0; \)
if \( (y_{\text{largest\_cluster\_max}} > (yhi-\text{wall\_threshold})) \& (y_{\text{largest\_cluster\_min}} < (ylo+\text{wall\_threshold})) \)
\[
y_{\text{percolation}} = 1;
\]
end
\( z_{\text{percolation}} = 0; \)
if \( (z_{\text{largest\_cluster\_max}} > (zhi-\text{wall\_threshold})) \& (z_{\text{largest\_cluster\_min}} < (zlo+\text{wall\_threshold})) \)
\[
z_{\text{percolation}} = 1;
\]
end

percolation\_flag = zeros(1,3);
percolation\_cluster\_size = zeros(1,3);
percolation\_cluster\_size(1) = largest\_cluster;
if x\_percolation == 1 | y\_percolation == 1 | z\_percolation == 1
\%
if x\_percolation == 1 | y\_percolation == 1 (For thin plate, only x and y are considered)
    percolation\_flag(1) = 1;
\%
    percolation\_cluster\_size(1) = largest\_cluster;
end
% Find 2nd largest cluster
cluster\_takeaway\_max = cluster;
cluster\_takeaway\_max(n\_max\_cluster) = 0;
[second\_largest\_cluster, n\_second\_max\_cluster] = max(cluster\_takeaway\_max);
% Check percolation
x\_2nd\_largest\_cluster\_min = xhi;
x\_2nd\_largest\_cluster\_max = xlo;
y\_2nd\_largest\_cluster\_min = yhi;
y\_2nd\_largest\_cluster\_max = ylo;
z_2nd_largest_cluster_min=zhi;
z_2nd_largest_cluster_max=zlo;
for i=1:n_tube
    i_sorted=x_sort_lst(i);
    if tube_cluster(i)==n_second_max_cluster
        x1=sc(i_sorted,1) - 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*cos(sc(i_sorted,7));
        if x1 < x_2nd_largest_cluster_min
            x_2nd_largest_cluster_min=x1;
        end
        if x1 > x_2nd_largest_cluster_max
            x_2nd_largest_cluster_max=x1;
        end
        x2=sc(i_sorted,1) + 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*cos(sc(i_sorted,7));
        if x2 < x_2nd_largest_cluster_min
            x_2nd_largest_cluster_min=x2;
        end
        if x2 > x_2nd_largest_cluster_max
            x_2nd_largest_cluster_max=x2;
        end
    end
    %output=[i_sorted,x1,x2,x_2nd_largest_cluster_max]
    %pause
    y1=sc(i_sorted,2) - 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*sin(sc(i_sorted,7));
    if y1 < y_2nd_largest_cluster_min
        y_2nd_largest_cluster_min=y1;
    end
    if y1 > y_2nd_largest_cluster_max
        y_2nd_largest_cluster_max=y1;
    end
    y2=sc(i_sorted,2) + 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*sin(sc(i_sorted,7));
    if y2 < y_2nd_largest_cluster_min
        y_2nd_largest_cluster_min=y2;
    end
    if y2 > y_2nd_largest_cluster_max
        y_2nd_largest_cluster_max=y2;
    end
    z1=sc(i_sorted,3) - 0.5*sc(i_sorted,4)*sin(sc(i_sorted,6));
    if z1 < z_2nd_largest_cluster_min
        z_2nd_largest_cluster_min=z1;
    end
    if z1 > z_2nd_largest_cluster_max
        z_2nd_largest_cluster_max=z1;
    end
    z2=sc(i_sorted,3) + 0.5*sc(i_sorted,4)*sin(sc(i_sorted,6));
    if z2 < z_2nd_largest_cluster_min
        z_2nd_largest_cluster_min=z2;
    end
    z_2nd_largest_cluster_max
end
if z2 > z_2nd_largest_cluster_max
   z_2nd_largest_cluster_max = z2;
end
end
end
x_percolation = 0;
if (x_2nd_largest_cluster_max > (xhi-wall_threshold)) & (x_2nd_largest_cluster_min < (xlo+wall_threshold))
   x_percolation = 1;
end
y_percolation = 0;
if (y_2nd_largest_cluster_max > (yhi-wall_threshold)) & (y_2nd_largest_cluster_min < (ylo+wall_threshold))
   y_percolation = 1;
end
z_percolation = 0;
if (z_2nd_largest_cluster_max > (zhi-wall_threshold)) & (z_2nd_largest_cluster_min < (zlo+wall_threshold))
   z_percolation = 1;
end
% percolation_cluster_size(2)=second_largest_cluster;
if x_percolation==1|y_percolation==1|z_percolation==1
    percolation_flag(2)=1;
end
% Find 3rd largest cluster
cluster_takeaway_max(n_second_max_cluster)=0;
[third_largest_cluster,n_third_max_cluster]=max(cluster_takeaway_max);
% Check percolation
x_3rd_largest_cluster_min=xhi;
x_3rd_largest_cluster_max=xlo;
y_3rd_largest_cluster_min=yhi;
y_3rd_largest_cluster_max=ylo;
z_3rd_largest_cluster_min=zhi;
z_3rd_largest_cluster_max=zlo;
for i=1:n_tube
    i_sorted=x_sort_lst(i);
    if tube_cluster(i)==n_third_max_cluster
        x1=sc(i_sorted,1) - 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*cos(sc(i_sorted,7));
        if x1 < x_3rd_largest_cluster_min
            x_3rd_largest_cluster_min=x1;
        end
end
if x1 > x_3rd_largest_cluster_max
    x_3rd_largest_cluster_max=x1;
end

x2=sc(i_sorted,1) + 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*cos(sc(i_sorted,7));
if x2 < x_3rd_largest_cluster_min
    x_3rd_largest_cluster_min=x2;
end
if x2 > x_3rd_largest_cluster_max
    x_3rd_largest_cluster_max=x2;
end

%output=[i_sorted,x1,x2,x_3rd_largest_cluster_max]
%pause

y1=sc(i_sorted,2) - 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*sin(sc(i_sorted,7));
if y1 < y_3rd_largest_cluster_min
    y_3rd_largest_cluster_min=y1;
end
if y1 > y_3rd_largest_cluster_max
    y_3rd_largest_cluster_max=y1;
end

y2=sc(i_sorted,2) + 0.5*sc(i_sorted,4)*cos(sc(i_sorted,6))*sin(sc(i_sorted,7));
if y2 < y_3rd_largest_cluster_min
    y_3rd_largest_cluster_min=y2;
end
if y2 > y_3rd_largest_cluster_max
    y_3rd_largest_cluster_max=y2;
end

z1=sc(i_sorted,3) - 0.5*sc(i_sorted,4)*sin(sc(i_sorted,6));
if z1 < z_3rd_largest_cluster_min
    z_3rd_largest_cluster_min=z1;
end
if z1 > z_3rd_largest_cluster_max
    z_3rd_largest_cluster_max=z1;
end

z2=sc(i_sorted,3) + 0.5*sc(i_sorted,4)*sin(sc(i_sorted,6));
if z2 < z_3rd_largest_cluster_min
    z_3rd_largest_cluster_min=z2;
end
if z2 > z_3rd_largest_cluster_max
    z_3rd_largest_cluster_max=z2;
end

x_percolation=0;
if (x_3rd_largest_cluster_max > (xhi-wall_threshold))&(x_3rd_largest_cluster_min < (xlo+wall_threshold))

x_percolation=1;
end
y_percolation=0;
if (y_3rd_largest_cluster_max > (yhi-wall_threshold))&(y_3rd_largest_cluster_min < (ylo+wall_threshold))
    y_percolation=1;
end
z_percolation=0;
if (z_3rd_largest_cluster_max > (zhi-wall_threshold))&(z_3rd_largest_cluster_min < (zlo+wall_threshold))
    z_percolation=1;
end

% percolation_cluster_size(3)=third_largest_cluster;
if x_percolation==1|y_percolation==1|z_percolation==1
    percolation_flag(3)=1;
end

fid=fopen('main_all_output.txt','a');

fprintf(fid,'n_tube = %6d random_seed = %6d
',n_tube,random_seed);
fclose(fid);

return;

function sc = set_tube ( idebug, data_dim, n_tube, xlo, xhi, ylo, yhi, zlo, zhi, hlo, hhi, rlo, rhi, filo, fihi, theta_lo, theta_hi, random_seed )
    xlo, xhi, ylo, yhi, zlo, zhi, hlo, hhi, rlo, rhi, filo, fihi, theta_lo, theta_hi, random_seed)

% float ran4(long *idum);
% long idum;
% FILE *fp_out_tube;

% fprintf('n To check_and_remove tubes extending out in six directions\n\n');
% fprintf(' 1 : yes, Else ( nonzero ) not, \n\n');
% fprintf('t Answer in condensed way, as follows : \n\n');
% fprintf(' 111111 : yes, in all six directions ; \n\n');
% fprintf(' 222222 : No, in none of the six directions ; \n\n');
% fprintf(' 121212 : make sure  0 < x   0 < y   0 < z   \n\n');
% fprintf(' 212121 : make sure   x < 1   y < 1   z < 1 \n\n');
fprintf(' 112222 : make sure 0 < x < 1  
');
chk = input('Enter a six-digit number to check some, check all, check none : 
');
chk=222222;

rand('twister',random_seed); % Set random seed
sc = zeros(n_tube, 7);

% ----------------------------------- examine the random data ----------
for i = 1 : n_tube
i_random = 0;
chk_valid = 0;
sc(i,:) = rand(1,7);
while ( chk_valid ~= 1 )
i_random = i_random + 1;
for j = 1 : 7
if ( sc(i,j) < 0.0)
    fprintf(' sc(%d,%d) = %e ?
',i,j,sc(i,j));pause
end
if ( sc(i,j) > 1.0)
    fprintf(' sc(%d,%d) = %e ?
',i,j,sc(i,j));pause
end
end
sc(i,1) = xlo      + ( xhi      - xlo     )*sc(i,1) ;
sc(i,2) = ylo      + ( yhi      - ylo     )*sc(i,2) ;
sc(i,3) = zlo      + ( zhi      - zlo     )*sc(i,3) ;
sc(i,4) = hlo      + ( hhi      - hlo     )*sc(i,4) ;
sc(i,5) = rlo      + ( rhi      - rlo     )*sc(i,5) ;
sc(i,6) = filo     + ( fihi     - filo    )*sc(i,6) ; % latitude angle
sc(i,7) = theta_lo + ( theta_hi - theta_lo)*sc(i,7) ; % longitude angle
end
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if ( data_dim == 2 )
    sc(i,3) = zlo ;
    \% old sc(i,6) = pi/2 ;
    sc(i,6) = 0.0 ; \% latitude angle
end

\% ------------------------ must pause to check ---------------------
\% ------------------------ must pause to check ---------------------
chk_valid = 1 ;

if ( chk_valid < 1 )
    sc(i,1:7) = rand(1,7) ;
end
end % --- end while ---
end % --- end while ---

return ;

\% ------------------- current version : linear sort ---

function x_sort_lst = linear_sort ( idebug, n_tube, sc )

i_select = 0 ;
mark_lst = zeros ( 1, n_tube ) ;

fprintf('
 --- Start sorting by x-coordinates ... \n') ;

for i_least = 1 : n_tube
    \% -------- assign an initial candidate, then compare and update ---
    for k = 1 : n_tube
        if ( mark_lst(k) == 0 )
            i_select = k ;
            break ;
        end
    end
    for j = 1 : n_tube
        if ( mark_lst(j) == 0 )
            if ( sc(j,1) < sc(i_select,1) )
                i_select = j ;
            end
        end
    end
end
x_sort_lst(i_least) = i_select;
mark_lst(i_select) = 1;
end

return;

%* --------------------------------------------------------------------
% Input:
% Four points in space
% P1(x1,y1,z1),
% P2(x2,y2,z2),
% P3(x3,y3,z3),
% P4(x4,y4,z4),
% Output: the distance between two line segments P1_P2, P3_P4,
% --------------------------------------------------------------- */

function dist_3 = dist_3D_2_segments ( idebug, P1, P2, P3, P4, A, B )

% int i, k, itmp, ierr;
% double p, q, r, s, ax, ay, bx, by, dist_2, l_half, dist_common, t3,t4, tmp;
% double ratio_1, ratio_2;

% */* ----------------------------------------------------- */
% PO : mid-point of segment P1_P2
% H : the plane passing through PO with normal N[]
% P5 : orthogonal projection of P3 onto H, with relative coordinate (p,q)
% P6 : orthogonal projection of P4 onto H, with relative coordinate (r,s)
% U : orthogonal basis { U, V, N } for plane through P0
% V : orthogonal basis { U, V, N } for plane through P0
% N : orthogonal basis { U, V, N } for plane through P0
% --------------------------------------------------------------- */

% /* ------- global allocated in main() ------- */
% /* ------- global allocated in main() ------- */

% /* ----------------------------------------------------- mid-point of P1, P2 --- */
PO = ( P1 + P2 ) / 2;

P1 = P1 - PO ; % local coordinate: shift origin to P0
P2 = P2 - PO ; % local coordinate: shift origin to P0
P3 = P3 - PO ; % local coordinate: shift origin to P0
P4 = P4 - PO ; % local coordinate: shift origin to P0
U = P2 ; % U = (P2-P1)/2
tmp = sqrt((U')*U);
U = U /tmp ;  % now a unit vector, done
1_half = tmp ;  % half length of P1_P2
V = P4 - P3 ;
tmp = sqrt ( (V' )*V ) ;
V = V /tmp ;  % now a unit vector, done

N = tensor_3D(U,V) ;   % N = P2_P1 x P3_P4 as normal to plane through P1
tmp = sqrt ( (N'*N ) ) ;
if ( tmp > 1.0e-12 )
    N = N /tmp ;  % now a unit vector, done
end

V = tensor_3D( N, U ) ;  % (re)generate V
tmp = sqrt ( (V'*V) ;
V = V /tmp ;  % now a unit vector, done

% ---------------------------- always check -------
t3 = (N'*P3 ) ;
t4 = (N'*P4 ) ;
% ------------------------------------------ always check ------------
t3 = 0.5*( t3 + t4 ) ;
t4 = t3 ;
dist_common = abs(t3) ;
P5 = P3 - t3*N ;  % P5-PO = P3-PO - t*N
P6 = P4 - t4*N ;  % P6-PO = P4-PO - t*N

% ------------------------------------------- always check ------------
[p q r s] = dist_2D_2_segments ( idebug, 1_half, p, q, r, s ) ;
dist_3 = sqrt ( dist_2*dist_2 + dist_common*dist_common ) ;

%悟atorium             bug here ---
B = PO + bx*U + by*V + t3*N ;
A = PO + ax*U + ay*V ;
%悟atorium             missing was below ---
P1 = P1 + PO ;
P2 = P2 + PO ;
P3 = P3 + PO ;
P4 = P4 + PO ;

%  ------------------------------- always check --------------
%  ------------------------------- always check --------------

return ;

%  ---------------------------------------------------------------------
%  ------------------------------ 2D configuration ---------------------
%  ---------------------------------------------------------------------

function [ax ay bx by dist_2] = dist_2D_2_segments ( idebug, l_half, p,q,r,s )

% /*----------------------------------------------------------------------
% Reduction : P1 = ( -l_half, 0 ) , P2 = ( l_half, 0 ), 
% P5 = ( p/l_half, q/l_half ), P6 = ( r/l_half, s/l_half )

p_sav = p ;
q_sav = q ;
r_sav = r ;
s_sav = s ;

p = p / l_half ;
q = q / l_half ;
r = r / l_half ;

% OK for C, passed by value
% OK for C, passed by value
% OK for C, passed by value

s = s / l_half; % OK for C, passed by value

if ( ( p >= 1.0 ) && ( r >= 1.0 ) )
    subcase = 10;
elseif ( ( p <= -1.0 ) && ( r <= -1.0 ) )
    subcase = 20;
elseif ( ( abs(p) <= 1.0 ) && ( abs(r) <= 1.0 ) )
    subcase = 30;
elseif ( ( min(p,r) < 1.0 ) && ( 1.0 < max(p,r) ) )
    subcase = 40;
elseif ( ( min(p,r) < -1.0 ) && ( -1.0 < max(p,r) ) )
    subcase = 50;
else
    subcase = -1;
pause
end

switch ( subcase )

    case 10 % --- P5, P6 are both to the right of line x=1 ---
        % ----- A = P2 */
        ax = 1.0;
        ay = 0.0;
        [bx by dist_2] = dist_2D_pt_to_segment ( 1.0, 0.0, p, q, r, s ) ;

    case 20 % --- P5, P6 are both to the left of line x=-1 ---
        % ----- A = P1 */
        ax = -1.0;
        ay = 0.0;
        [bx by dist_2] = dist_2D_pt_to_segment (-1.0, 0.0, p, q, r, s ) ;

    case 30 % --- P5, P6 are both between lines x=1, x=-1 ---
        % --- recall q < s, always --- *
        if ( 0.0 < q )
            % --- (p,q)=P5 is closest to segment P1_P2, on plane H ---
            % --- A = (p,0), B = P5 = (p,q) --- */
            ax = p;
            ay = 0.0;
            bx = p;
            by = q;
            dist_2 = q ;

        elseif ( s < 0.0 )
            % --- (r,s)=P6 is closest to segment P1_P2, on plane H ---
            % --- A = (r,0), B = P6 = (r,s) --- */

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ax = r;
ay = 0.0;
bx = r;
by = s;
dist_2 = -s;

else
  ay = 0.0;  % A is on P1_P2
  ax = (p*s-q*r) / (s - q);  % some math
  bx = (ax);  % B = A
  by = 0.0;  % B = A
  dist_2 = 0.0;
end

case 40  % --- segment P5_P6 intersects with line x=1, and also x=-1 ?
  [ax ay bx by dist_2] = dist_2D_case_40 (p, q, r, s);

case 50  % --- segment P5_P6 intersects with line x=-1, and also x=1 ?
  % --- apply reflection wrt x=0 and appeal to case 40 ---
  [ax ay bx by dist_2] = dist_2D_case_40 (-p, q, -r, s);
  ax = -ax;
  bx = -bx;
end

%  --------------------------------- re-scale ----------------
ax = ax*l_half;
ay = ay*l_half;
bx = bx*l_half;
by = by*l_half;
dist_2 = dist_2*l_half;

% /* ---------------------------------- re-scale --------------------
% %  prefer the next block, for accuracy
% % p *= l_half;
% % q *= l_half;
% % r *= l_half;
% % s *= l_half;
% % ----------------------------------
% p = p_sav;
% q = q_sav;
% r = r_sav;

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s = _sav ;  

% /* --------------------------------------------- always check ----- 

return ;

function N = tensor_3D ( U, V )

N(3,1) = U(1,1)*V(2,1) - U(2,1)*V(1,1) ;
N(1,1) = U(2,1)*V(3,1) - U(3,1)*V(2,1) ;
N(2,1) = U(3,1)*V(1,1) - U(1,1)*V(3,1) ;

return ;

% --------------------------------------------------------------------
% ---------------------------- 2D configuration ---------------------
% --------------------------------------------------------------------
% Input     : point P and line segment A_B, with 
% P = ( px, py )
% A = ( ax, ay )
% B = ( bx, by )
% Output    : distance from  P  to  A_B

% Algorithm : transform to case P=(0,0), ... for code efficiency
% Reduction : P = (0,0), A = (ax-px,ay-py), B = (bx-px,by-py)
% Projection of P(0,0) to line A_B with A(ax,ay) B(bx,by) is Q(qx,qy), by

% choosing   n = (nx,ny), a unit normal to line A_B
% then 
% dist = | nx*ax+ny*ay | or | nx*bx+ny*by | 
% and
% qx = nx/(nx*nx+ny*ny)*(nx*ax+ny*ay) ;
% qy = ny/(nx*nx+ny*ny)*(nx*ax+ny*ay) ;
% transform back
% ( qx, qy ) += ( px, py )
% --------------------------------------------------------------------

function [qx qy dist] = dist_2D_pt_to_segment ( px, py, ax, ay, bx, by )

global idebug ;

% int i, itmp, numerical_chk, local_debug, ierr ;
%  double nx, ny, ab_x, ab_y, tmp1, tmp2, x_chk, y_chk;

%  ---------------------------------------------------------------
local_debug = 0;

ab_x = bx - ax;
ab_y = by - ay;

ax = ax - px; % OK for C, passed by value
ay = ay - py; % OK for C, passed by value
bx = bx - px; % OK for C, passed by value
by = by - py; % OK for C, passed by value

%  ---------------------------------------------------------------------
dist = sqrt ( ab_x*ab_x + ab_y*ab_y ); % temporary usage
nx = -ab_y/dist;
y = ab_x/dist;

%  ------- bug was below ---
%  ------ bug was below ---
if ( tmp1 >= ( 0.0 - 1.0e-10 ) )
%  --- exterior angle(A-P,B-A) <= 90, interior >= 90, then Q = A ---
dist = sqrt ( ax*ax + ay*ay );
qx = ax + px;
yq = ay + py;
else %  ----------- angle(A-P,B-A) <= 90 ---
tmp2 = bx*ab_x + by*ab_y; % < B-P, B-A >
if ( tmp2 <= ( 0.0 + 1.0e-10 ) )
%  --- angle(B-P,B-A) >= 90_degree, then Q = B ---
dist = sqrt ( bx*bx + by*by );
qx = bx + px;
yq = by + py;
else
%  --- triangle PAB is acute
%  ------------------------------- case : Q is bewteen A, B ---
dist = nx*ax + ny*ay; % temporary usage for < n, a >
qx = nx*(dist) + px;
yq = ny*(dist) + py;
dist = abs(dist);

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function [ax ay bx by dist_2] = dist_2D_case_40 ( p, q, r, s )

global idebug ;
if ( ( r < 1.0 ) && ( 1.0 < p ) && ( 0.0 < q ) )
    %* --- case 4a ------- segment P5_P6 intersects with line x=1 ---
    %*----------------------------------------------------------------------
    % dist ( P1_P2, P5_P6 ) = dist ( P1_P2, P5_Bstar )
    % = dist ( P2, P5_Bstar )
    %----------------------------------------------------------------------
    subcase = 41 ;
    b_star = ( (p-1)*s + (1-r)*q ) / (p-r) ;
    ax = 1.0 ; % i.e., A = P2
    ay = 0.0 ;
    [bx by dist_2] = dist_2D_pt_to_segment (1.0,0.0,p,q,1.0,b_star) ;
elseif ( ( p < 1.0 ) && ( 1.0 < r ) && ( s < 0.0 ) )
    % --- recall q < s, always ---
    % --- case 4b ------- segment P5_P6 intersects with line x=1 ---
% -----------------------------
%  dist ( P1_P2, P5_P6 ) = dist ( P1_P2, Bstar_P6 )
%  = dist ( P2, Bstar_P6 )
% -----------------------------
subcase = 42 ;
% bug b_star = ((1-p)*s - (r-1)*q) / (r-p);
b_star = ((1-p)*s + (r-1)*q) / (r-p);
ax = 1.0 ; % i.e., A = P2
ay = 0.0 ;
[ bx by dist_2 ] = dist_2D_pt_to_segment (1.0, 0.0, 1.0, b_star, r, s) ;

elseif ( ( p < 1.0 ) && ( 1.0 < r ) && ( 0.0 < q ) )
% --- case 4b ------- segment P5_P6 intersects with line x= 1 ---
% ----------------------
%  dist ( P1_P2, Bstar_P6 ) = dist ( P2, Bstar_P6 )
% s.t. dist ( P1_P2, P5_P6 ) = dist ( P1_P2, P5_Bstar ) OK
%      = dist ( P1_P2, P5 ) wrong
%    i.e., dist ( A , B ) wrong
%          with A=(p,0) B=P5 wrong
% ----------------------
% ------------------- was bug --------
if ( p >= -1.0 )
  subcase = 43 ;
  % --------------- A=(p,0) B=P5 ---
  bx = p ;
  by = q ;
  ax = bx ;
  ay = 0.0 ;
  dist_2 = by ;
else
  subcase = 44 ;
  % --------------- A=P1, B=? ---
  ax = -1.0 ;
  ay = 0.0 ;
  b_star = q + ( s - q ) / ( r - p ) * ( -1.0 - p ) ;
  [ bx by dist_2 ] = dist_2D_pt_to_segment(-1.0, 0.0, p, q, -1.0, b_star);
end

elseif ( ( p > 1.0 ) && ( 1.0 > r ) && ( s < 0.0 ) )
% --- case 4g ------- segment P5_P6 intersects with line x= 1 ---
% -------------------------
%  dist ( P1_P2, P5_P6 ) = dist ( P1_P2, Bstar_P6 ) OK
%    = dist ( P1_P2, P6 ) wrong
%   with B = P6, A = (r,0) wrong
% -------------------------
%* ---------------- was bug ---------
if ( r >= -1.0 )
    subcase = 45 ;
    bx = r ;
    by = s ;
    ax = bx ;
    ay = 0.0 ;
    dist_2 = -by ;
else
    subcase = 46 ;
    ax = -1.0 ;
    ay = 0.0 ;
    b_star = q + ( s - q ) / ( r - p ) * ( -1.0 - p ) ;
    [bx by dist_2] = dist_2D_pt_to_segment(-1.0,0.0,-1.0,b_star,r,s);
end

elseif ( ( r < 1.0 ) && ( 1.0 < p ) && ( q < 0.0 ) && ( 0.0 < s ) )

% ---------------- bug was here ----- 
% solve for the intersect (*,0) of the line (p,q)_(r,s) and line y=0 
ax = p - q*(r-p)/(s-q) ; %* may be of temporary storage
if ( ax > 1.0 )
% *------------------------------------------------------------------
%  case 4c : P5_P6 intersects with both lines x=1, y=0, but not P1_P2
% dist( P1_P2, P5_P6 ) = dist( P2, P5_P6 ), with 
% A = P2, and B in interior of P5_P6, in quadrant I 
% *------------------------------------------------------------------
    subcase = 47 ;
    ax = 1.0 ;
    ay = 0.0 ;
    n1 = (q-s)/sqrt((r-p)*(r-p)+(s-q)*(s-q)) ;
    n2 = (r-p)/sqrt((r-p)*(r-p)+(s-q)*(s-q)) ;
    dist_2 = n1*(p-1.0)+n2*(q-0.0) ;
    bx = ax + dist_2*n1 ;
    by = ay + dist_2*n2 ;
    dist_2 = abs(dist_2) ;
elseif ( ( -1.0 <= ax ) && ( ax <= 1.0 ) )
% *------------------------------------------------------------------
% case 4e : P5_P6 intersects with both lines x=1, y=0, also P1_P2 
% dist( P1_P2, P5_P6 ) = dist( A, B ) = 0, since A = B 
% *------------------------------------------------------------------
    subcase = 48 ;
    dist_2 = 0.0 ;
ay = 0.0
bx = ax
by = 0.0

elseif ( ax < -1.0 )
  % case 4+: P5_P6 intersects with both lines x=1, y=0, but not P1_P2
  % dist( P1_P2, P5_P6 ) = dist( P1, p5_P6 ), with
  % A = P1, and B in interior of P5_P6, in quadrant III
  % --------------------------------------------------------------
  subcase = 49;
  ax = -1.0
  ay = 0.0;
  n1 = (q-s)/sqrt((r-p)*(r-p)+(s-q)*(s-q));
  n2 = (r-p)/sqrt((r-p)*(r-p)+(s-q)*(s-q));
  dist_2 = n1*(p-(-1.0))+n2*(q-0.0);
  bx = ax + dist_2*n1;
  by = ay + dist_2*n2;
  dist_2 = abs(dist_2);
end

elseif ( ( p < 1.0 ) && ( 1.0 < r ) && ( q < 0.0 ) && ( 0.0 < s ) )
  % ------------------------------------ bug was here ----------------
  % solve for the intersect (*,0) of the line (p,q)_(r,s) and line y=0
  ax = p - q*(r-p)/(s-q);          % may be of temporary storage
if ( ax > 1.0 )
  % case 4f: P5_P6 intersects with both lines x=1, y=0, but not P1_P2
  % dist( P1_P2, P5_P6 ) = dist( P2, p5_P6 ), with
  % A = P2, and B in interior of P5_P6, in quadrant IV
  % --------------------------------------------------------------
  subcase = 50;
  ax = 1.0;
  ay = 0.0;
  n1 = (q-s)/sqrt((r-p)*(r-p)+(s-q)*(s-q));
  n2 = (r-p)/sqrt((r-p)*(r-p)+(s-q)*(s-q));
  dist_2 = n1*(p-1.0)+n2*(q-0.0);
  bx = ax + dist_2*n1;
  by = ay + dist_2*n2;
  dist_2 = abs(dist_2);
elseif ( ( -1.0 <= ax ) && ( ax <= 1.0 ) )

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% case 4d : P5_P6 intersects with both lines x=1, y=0, also the segment P1_P2
% such that
% dist( P1_P2, P5_P6 ) = dist( A, B )
% with (A,B) in segment P1_P2,

subcase = 51;
dist_2 = 0.0;
ay = 0.0;
bx = ax;
by = 0.0;

elseif ( ax < -1.0 )

% case 4+ : P5_P6 intersects with both lines x=1, y=0, but not P1_P2
% dist( P1_P2, P5_P6 ) = dist( P1, p5_P6 ), with A = P1, and B in interior of P5_P6, in quadrant II

subcase = 52;
ax = -1.0;
ay = 0.0;
n1 = (q-s)/sqrt((r-p)*(r-p)+(s-q)*(s-q));
n2 = (r-p)/sqrt((r-p)*(r-p)+(s-q)*(s-q));
dist_2 = n1*(p-(-1.0))+n2*(q-0.0);
bx = ax + dist_2*n1;
by = ay + dist_2*n2;
dist_2 = abs(dist_2);

end
end

return;
REFERENCES


Hansen, G., 2005, High aspect ratio sub-micron and nano-scale metal filaments, SAMPE J. 41. 24-33.


