THE STRUCTURAL INTEGRITY OF NANOCLAY FILLED EPOXY POLYMER UNDER CYCLIC LOADING

Submitted in fulfillment of the requirements of the degree of

Master of Engineering: Mechanical Engineering
In the Faculty of Engineering and the Built Environment at the

Durban University of Technology

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April 2017
ABSTRACT

Fatigue crack initiation and propagation behaviour of CFRP have been of great importance because such composites are often used in engineering components that are subjected to continuous cyclic loading. The objective of this thesis work was to investigate the damage characteristics of the fatigue properties of CFRP composites by the modification of the polymer matrix with nanoclay addition. Carbon fibre reinforced epoxy was produced via vacuum assisted resin infusion moulding method (VARIM) with nanoclay concentrations of 0wt%, 1wt%, 3wt% and 5wt%. Tension-tension fatigue tests were conducted at loading levels of 90%, 75% and 60%. The frequency that was used was 3Hz with R value of 0.1. The results showed that at nanoclay percentages of 0wt%, 1wt% and 3wt% there was a consistent trend, where the number of cycles increased in fatigue loading percentages of 90%, 75% and 60%. At 5wt% nanoclay percentage the number of fatigue cycles dropped significantly at the 90% fatigue loading. The brittle nature of the 5wt% laminate became dominate and the sample fractured early at low fatigue cycle numbers. At the 75% fatigue loading, the number of cycles increased and at 60% fatigue loading the 5wt% nanoclay sample exceeded the number of cycles of all the nanoclay percentages by 194%. This was due to the intercalated arrangement of the nanoclays favouring the slow rate of surface temperature increase, during fatigue testing, at low fatigue cycle loading. The Crack Density analysis was performed and showed that at the same time in the fatigue cycle life, the 1wt% had 55 cracks, 3wt% had 52 cracks and the 5wt% had 50 cracks, for the 60% fatigue loading. This proved that it took longer for the cracks to initiate and propagate through the sample as the nanoclay percentage increased. Impact and hardness testing showed that the 5wt% exhibited brittle behaviour, which contributed to the results above. Scanning electron microscopy examination highlighted that the agglomeration of nanoclays delayed the crack initiation and propagation through the specimen and that the extent of fatigue damage
decreased as the nanoclay percentage increased. A fatigue failure matrix was developed and showed that delamination, fibre breakage and matrix failure were the predominate causes for the fatigue failure.
DECLARATION

I hereby declare that the work described in this thesis is my original work and has not previously been submitted in its entirety or in part for a degree in any other university. I further declare that this work does not infringe or violate the right of others, as all the sources cited or quoted are indicated and acknowledged by means of a comprehensive list of references.

Sathievelli Chetty
ACKNOWLEDGEMENTS

I would like to express my sincere appreciation and gratitude to the following individuals and institutions that formed an integral part of the successful completion of this research.

Prof Krishnan Kanny for the supervisory role that he played throughout the course of the research period. He has provided some guidance, assistance and motivation throughout the duration of this study. His critical appraisals and suggestions have been welcomed at every phase of the study.

Dr Mohan Turup Pandurangan for co-supervising this work and for providing excellent guidance at every step of the way. His knowledgeable hands-on feedback was insightful and accelerated the process. His persistence and perseverance ensured that a steady progress was maintained that lead to the successful submission of this thesis. He was a source of encouragement and motivation when spirits were low, for that I am very grateful.

My mother, Mrs. P Chetty, my family and my little angel and savior, my son, Lolan who kept me sane through this whole process, who gave me the strength to forge ahead every day. His enthusiasm for life and excitement for my research work kept me headstrong, motivated me and never allowed me to give up. I thank them all for the support and warm encouragement they offered throughout the years. Thank you for believing in me. You are simply the best. I love you all.

Mr. Avinash Ramsuroop for his assistance with writing, testing, advice and overall running of the department. Assisting with procedures in place, procuring items and sometimes going the extra mile by collecting items required personally himself. Keeping me on the right track, keeping me positive and motivating me to be the best that I can be. For showing me that hard work and dedication pays off and is rewarded.

Dr Vimla Paul for guidance and assistance in writing and motivation to never give up and hard work will be rewarded. For the long talks and apt advice and
friendship offered and for always believing in me. Your critical suggestions and constructive criticism further invigorated my drive towards reaching the optimum of this study.

I would like to thank Vishnu Kribagaran Moodley (Chemical Engineering Department) for assistance with consumables, advise and general assistance, Ravi Veerasamy (Manufacturing Workshop) for use of the equipment at the workshop, i.e., the Drill, sander, polisher, band saw, handsaw and general consumables and the Technology Station for use of the oven, band saw, sander, angle grinder, Barcol meter and the CNC machine. All this assistance minimized delays and ensured that the samples created were free of defect and at the highest level of accuracy.

Kentron (Denel Dynamics) and the National Research Foundation (NRF) for providing the funding for the research project. This scholarship assisted in making a dream come true to pursue and attain this qualification.

My fellow diverse team members, for the constant guidance and motivation and help with keeping me grounded. For assistance with no hesitation and for always having encouraging words. I say a BIG thank you to you all.
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<th>Description</th>
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<tbody>
<tr>
<td>VHCF</td>
<td>Very high cycle fatigue</td>
</tr>
<tr>
<td>S-N</td>
<td>Stress vs number of fatigue cycles</td>
</tr>
<tr>
<td>UTS</td>
<td>Ultimate Tensile Strength</td>
</tr>
<tr>
<td>AMT</td>
<td>Advanced Materials Technology</td>
</tr>
<tr>
<td>CTE</td>
<td>Coefficient of thermal expansion</td>
</tr>
<tr>
<td>HDT</td>
<td>Heat deflection temperature</td>
</tr>
<tr>
<td>DUT</td>
<td>Durban University of Technology</td>
</tr>
<tr>
<td>CNC</td>
<td>Computer numerical control</td>
</tr>
<tr>
<td>HNO₃</td>
<td>Nitric Acid</td>
</tr>
<tr>
<td>wt%</td>
<td>Weight percentage</td>
</tr>
<tr>
<td>A</td>
<td>Angstrom</td>
</tr>
<tr>
<td>kPa</td>
<td>Kilo Pascal</td>
</tr>
<tr>
<td>MPa</td>
<td>Mega Pascal</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission Electron Microscopy</td>
</tr>
<tr>
<td>CF</td>
<td>Carbon Fibre</td>
</tr>
</tbody>
</table>

**Mag**

magnification

**UTS**

Charge-coupled device

**AMT**

American society for testing and materials

**CTE**

Complementary metal oxide semi-conductor

**HDT**

Nanometer

**DUT**

Glass Fibre Reinforced Polymer

**CNC**

Montmorillonite

**HNO₃**

Mechanical testing system

**wt%**

Optical Microscope

**A**

Scanning electron microscope

**kPa**

Giga Pascal

**MPa**

United States of America

**TEM**

Vacuum assisted resin infusion method

**CF**

Nanoclay
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
<th>Expanded</th>
<th>Acronym</th>
</tr>
</thead>
<tbody>
<tr>
<td>mm</td>
<td>Millimeter</td>
<td>XRD</td>
<td>X-ray diffraction</td>
</tr>
<tr>
<td>FRP</td>
<td>Fibre Reinforced Plastic</td>
<td>PCM</td>
<td>Polymer matrix Composite</td>
</tr>
<tr>
<td>MMC</td>
<td>Metal matrix Composite</td>
<td>CMC</td>
<td>Ceramic matrix Composite</td>
</tr>
<tr>
<td>ROM</td>
<td>Rule of mixture</td>
<td>$E_1$</td>
<td>Elastic Modulus</td>
</tr>
<tr>
<td>$\varepsilon$</td>
<td>Strain</td>
<td>$\sigma$</td>
<td>Stress</td>
</tr>
<tr>
<td>$V_F$</td>
<td>Fibre Volume Fraction</td>
<td>$V_M$</td>
<td>Matrix Volume Fraction</td>
</tr>
<tr>
<td>$A$</td>
<td>Cross Sectional Area</td>
<td>SCFM</td>
<td>Self-consistent field method</td>
</tr>
<tr>
<td>H-T</td>
<td>Halpin–Tsai</td>
<td>$\mu$</td>
<td>Poisson’s ratio</td>
</tr>
<tr>
<td>$T_g$</td>
<td>Glass transition Temperature</td>
<td>$\tau$</td>
<td>Shear Stress</td>
</tr>
<tr>
<td>$\gamma$</td>
<td>Engineering Shear Stress</td>
<td>$kg$</td>
<td>Kilogram</td>
</tr>
<tr>
<td>$N_i$</td>
<td>Number of Cycles at the point of fracture</td>
<td>$N_p$</td>
<td>Number of cycle for the crack growth to occur</td>
</tr>
<tr>
<td>$\sigma_{UTS}$</td>
<td>Ultimate Tensile Stress at the fatigue loading percentage</td>
<td>$N_i$</td>
<td>Number of cycle for the crack to initiate</td>
</tr>
<tr>
<td>$\sigma_{max}$</td>
<td>Maximum Stress</td>
<td>PSB</td>
<td>Persistent Slip Band</td>
</tr>
<tr>
<td>Hz</td>
<td>Hertz</td>
<td>$^\circ C$</td>
<td>Degrees Celsius</td>
</tr>
<tr>
<td>$b$</td>
<td>Slope of the fatigue graph</td>
<td>EDX</td>
<td>Dispersion X-ray</td>
</tr>
<tr>
<td>NIOSH</td>
<td>National Institute for Occupational Safety and Health</td>
<td>HEPA</td>
<td>High-Efficiency Particulate Air</td>
</tr>
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</table>
CHAPTER 1

1.1 BACKGROUND

The primary structural material of World War One aircrafts were wood however, the anisotropic nature and strength characteristics of wood varied, depending on the source of supply [1]. Metal was a potential alternative [1]. Metals are strong, hard and durable and were therefore sought after materials in the manufacture of military and space products. Military vehicles, such as airplanes, helicopters, and rockets placed a premium on high-strength, light-weight materials [2] but it was realized that it would not be a simple matter of substituting metal for wood. Attempts to use metals such as aluminum and steel did not immediately satisfy the desired goal of weight reduction. The higher the weight of the aircraft itself, the less cargo its engines could carry. This led to the exploration into new polymer products, which could be lighter while affording the same strength and durability as metal and metal alloys [2]. In the aero vehicle industry, weight is money. The heavier an aircraft is, the more fuel it requires to fly. The more fuel it consumes, the more the running costs will be. In order to decrease fuel consumption and improve the aerodynamic performance of the aircrafts, engineers moved away from using aluminum. Instead today's latest aircrafts like the Boeing's 787 Dreamliner and Airbus's A350 [3], rely on lightweight carbon fibre composites i.e. woven mats of carbon which are embedded into a polymer.

A polymer [4] is a substance which consists of a large number of chains that come together to form a complex structure and is typically used in the manufacture of plastics and resins. Polymers were used as lightweight materials for the replacement of metal and metal alloys, provided that these polymeric materials possessed improved mechanical properties [2]. Hence fibre like materials which had high strength were incorporated into polymers to achieve comparable properties with metal and metal alloys. This discovery in turn lead to the discovery of materials like fibres which have an extremely high strength.
Engineers soon realized that by incorporating fibres in a matrix of a lightweight, lower-strength polymer, they could obtain a stronger material called a composite [5]. The fibres delayed the propagation of the cracks in the matrix. According to Reade [6], this implied that a polymer with insufficient strength or stiffness intended for use as an airplane wing could be reinforced with fibres to produce a stronger, stiffer and light-weight product. While these materials used in the form of long fibres were strong they were also brittle and hence when they failed, they did so catastrophically [2].

Defects, such as micro cracks in the material, could severely undermine the high strengths, since the defect sizes are different for each manufactured piece. Also since the number of defects generally scale with the size of the component, the only solution was to use short fibres [2], for the increased strength, the defects were generally minimized in the system.

Major world events prompted the use of new stronger reinforcement fibres. Consequently, graphite (carbon) fibres were produced using rayon as the starting raw material [6]. While carbon and boron fibres were developed around the same time, carbon took the lead due to its superior processing capabilities and its lower cost [5]. Carbon fibres were of use only in polymer matrices at this time. According to Koo [7], carbon fibre and a polymer matrix gave far superior mechanical strength properties when compared to that of glass fibre. Currently, the 787 Dreamliner which is Boeing’s latest plane, used composites for more than half of its airframe and that is including the fuselage and the wing. The Airbus’s A350 XWB [3] had both its wings and fuselage made from carbon fibre. Carbon fibre reinforced polymer composites used for the creation of sweeping wing tips reduced fuel consumption by up to 5% and this was achieved while keeping the aircraft still conventionally shaped.

Both academic and industrial researchers extended the composite paradigm to smaller scales, due to the need for minute and more efficient options and hence the era of the nanocomposites was born. Nanocomposites entail infusing nanoparticles, such as nanoclays, into the polymer/fibre matrix system.
Nanoclays are nanoparticles of layered silicates. Koo [7], further stated that a carbon fibre and polymer matrix infused with nanoclays results in enhanced improvement in temperature performance, mechanical properties, damage resistance and environment corrosion resistance of materials. A composite with carbon fibre added to polymer infused nanoclay is called a polymer hybrid composite.

1.2 THE COMPOSITE SYSTEM

A composite is any material made of more than one component. There are many examples in everyday life, for example concrete is a composite. It is made of cement, gravel and sand and sometimes steel rods to reinforce it. Each component is added in a particular quantity, at specific time for specific period to serve a specific purpose. The composite structure can be created when a polymer resin and hardener mixture is passed through the fibres using methods such as vacuum bagging and hand lay-up processes. In some instances composite materials may require heat and pressure during processing [8]. This compound is generally referred to as Fibre Reinforced Plastic (FRP).

The classification of a composite is as per the diagram shown in Figure 1.1. There are 3 matrix composites, polymer matrix composites (PMC), metal matrix composites (MMC) and ceramic matrix composite (CMC). The composites that will be studied in this thesis will be polymer composites that are made from polymer matrix, fibres and nanoclays. Fibres used in the production of composites are fiberglass, carbon (graphite), Kevlar, boron, diolen and vectran. The fibre most often utilized is glass fibre however sometimes; Kevlar and carbon are used, depending on the application. Boron, diolen and vectran although have their own unique properties, however they are less desirable as compared to glass, Kevlar and carbon, due to cost, processing and temperature resistance.
Fibres are embedded into the matrix in order to make the composite stronger. The fibres themselves are strong and light and are often stronger than steel and weigh much less [5]. This means that composites can be used to make aircrafts and automobiles lighter and thus more fuel efficient and hence means that pollution will be reduced [4].

Global polymer production began in the mid-20th century, when low material and production costs, new production technologies and new product categories combined to make polymer production economical. The industry finally matured in the late 1970s as seen in Figure 1.2, when world polymer production surpassed that of steel, making polymers the ubiquitous material that it is today. From the 1970s the usage of carbon fibre in the Airbus 300 started off at 4t/unit and increased gradually to 9t/unit by 1995. And in 2015, the Airbus A350 consisted of 32t/unit of carbon fibre [9].

![Classification based on Matrices](image)

**Figure 1.1: Classification of Composites**
The growth of the use of composites in the Airbus’s from the 1990’s till present has increased steadily, from the Rudder, Spoilers, and Airbrakes to the Wing box and Fuselage. The Airbus is now made up of 50% composites and this percentage will continue growing. Fibre reinforced plastics have been a significant aspect of this industry from the beginning.

The trailing edges of the wings, that extends back and down takes the full load of forces when the airplane is landing. This is the component that takes the full load of the forces when travelling through the air at 250mph and it is light enough to be lifted by a human being. This is one of the recent developments of the Airbus’s latest plane, the A350 [3]. As per Sirett [11], if a kilogram of weight is removed, that is a major saving over the life span of an aircraft.

![Evolution composite application at Airbus](image)

**Figure 1.2:** Showing increasing Composite usage from 1970 [10]

With this kind of technology, the parts on an airbus can be cut down considerably from six million parts. With fewer parts, it means manufacturing time and costs
will be cut down considerably. Sirett goes further to state that with each kilogram saving on an Airbus, this equates to roughly $1M in costs over the lifetime of the aircraft and that with use of composites the weight of the aircraft can be reduced by up to 20% [11].

### 1.2.1 Properties of Composites

Composites have this unique ability to offer strength and light weight to a product. The properties that contribute to these qualities are described below. The rapid growth in the use of composite materials in structures requires the development of structure mechanics for modelling the mechanical behaviors and the analysis of structural elements made of composite material.

The properties of composites may be estimated by the application of the simple rule of mixture (ROM) theory. These rules can be used to estimate the average composite mechanical and physical properties along different directions. They are therefore extremely useful in the assessment of the combinations of basic mechanical/physical properties that can be engineered via composite reinforcement [12]. In order to calculate the elastic modulus of the composite material in the one-direction (E₁), Voigt postulated that both the matrix and fibre experience the same strain (ε₁) as shown in Figure 1.3. This strain is a result of a uniform stress (σ₁) being applied over a uniform cross-sectional area, A [13].

*Equation 1.1: ROM equation for the apparent Young’s modulus in the fibre direction [13]*

\[
E_1 = E_F V_F + E_M V_M
\]  

(1.1)

Where \( E_F, E_M, V_F \) and \( V_M \) are the moduli and volume fractions of the fibre and matrix materials respectively. This model works extremely well for aligned continuous fibre composites where the assumption of equal strain in the two components is correct.
The elastic modulus, $E$ of the composite in the two-direction ($\varepsilon_2$) is determined by assuming that the applied transverse stress ($\sigma_2$) is equal in both the fibre and the matrix as shown in Figure 1.4 (Reuss assumption) [13]. As a result, $E_2$ is determined by an inverse rule of mixtures equation that is given as [13].

![Figure 1.3: Representation of matrix and fibre stressed in one direction](image)

**Equation 1.2: Inverse rule of mixtures equation**

$$E_2 = \frac{E_F E_M}{V_M E_F + V_F E_M}$$  \hspace{1cm} (1.2)

For all composites with well-bonded reinforcements, Young’s modulus in the principle fibre direction will be somewhere in between the extreme values predicted by the ROM equations.

The mathematical model, Halpin–Tsai, (H-T) is used for the prediction of elasticity of composite materials based on the geometry and orientation of the filler and the elastic properties of the filler and matrix [14]. The model is often considered to be empirical, although it is based on the self-consistent field method (SCFM), which is also called the Hartree-Fock Method. These relations, which are called semi-
empirical relations because the parameters involved in these relations have some physical significance, were developed by curve fitting to the results that’s are based on elasticity [15].

Figure 1.4:  Representation of matrix and fibre stressed in two directions

The assumption for the creation of the ROM and the H-T equations are the basics, i.e. both the matrix and fibres are linearly elastic, isotropic and homogeneous, fibres are perfectly aligned and spaced, matrix is void free and the bonding between the matrix and the fibres is perfect [15].

1.2.2 Poisson Ratio

When a composite material is stretched, longitudinally, it usually tends to contract in the direction transverse to the direction of stretching, which is lateral as depicted in figure 1.5. Poisson's ratio, μ is a measure of this effect that is it is the negative ratio of the lateral strain and the longitudinal strain.
The Poisson’s ratio of the carbon fibre is 0.30. This means that the material has a greater ability to stretch in one direction than it is able to contract in the other direction. The bonds between the atoms in the structure become realigned during the process of stretching and compressing. In most cases, a material’s Poisson ratio will range between 0 and 0.5. Among common materials, rubber has a Poisson ratio very close to 0.5, whereas steel has one of 0.3 and cork is much closer to 0 [16]. This is why wine corks are made of cork: it can withstand the pressure from the neck of the bottle without stretching vertically and jamming in place. Hence it can be seen that the composite can stretch and allow contraction sufficient to adapt to any situation.
1.3 COMPONENTS OF THE COMPOSITE

As already discussed previously the main components of the composite are the matrix and the fibre reinforcement. These are broad terms and the different categories will be discussed further.

1.3.1 Types of resin matrices

The resins that are used in fibre reinforced composites can also be referred to as polymers. All polymers exhibit an important common property, that is they are composed of long chain-like molecules consisting of many simple repeating units. Man-made polymers are generally called 'synthetic resins' or simply 'resins'. Polymers can be classified under two types, 'thermoplastic' and 'thermosetting', according to the effect of heat on their properties. Figure 1.6, shows the difference between thermoplastic and thermoset resins. The black dots represent the permanent bonds that are formed, that heat cannot destroy.

Thermoplastics will soften with heating and eventually melt and harden again upon cooling. This repetition of the process of crossing the softening or melting point can be repeated as often as desired, without any appreciable effect on the material properties in either state. Typical thermoplastics include nylon, polypropylene and ABS, and these can be reinforced, although usually only with short, chopped fibres such as glass.

Thermosetting materials, or 'thermosets', are formed from a chemical reaction in situ, where the resin and hardener or resin and catalyst are mixed and then they undergo a non-reversible chemical reaction to form a hard, infusible product. Once cured, thermosets will not become liquid again if heated, although above a certain temperature their mechanical properties will change significantly. This temperature is known as the glass transition temperature (Tg), and varies widely according to the particular resin system used, its degree of cure and whether it
was mixed correctly. Above the Tg, the molecular structure of the thermoset changes from that of a rigid crystalline polymer to a more flexible, amorphous polymer. This change is reversible on cooling back below the Tg. [18]. For this study, a thermoset, polyester resin will be utilised, as it is not desired, for example for the wing of an aircraft to become flexible when the polymer resin becomes liquid.

![Figure 1.6: Showing the difference between thermoplastic and thermoset resin [17]](image)

Thermoset materials can be brittle due to high crosslink density and have high service temperature. Due to presence of strong covalent bonds as crosslinks, thermoset cannot be recycled easily. Incorporation of fillers in form of particulates and fibres help to reduce transfer and distribute stress applied, and can be used in high end applications. [19].

There are three main types of resin matrices; namely; polymer matrix composite, metal matrix composite and ceramic matrix composite.
The matrix is a major component of any composite, hence a polymer matrix composite is one in which resins and polymers are present in major quantity and fillers are added to improve stiffness of the polymer. For example, reacting polyester with carbon-carbon double bonds in its backbone and styrene. This mixture is then mixed with the fibres and allowed to cure [20].

![Chemical structure of polyester and styrene reaction]

*Figure 1.7: Showing the polyester and styrene reaction [20]*

Figure 1.7 shows the reaction that occurs when polyester and styrene are mixed to create a resin matrix. The styrene and the double bonds react by “free radical vinyl polymerization” to form a cross-linked resin [20]. As the reaction continues and more cross links are generated, a network of molecules is created. This is an example of the thermosetting resin.

A metal matrix composite (MMC) is a composite material with at least two constituent parts, one being a metal and the other material may be a different metal or another material, such as a ceramic or organic compound. MMCs are made by dispersing a reinforcing material into a metal matrix. The reinforcement surface can be coated to prevent a chemical reaction with the matrix. For example, carbon fibre is commonly used in aluminum matrix to synthesize composites showing low density and high strength [21]. However, carbon reacts
with aluminum to form a brittle and water-soluble compound $\text{Al}_4\text{C}_3$ on the surface of the fibre. To prevent this reaction, the carbon fibres are coated with nickel or titanium boride. The carbon fibres here are not coated with any inhibitors.

Ceramic-matrix composites (CMCs) comprise of a ceramic matrix reinforced by a refractory fibre, such as silicon carbide (SiC) fibre. CMCs offer low density, high hardness and superior thermal and chemical resistance. These qualities coupled with their intrinsic ability to be tailored as composites, make CMCs highly attractive in an array of applications, most notably internal engine components, exhaust systems and other “hot-zone” structures, where CMCs are envisioned as lightweight replacements for metallic superalloys [22]. CMCs have been largely limited to missile structures, radomes and exhaust systems for fighter jets.

The three classifications described above offer a brief description of the available composite systems that are suitable for use in the military and space environment. Polymer matrix composites based on thermoset polymers i.e. polyester (epoxy) is chosen to work with, with an aim to improve stiffness of resin by incorporating nanoclays. This study will investigate the improvement of this brittle quality of the carbon fibre reinforced plastic through the use of nanoclays.

### 1.3.2 Reinforcement

Reinforcement provides strength and stiffness while polymer matrix protects and transfers loads between fibres. Long thin fibres possess high strength and stiffness. Particulate filler is a solid material which is capable of changing the physical and chemical properties of materials by surface interaction or by its own physical characterization. The combination of long thin fibres and particulate fillers will give rise to a hybrid structure with network links between long thin fibres-particulate fillers-polymer which shall impart high impact strength as well as high stiffness to a composite material.
Polymer resin has been used for the manufacture of products; however, its strength was very low. When fibres were added to the resin they became reinforced and as such the composite became mechanically stronger [2]. The fibres are processed along with resins, which develops a synergistic effect and leads to reinforcement of resins. Composites can be made stronger by lining up all the fibres in the same direction. When there is a force applied to the composite, see Figure 1.8, in the direction of the fibres, the composite is strong and will be able to withstand a higher force. However, if the force is applied to the composite at 90° to the fibre direction that is through the thickness of the fibre the composite strength is low.

![Diagram showing fibre direction and strength](image)

*Figure 1.8: Showing fibre direction and strength*

Sometimes strength is needed in more than one direction. In order to achieve this, the fibres are aligned in more than one direction, as shown in Figure 1.9. This type of fibre orientation gives strength in all directions. This is useful in aircraft design, as it cannot be anticipated where the point of stress or force to the material can occur.
Furthermore when an on-axis composite configuration is studied, the following can be seen, as shown in Figure 1.10, the principle axes, 1, 2 and 3 are aligned with the co-ordinate system. This also shows the definition of stress components associated with the material coordinate system, 1, 2, 3, where the stresses are volume averages over the fibre and matrix domains. The normal stresses are $\sigma_1$, $\sigma_2$, and $\sigma_3$, while the shear stresses are $\tau_{12}$, $\tau_{13}$, and $\tau_{23}$. Corresponding normal strains are $\varepsilon_1$, $\varepsilon_2$, and $\varepsilon_3$, and the engineering shear strains are $\gamma_{12}$, $\gamma_{13}$, and $\gamma_{23}$.

In thin, sheet-like structures such as a laminate, it is common to assume a state of plane stress by setting:

**Equation 1.4: State of plane stress**

$$\sigma_3 = \tau_{13} = \tau_{23} = 0. \quad [1.4]$$
Figure 1.10: Showing the element of the on-axis composite and definition of normal and shear stress components [23]

It may be shown that such a state of stress leads to vanishing of the out of plane shear strains, i.e.

**Equation 1.5: Out of plane shear strains**

\[
\gamma_{13} = \gamma_{23} = 0. \tag{1.5}
\]

The out-of-plane extensional strain, \( \varepsilon_3 \), does not vanish but becomes coupled to the in-plane stresses \( \sigma_1 \) and \( \sigma_2 \) and does not remain an independent quantity. The stress-strain relation for plane stress becomes:

**Equation 1.6: Stress-strain relation for plane stress**

\[
\begin{bmatrix}
\sigma_1 \\
\sigma_2 \\
\tau_{12}
\end{bmatrix} =
\begin{bmatrix}
Q_{11} & Q_{12} & 0 \\
Q_{12} & Q_{22} & 0 \\
0 & 0 & Q_{66}
\end{bmatrix}
\begin{bmatrix}
\varepsilon_1 \\
\varepsilon_2 \\
\gamma_{12}
\end{bmatrix} \tag{1.6}
\]
A common approach to achieve a set of more balanced mechanical properties is to utilize woven fabric composites, where tows of several thousand fibres are arranged in a specific pattern, such as plain weave fabrics consisting of “one fibre bundle over-one under” as shown in Figure 1.11. Figure 1.11 shows the various types of woven structures that are available, that is; plain weave, satin 5HS, 2x2 twill weave and non-crimp weave. Each type of weave has their own use for a particular application, with their own set of advantages and disadvantages. Direction of the individual fibre strands is referred to as the warp and weft (or fill), where the warp is the vertical fibre strand and the weft is the horizontal fibre. In this dissertation, the plain woven weave will be discussed.

Unidirectional composites, such as shown in Figure 1.11, are rarely utilized in actual structures because of their extremely anisotropic properties and weak failure planes. Most composite structures are multi-directional laminates, where unidirectional plies are oriented in different directions $\theta$, e.g., $\theta = 0^\circ$, $45^\circ$, $-45^\circ$, and $90^\circ$ Figure 1.12. It is observed that each ply orientation $\theta$ is obtained by a suitable rotation of the ply around the $3$ axis.
Figure 1.11: The different types of Woven Weaves available. [105]

Figure 1.12: (a) A multidirectional laminate is built by a stack of unidirectional composites bonded together. (b) Definition of ply orientation angle $\theta$. [23]
1.3.2.1 Glass fibre

Glass wool, which is also referred to as "glassfibre", is a material that is used as thermal building insulation. Glass fibre has roughly comparable mechanical properties to other fibres such as polymers and carbon fibre [24]. Although not as strong or as rigid as carbon fibre, it is much cheaper and significantly less brittle when used in composites. Glass fibre is therefore used as a reinforcing agent for many polymer products; to form a very strong and relatively lightweight fibre-reinforced polymer (FRP) composite material called glass fibre reinforced plastic (GFRP). Figure 1.13 shows the 3 different types of fibres; glass, carbon and Kevlar.

1.3.2.2 Kevlar fibre

Kevlar is the registered trademark for a para-aramid synthetic fibre. This high-strength material was first commercially used in the early 1970s as a replacement for steel in racing tyres [26]. Typically it is spun into ropes or fabric sheets that can be used as such or as an ingredient in composite material components.

![Figure 1.13: Glass, Carbon and Kevlar fibres [25]](image)
Considering its high tensile strength-to-weight ratio, currently, Kevlar has many applications, ranging from bicycle tyres and racing sails to body armour because of it being five times stronger than steel [4]. When used as a woven material, it is suitable for mooring lines and other underwater applications [26].

1.3.2.3 Carbon fibre

Carbon fibre is also called graphite fibre or carbon graphite and consists of very thin strands of the element carbon. These fibres have high tensile strength and are very strong for their size and is known to be the strongest material available [8]. Each fibre strand is 5-10 microns in diameter. To give a sense of how small that is, one micron (µm) is 0.000039 inches. One strand of spider web silk is usually between 3-8 microns. Carbon fibres are twice as stiff as steel and five times as strong as steel (per unit of weight) and is also highly chemically resistant and has high temperature tolerance with low thermal expansion [8].

Carbon fibre is created through a series of processes which are graphically depicted in figure 1.14. Polyacrylonitrile fibres (PAN), pitch resins, or rayon are carbonized (through oxidation and thermal pyrolysis) at high temperatures. Through further processes of graphitizing or stretching the fibres, strength or elasticity can be enhanced respectively. Carbon fibres are manufactured in diameters analogous to glass fibres with diameters ranging from 9 to 17 µm. There are typically five segments in the manufacturing of carbon fibres from the PAN process [8]. These are:

1. PAN Precursor Spinning: PAN mixed with other ingredients and spun into fibres, which are washed and stretched.

2. Oxidizing: Chemical alteration to stabilize bonding.

3. Carbonizing: Stabilized fibres heated to very high temperature forming tightly bonded carbon crystals.
4. Treating the Surface: Surface of fibres oxidized to improve bonding properties.

5. Sizing: Fibres are coated and wound onto bobbins, which are loaded onto spinning machines that twist the fibres into different size yarns. Instead of being woven into fabrics, fibres may be formed into composites. To form composite materials, heat, pressure, or a vacuum binds fibres together with a plastic polymer. These steps are graphically displayed in figure 1.14.

![Figure 1.14: Carbon Fibre production process [27]](image)

These fibres are wound into larger threads during a spool winding process for transportation and further production processes [28]. Further production processes include weaving or braiding into carbon fabrics, cloths and mats which are then used in actual reinforcement processes [29].

There are many challenges in the manufacturing processes, but the most important and relevant is the need for more cost effective recovery and repair. The surface treatment process must be carefully regulated to avoid creating pits that could result in defective fibres. Close control is required to ensure consistent quality and to minimise skin irritation, breathing irritation, arcing and shorts in electrical equipment because of the strong electro-conductivity of carbon fibres and of the health and safety issues [30].
1.3.2.3 Other Types of Reinforcements

There are other types of reinforcements that can be utilized. They are:

- Polyester – Due to lack of stiffness, it is usually excluded from being used in composites. It is however very useful due to its low cost and where high impact, low weight or abrasion resistance is desired [31]. It has low modulus, low density and high tenacity with good impact resistance. It is useful as a surfacing material as it can be smoothened and is compatible with most resin systems.

- Boron
In order to improve the overall fibre properties, metal or carbon fibres are often coated with a layer of boron. A boron/carbon hybrid, composed of carbon fibres among 80-100μm boron fibres, in an epoxy matrix, can achieve properties greater than either fibre alone, with flexural strength and stiffness twice that of carbon and 1.4 times that of boron, and shear strength exceeding that of either fibre. Its use is restricted in high temperature aerospace applications and in specialized sporting equipment, due to its extremely high cost.

- Ceramics
These are most frequently linked with non-polymer matrices such as metal alloys. Ceramic fibres, are mainly used in areas requiring high temperature resistance, usually in the form of very short ‘whiskers’.

- Natural
It is possible to use silk, wool and fibrous plant materials such as jute and sisal as reinforcements in ‘low-tech’ applications. Fairly high specific strengths can be achieved due to the fibre’s low specific gravity which is typically between 0.5-0.6.
1.3.3 Nanoclays

The nanoclay utilized was Cloisite®15A which was obtained from Southern Clay Products Inc., USA. Cloisite 15A is a dimethyl, dehydrogenated tallow, quaternary ammonium with Chloride as its anion. Cloisite 15A improves various plastic physical properties, such as heat deflection temperature (HDT), coefficient of thermal expansion (CTE) and thermal barrier [32]. There are various nanoclays available, Cloisite Na+, 10A, 15A, 20A, 25A, 93A AND 30B, each with their own unique chemical structure. Figure 1.15 below, illustrates the structure of Cloisite 15A. HT denotes hydrogenated tallow which consists of ~65% C18, ~30% C16 and ~5% C14 [33].

![Chemical structures of Cloisite 15A](image)

*Figure 1.15: Chemical structures of Cloisite 15A [33]*

The interlayer distance for Cloisite 15A is higher than the other Cloisite. This implies that the desorption of organic compounds will be easier. It is for this reason that Cloisite 15A was chosen as the nanoclay to use during the processing of the composite structure.

At the nanoscale level, unusual physical, chemical, and biological properties can emerge in materials. Nanomaterials behave differently from bulk materials of the same chemical because of the increased surface area – or the area of an object that is an exposed surface.
In this thesis the type of clay that is used is called nanoclay which are clay minerals optimized for use in clay nanocomposites—multi-functional material systems with several property enhancements targeted for a particular application. Nanoclays are a broad class of naturally occurring inorganic minerals, of which plate-like montmorillonite is the most commonly used in materials applications. Montmorillonite consists of ~1 nm thick aluminosilicate layers surface-substituted with metal cations and stacked in ~10 µm-sized multilayer stacks. The stacks can be dispersed in a polymer matrix to form polymer-clay nanocomposite, as shown in Figure 1.16. Within the nanocomposite, individual nm-thick clay layers are fully separated to form plate-like nanoparticles with very high (nm × µm) aspect ratio. Even at low nanoclay loading (a few weight %), the entire nanocomposite consists of interfacial polymer, with majority of polymer chains residing in close contact with the clay surface. Potential benefits include increased mechanical strength, decreased gas permeability, superior flame-resistance, and even enhanced transparency when dispersed nanoclay plates suppress polymer crystallization. [3]

Nanomaterial-enabled products such as surface-coated materials, nanocomposites, and materials comprised of nanostructures (Integrated circuits) are unlikely to pose a risk of exposure during their handling and during their use as the materials are of a non-inhalable size. However, some of the processes, like the cutting or grinding of the products could release respirable-sized nanoparticles used in their production which may lead to exposure to nanomaterials.

Based on studies done on animals, nanoparticles can enter the blood stream, and translocate to other organs. Results from human and animal studies carried out by NIOSH [11] documented that airborne nanoparticles can be inhaled and deposited into the respiratory tract. Engineering control techniques which are effective for capturing airborne nanoparticles and source enclosure i.e., isolating the generation source from the worker are some examples of the precautions that can be carried out when handling nanoparticles.
Exhaust ventilation systems with a high-efficiency particulate air (HEPA) filter are known to effectively remove nanomaterials. Good basic work practices e.g. washing hands, wearing appropriate personal protection equipment and no eating/drinking in the work area can help to minimize the workers exposures to nanomaterials.

1.4 HYBRID COMPOSITES

A hybrid composite is one where there are three or more components. They could be the resin matrix, the fibre reinforcement and particles, such as nanoclays. Figure 1.17 shows the process of fabricating a hybrid composite. The nanoclays are added to the resin mixture and they are allowed to interact. After this interaction they create either an intercalated or exfoliated arrangement. This resin mixture is now added to the fibre reinforcement, via a processing technique, for example, resin transfer moulding. The composite is cured and is ready for fabrication and testing.
Figure 1.17: Schematic showing the process of creating a hybrid Nanocomposite [35]

1.4.1 Processing of hybrid composites

Composites have been known for years. Their use in high end applications like aerospace, motor vehicles and sporting equipment is extensive, as indicated by Tong et al. It is with this growing confidence that research in this field is still continuing.

The processing techniques of these composites can be varied depending on their application. There are two distinct types of moulding, namely, composite moulding and wet moulding. In composite moulding, the resin is fibre reinforced first before being put through further moulding processes. Sheets of prepreg (pretreated with resin) FRP are heated or compressed in different ways to create geometric shapes. In wet moulding, the fibre reinforcement and the matrix are combined during the moulding process [36], namely, Resin Transfer Molding (RTM), High pressure Moulding (HPM) & Vacuum Assisted Resin Transfer Molding (VARTM).
Fibres are placed into a mould into which wet resin is injected. The resin is pressurized and forced into a cavity which is under vacuum. Resin is pulled into cavity under vacuum in the VARTM process. This moulding process allows precise tolerances and detailed shaping [36].

In the aerospace industry, tensile testing is used for material strength testing of adhesive bonds, aircraft textiles and carpets, peel tests on the airframe composites and shear and tensile strength testing of fasteners, e.g. bolts, nuts and screws and tensile. In the Automotive Industry, quality assessments of interior fittings (airbags, carpets, dashboards, electrical harness, handles, laminated trim, mirrors, seals, seatbelts) and handbrake levers.

There are also benefits for the Beverage, Construction, Electrical, Medical, Packaging, Paper and pulp, pharmaceutical and Plastics, Rubber and Elastomer Industries. In all these applications, there are ASTM standards that govern the dimension of the test specimens, the frequency and the test speed [37]. The Mechanical Testing Equipment is the universal testing equipment that all the industries use in order to perform the tensile tests.

1.4.2.1 Fatigue Testing
Fibre reinforced polymer composites have become an attractive replacement for heavier metals due to their superior fatigue and corrosion properties. Although they may be less susceptible to fatigue failure than metals, fatigue can still occur. Composite fatigue failure is generally driven by fatigue failure in the polymer matrix, as such polymer kinetics becomes the obvious tool for predicting fatigue in composites [38].

Fatigue failures represent the greatest uncertainty with regard to the long-term service lifetime of the major structural components of a wind turbine, Boeing nose piece, satellites etc, often leading the designer to either add excess weight to compensate or to inadequately size a blade resulting in fracture – both costly errors limiting the cost advantage of wind energy.
Fatigue is the main reason that causes extensive damage throughout the specimen, which can lead to failure from general degradation of the material instead of a predominant crack. However in static loading of isotopic, brittle materials, a predominant single crack is the most common failure mechanism that can occur [39]. Matrix cracking, delamination, fibre breakage and interfacial debonding, are four basic failure mechanisms that occur in composite materials as a result of fatigue.

Despite the large amount of research carried out on fatigue failure, its true nature still remains unknown and damage, cracks or even complete failure due to cycling loads are constantly being reported. If the problem still exists after 100 years of research in the previous century, there is something to be explained. The idea behind this thesis work is not to provide answers to the unanswered questions, but to tackle the fatigue problem from a different perspective, ITO understanding the crack initiation and propagation characteristic during Fatigue testing.

This includes testing and analyzing specimens that have 0wt%, 1wt%, 3wt% and 5wt% nanoclays added into the matrix to further reinforce the composite, with addition of carbon fibre. Furthermore to analyze the complex nature of fatigue failure, highlight the crack initiation, propagation and arrest during the actual fatigue testing itself. These phenomena will allow the designer insight into the crack behavior when fatigue initiates.

1.4.2.2 Crack Initiation, propagation and arrest
Following on from the previous section, it can be noted fatigue failure does not happen suddenly, there are steps that occur prior to that, that build up to the final failure. These are highlighted in Figure 1.18. The fatigue life is generally divided into three stages/periods, Crack Initiation period, Crack Growth Period and then the final failure itself.
Figure 1.18: Different phases of the fatigue life

Crack Initiation is the number of cycles required to initiate a crack, $N_I$. It generally results from dislocation pile-ups and imperfections such as surface roughness, voids, scratch etc. hence; in this period fatigue is a material surface phenomenon.

Crack Growth is the number of cycles, $N_p$ required to grow the crack in a stable manner to a critical size, generally controlled by stress level. Since most common material contains flaws, the prediction of crack growth is the most studied aspect of fatigue. Crack growth resistance, when the crack penetrates into the material, depends on the material as a bulk property. It is no longer a surface phenomenon.

To further elaborate on the Crack initiation and growth, Figure 1.19, highlights the occurrences of the surface of a specimen during fatigue testing.

Intrusions or extrusions which are actually surface roughness, which can be created during manufacturing of the specimen itself, are the basis of determining crack initiation. From here the Persistent Slip Band, PSB is created which is the very start of the fatigue crack. Thereafter, the primary crack appears and after a long fatigue test cycle, ~ 1Million cycles, the secondary crack appears. These cracks will continue until the specimen has fractured. This phenomena occurs in a composite, however in a hybrid composite the mechanism will be slightly different. This will be discussed in this thesis.
However, for the purpose of this study the nanoclays have been added during the processing stage to the matrix itself. These nanoclays are thought to be able to arrest the crack and prolong the crack from actually fracturing the specimen and in essence actually prolonging the life span of the product.

1.4.3 Advantages and disadvantages of hybrid composites

CFRP has a higher performance for a given weight which leads to lighter aircrafts and hence fuel savings. Excellent strength to weight and stiffness to weight ratios can be achieved by employing the use of composite materials. Laminate patterns and ply build-up in a part can be tailored to give the required mechanical properties in various directions. It is easier to achieve smooth aerodynamic profiles for drag reduction. Complex double-curvature parts with a smooth surface
finish can be made in one manufacturing operation, hence reducing production cost. For instance, an A380 super-jumbo [3] has about six million parts which in the future can be reduced significantly. All the parts can be molded simultaneously. Manufacturing time will be shortened due to fewer components, saving money. Each precious kilogram cut, from an aircraft’s weight means a saving of approximately, $1m (£603,000) in costs over the lifetime of an aircraft and the use of such composites can reduce the weight of an aircraft by up to 20% [3].

Composites may be made from a wide range of processes. Composites offer excellent resistance to corrosion. According to Tong et al [36], chemical attack and outdoor weathering can be damaging to composites. New types of paint and stripper are being developed to deal with this problem.

Some thermoplastics are not resistant to some solvents. Detection of cracks and fatigue cracks are easily performed and the repair strategy for these cracks is equally as simple. Just the area where the crack has been detected can be repaired and the crack propagation can hence be stopped. By using composites, manufacturers can get a good component surface finish to optimize their aerodynamic performance.

Some disadvantages of composites are: Composites are more brittle than metals and thus are easily damaged. Cast metals also tend to be brittle and repair of the damages introduces new problems. Materials require refrigerated transport and storage and have limited shelf lives. Hot curing is necessary in many cases, requiring special equipment, curing either hot or cold takes time.

Keith et al [40] stated that the product is not finished until the last rivet has been installed. If rivets have been used and must be removed, this presents problems of removal without causing further damage.

Repair at the original cure temperature requires tooling and pressure. Composites must be thoroughly cleaned of all contamination before repair, composites must
be dried before repair because all resin matrices and some fibres absorb moisture.

During processing, pieces of carbon fibres can break off and circulate in the air in the form of a fine dust. Industrial health studies have shown that, unlike some asbestos fibres, carbon fibres are too large to be a health hazard when inhaled. They can be an irritant, however, and people working in the area should wear protective masks. The carbon fibres can also cause skin irritation, especially on the back of hands and wrists. Protective clothing or the use of barrier skin creams is recommended for people in an area where carbon fibre dust is present.

The sizing materials used to coat the fibres often contain chemicals that can cause severe skin reactions, which also requires protection. Carbon fibres are also good conductors of electricity. As a result, carbon fibre dust can cause arcing and shorts in electrical equipment. If electrical equipment cannot be relocated from the area where carbon dust is present, the equipment is sealed in a cabinet or other enclosure.

1.4.4 Application of hybrid composites

Every day, a new application is found for carbon fibre reinforced plastic. What started out forty years ago as a highly exotic material is now a part of everyday life. These thin filaments, a tenth the thickness of a human hair, are now available in a wide range of useful forms.

The first fibre reinforced plastic plane fuselage was used on a modified Vultee BT-13A designated the XBT-16 based at Wright Field in late 1942 [42]. In 1943 further experiments were undertaken to build structural aircraft parts from composite materials resulting in the first plane, a Vultee BT-15, with a Glass Fibre Reinforced Plastic (GFRP) fuselage being flown in 1944 [43]. A significant development in the tooling processes for GFRP components had been made by Republic Aviation Corporation in 1943 [44].
Carbon fibre has gone to the moon on a spacecraft, but it is also used widely in aircraft components and structures, where its superior strength to weight ratio far exceeds that of any metal. Thirty percent of all carbon fibre is used in the aerospace industry. From helicopters to gliders, fighter jets to microlights, carbon fibre is a make of the design and structure and hence increasing range and simplifying maintenance [45]. Carbon fibre application in sports goods ranges from the stiffening of running shoes to ice hockey stick, tennis racquets and golf clubs. ‘Shells’ (hulls for rowing) are built from carbon fibre, and many lives have been saved on motor racing circuits by its strength and damage tolerance in body structures. Carbon fibre is used in crash helmets too, for rock climbers, horse riders and motor cyclists – in fact in any sport where there is a danger of head injury.

The applications in the military are very wide ranging; from planes and missiles to protective helmets, providing strengthening and weight reduction across all military equipment. It takes energy to move weight; whether it is a soldier’s personal gear or a field hospital, and weight saved means more weight moved per gallon of gas. A new military application is announced almost every day. The latest and most exotic military application is for small flapping wings on miniaturized flying drones, used for surveillance missions [2]. The application and uses of carbon fibre are widely extensive and research is seeking to improve the technology and quality of products so that they can last longer, be cost effective and have minimal maintenance requirements. With the increasing use of CFRP, the future options are endless.

Future designs with composites will significantly improve a plane's lift-to-drag ratio, making it much more aerodynamically efficient, and reduce its overall weight. The Airbus unveiled its proposal for an airliner of the future and this too moves away from the traditional narrow tube-like fuselage. Instead, in its 2050 concept plane the wings are slimmer and longer to reduce drag as shown in figure 1.20. The new fuselage efficient design is fatter, curved and shaped to improve
airflow and to provide more internal space. The engines will need less frequent
maintenance by the ground crew as it will become more efficient.

![Airplane in flight]

*Figure 1.20: Versatile Carbon fibre airframes mean that aircraft can be very
different in shape in the future [3]*

The tail section is U-shaped, which acts as a shield, cutting down on engine noise.
This means they can be partially embedded in the airframe to improve fuel
consumption. Elements of these designs are certain to be incorporated in all
future aircraft designs, in large part due to the materials revolution which is rapidly
changing all aspects of manufacturing [3].

In addition to lowering the overall airplane weight, moving to a composite primary
structure promises to reduce both the scheduled and non-routine maintenance
burden on the airlines. Experience with the Boeing 777 proves that composite
structures require less scheduled maintenance [37] than non-composite
structures. For example, the Boeing 777 composite tail is 25% larger than the
Boeing 767’s aluminum tail, yet requires 35% fewer scheduled maintenance labor
hours. This labor hour reduction is due to the result of a reduced risk of corrosion
and fatigue of composites compared with metal. A composite structure also
results in less non-routine maintenance. The 777 floor structure is all composite and highlights the advantages of this material when applied in a harsh environment. Airline operators are aware of the undetectable fatigue cracking and corrosion difficulties experienced with traditional aluminum floor beams. The 777 model has been flying for more than 10 years [34] with more than 565 airplanes in the fleet and to date has not replaced a single composite floor beam.

The intended use for the CFRP for this dissertation is the aircraft for military, space and passenger travel. For all these applications, weight saving is of paramount importance. Weight saving will mean fuel saving, less emissions and the ability to transport more passengers and cargo. The passenger per kilometer consumption is currently at 5l/km. With the weight savings this value will be reduced considerably.

1.5 COMPOSITE HYBRID USED IN PRESENT STUDY

1.5.1 Aim of the Study:

The aim of this research was to fatigue test the test specimens, produced from composite panels processed via the VARIM process. The test specimens were infused with nanoclays at percentages of 0 wt%, 1wt%, 3wt% and 5 wt%. The fatigue tests were carried out to determine the Fatigue life of Carbon Fibre Reinforced Composites for aerospace applications.

The idea is to fabricate light weight composite panels with similar strength and improved stiffness of epoxy resin for airplane parts. This in turn can ensure that the aircraft can carry more fuel and can hence travel for longer periods or carry more passengers and hence the use of the aircrafts can become economically viable.
1.5.2 Objectives of the Study:

The objectives in order to achieve this research aim are as follows:

- Produce Carbon fibre composite panels with 0wt%, 1wt%, 3wt% and 5wt% nanoclay concentration.
- Fully characterize the nanoclays and the sample specimens.
- Perform dynamic and static testing on the sample specimens and analyse this data.

1.5.3 Problem statement

With growing technology, there is always a need to improve on existing products. Composites are widely used in aerospace; main body of the aircrafts, wings, satellites and automobile industry; door linings, engine covers. In 2011; A dream became a reality when a Boeing's new carbon-fibre 787 Dreamliner heralded a new age of air travel [24].

This Boeing’s structure comprised of 50% of composites parts, including the fuselage and the wings. Due to this new technology being used, this Boeing is now 20% more fuel-efficient than similar sized planes. However the question needs to be asked how this product can be made better, stronger, light weight and more durable that is to increase on the already phenomenal 20% fuel-efficient savings. The two main components of a composite are resin and the fibre itself. For the purposes of this study, carbon fibre will be utilized, since it is already a major component in high end applications and its mechanical strength is very high.

If more fibre or more resin is to be added then the product will be stronger however it will also get heavier. And with aircrafts ever increasing speeds and agility, this will certainly be a problem. However if a nanoclay is added to the composite with the existing processing methods, this composite will retain its same weight and
become stronger. However with the addition of the nanoclays into the composite, there is very limited knowledge about the behavior of the composite during fatigue conditions.

Research has been conducted with the epoxy only, with epoxy and nanoclay, with epoxy and fibre and then with the hybrid; fibre, epoxy and nanoclay. A similar trend is observed in results where the nanoclay addition enhances the static and dynamic properties of the material. However limited work has been performed on woven carbon fibre hybrid composite for aerospace application. This dissertation will focus on creating a woven CFRP in a thermoset resin with the addition of MMT nanoclays at percentages of 0wt%, 1wt%, 3wt%, and 5wt%. Aircrafts undergo shear forces on a daily basis, and it is these forces that can create stress points where the components can shear off and break away. Therefore, the focus of this study is to investigate the fatigue properties of nanoclay filled Carbon Fibre Re-inforced Polymer.
CHAPTER 2

Carbon Fibre Reinforced Polymer has been extensively researched over the past decade. Studies have shown that although they are lightweight, strong, durable and suitable for aerospace applications; their brittle nature and hence cracking effect, their delamination effects and shear stress effects decrease their life span. Previously, as discussed by Ratna et al [46], only metals have been used for aircraft construction, however when these structures have been failing untimely and the root cause was difficult to ascertain, the move was to use composites for aircraft structures.

There was an increasing trend to utilize composites in various parts of aircrafts, however, the need now is to improve on this further. If this material could be made lighter whilst offering the same strength to weight ratio, this would mean that the aircraft fuel efficiency would increase and hence the aircraft could carry less fuel. The increase in space area could accommodate more passengers and luggage. The hypothesis for this is to incorporate nanoclays during the processing of the composite, directly into the resin mixture. In current work, nanoclays were added in concentrations 0wt%, 1wt%, 3wt% and 5%, and improvement in fatigue life was observed as loading of nanoclay increased to 5%.

Numerous studies [46, 47, and 48] have been carried out by incorporating nanoclays into the composite during the processing stage to create a hybrid nanocomposite. Tensile (to obtain ultimate tensile strength), fatigue (to determine the breakage point under a constant strain) and fracture (to determine the cracking properties) testing were then carried out on these samples. Previous works carried out, report characterization of polymers only. Here in this work to strengthen these polymers, fibres were then added during processing and the orientation and direction of these fibres were varied to obtain the optimum strength that is required for the given application. Thereafter to provide sufficient strength along with stiffness a hybrid composite was fabricated where nanoclays at various percentages were added during processing.
2.1 BACKGROUND INTO FAILURE DUE TO FATIGUE

Grimmer and Dharan [49] stated that high cycle fatigue life in glass fibre reinforced composites was dominated by fatigue cracking in the matrix, which subsequently propagated and ruptured the main load bearing elements which are the fibres. They further stated that the low elasticity modulus of glass fibres, when compared to the high modulus of carbon fibre composites, may impose higher strains in the matrix leading to failure by fatigue. Therefore the addition of nanoparticles like organoclay was expected to contribute to decrease the scale of damage mechanisms, leading to an increased absorption of the strain energy through the creation of fine nano-scale cracks. Carbon fibre is already a preferred fibre over glass fibre and adding nanoclays to the composite will result in a hybrid nanocomposite being able to withstand a higher degree of fatigue before rupturing and failing.

The use of composite materials has increased in aircraft structures where reducing the structure of non-useful weight becomes a crucial design criterion so as to maximize the weight of the useful structures. According to Helmy and Hoa [50], tapered composite structures produced by terminating or dropping off some of the laminate plies have received much attention, since fatigue failure on these type of structures were on the increase. Kabir and Hoa [51] further stated that, it can be expected that adding nanoparticles in polymer matrix would improve the ability of energy dissipation under dynamic loading and thus enhance the damping properties. If energy dissipated is improved, then the fatigue life of the structure must also be enhanced.

2.2 FAILURE OF POLYMER STRUCTURES

Polymer structures on their own without any reinforcements are rarely ever used in aerospace application, since they are very brittle and fracture easily. Hence, work has been done in this area with the addition of nanoclays to strength the polymer.
2.2.1 The Effect of Nanoclay on Polymer structure

There are two particular characteristics of clays that are generally considered for nanocomposites; the first is the ability of the silicate platelets to disperse into individual layers and the second is the ability to fine-tune their surface chemistry through ion exchange reactions with organic and inorganic cations. These two characteristics are, of course, interrelated since the degree of dispersion of silicate platelets in a particular polymer matrix depends on the interlayer cations.

A nanocomposite may not be formed by the physical mixture of a polymer and clay. This situation is analogous to polymer blends and in most cases separation into discrete phases takes place. The poor physical interaction between the organic and the inorganic components leads to poor mechanical and thermal properties which occurs mostly in immiscible systems. In contrast, strong interactions between the polymer and the organo-modified clay in nanocomposites leads to the dispersion of organic and inorganic phases at nanometer level. As a result, nanocomposites exhibit unique properties not shown by their micro counterparts or conventionally filled polymers [52-56].

Alkylammonium or alkylphosphonium cations in the organomodified clays lower the surface energy of the inorganic host and improve the wetting characteristics of the polymer matrix, and result in a larger interlayer spacing. Additionally, the alkylammonium or alkylphosphonium cations can provide functional groups that can react with the polymer matrix, or in some cases initiate the polymerization of monomers to improve the strength of the interface between the inorganic and the polymer matrix [57].

2.2.2 Types of Nanoclay

Clays have silicate layers of thickness 1 nm and high aspect ratio (e.g. 10-1000). A few weight percent of clay that are uniformly dispersed throughout the polymer matrix creates a much higher surface area for clay/polymer interaction as compared to conventional composites. Depending on the strength of interfacial
interactions between the polymer matrix and clay, two different types of nanocomposites are thermodynamically achievable as described in Figure 2.1.

In an intercalated nanocomposite the insertion of a polymer matrix into the intergallery spacing of clay occurs in a regular crystallographic fashion, regardless of the clay to polymer ratio. Intercalated nanocomposites normally have interlayer of few molecular layers of polymer.

The second type is the exfoliated nanocomposites: the individual clay layers are separated in a continuous polymer matrix, and the average distance depends on clay loading. Usually, the clay content of an exfoliated nanocomposite is much lower than that of an intercalated nanocomposite.

![Figure 2.1: Showing the different types of nanoclay infused structures](image)

**Figure 2.1:** Showing the different types of nanoclay infused structures [57]

### 2.2.3 Techniques for Characterizing Nanocomposites

The structure of nanocomposites has typically been established using X-Ray Diffraction (XRD) analysis and Transmission Electron Microscopic (TEM) observation. Monitoring the shape, position, and intensity of the basal distance within the distributed silicate layers, the nanocomposite structure may be
identified as to whether it is intercalated, exfoliated or a combination of both. For example, in an exfoliated nanocomposite, the extensive layer separation associated with the delamination of the silicate layers in the polymer matrix results in the eventual disappearance of any coherent X-ray diffraction from the distributed silicate layers. On the other hand, for intercalated nanocomposites, the finite layer expansion associated with the polymer intercalation results in the appearance of a new basal reflection corresponding to the larger gallery height. Although XRD offers a convenient method to determine the interlayer spacing of the silicate layers in the clay, and in the intercalated nanocomposites (within 1–4 nm), little can be said about the spatial distribution of the silicate layers or any structural non-homogeneities in nanocomposites. Additionally, some clays initially do not exhibit well-defined basal reflections. Thus, peak broadening and decrease in intensity are very difficult to assess. Therefore, conclusions concerning the mechanism of nanocomposites formation and their structure based solely on XRD patterns are only tentative.

TEM allows a qualitative understanding of the internal structure, spatial distribution of the various phases, and views of the defective structure through direct visualization. However, special care must be exercised to guarantee a representative cross-section of the sample. Typically, when inter-layer spacing exceed 6–7 nm in intercalated nanocomposites or when the inter layers become relatively disordered in exfoliated nanocomposites, associated XRD features weaken to the point of not being useful.

2.2.4 Testing of Polymer Structures
Moodley and Kanny [48] showed that the tensile modulus and tensile strength of polypropylene increased by 120% and 160% respectively with 2wt% nanoclays and that degradation of these properties was noticed for 3%, 4% and 5% clay loading. It can be further noted that in this study polypropylene was utilized, prepared by the melt blend technique, with Cloisite 15A Montmorillonite (MMT) nanoclay. According to Mohan et al [46], tensile modulus of nanoclay filled resin
was significantly increased from 1.1Gpa to 3.59Gpa (226% increase) at 3% nanoclay concentration and the tensile strength was increased from 61.1MPa to 66.3MPa (8.5% increase). Here there was no fibre included, just the resin and the nanoclays. Testing was carried out up to 10wt% nanoclays, however the 3% showed the best results. The effect of nanoclays on specimens even without fibre as a reinforcing agent still showed amassed results.

2.3 FAILURE OF COMPOSITE STRUCTURES

When fibres are added to the polymer it becomes a fibre re-inforced composite. Fibres bring their own re-in forcing properties and these need to be explored.

Gamstedt and Talerja [58] proved that a thermoset based composite showed a higher fatigue resistance with few micro cracks initiated at distributed fibre breaks growing at a decelerating rate. The thermoplastic composite had a more pronounced fatigue degradation with a steeper fatigue life curve which was caused by widespread propagating debonds and the matrix cracks. So the use of a tougher and more ductile matrix results in a superior fatigue life performance.

Backe et al [59] studied very high cycle fatigue (VHCF) of CFRP and showed that CFRP undergoes fatigue damage at low loading amplitude and at very long lifetimes in the VHCF regime.

2.3.1 Fibre Orientation

There are various fibre orientations that are available, unidirectional, cross-ply and woven. There are also multi directional fibres. Research done by Kawai et al [60] on the fatigue studies of cross-ply and unidirectional CFRP, showed that the static tensile strength of unidirectional laminate increases with decreasing number of plies however by contrast, the fatigue behavior is insensitive to the number of plies. The cross ply laminate behaves in a similar way as when compared to the unidirectional laminate. The S-N relationship for the cross ply laminate is almost parallel to the result on the unidirectional laminate. This
observation indicates that the fatigue strength of the cross-ply laminate is determined by the fatigue strength of the axial plies. Hence since woven fibre is used here, the S-N relationship will be similar to that as studied by Kawai, hence it can be implied that the cycles will be greater than $10^6$ range.

### 2.3.2 Fibre Volume Fraction

Tensile tests have already been conducted on CFRP, with variations; for example with different void fractions, with different fibre orientation (in the case of unidirectional fibre) and with different thicknesses. In most cases tensile tests are carried out to determine the UTS and stiffness and then further testing takes place from here on, it is most often not the final test.

As per Brunbauer et al [61], the fibre volume fraction of the composite plays a major role in the stiffness and strength during tensile testing. It was concluded that fibre volume fraction is directly proportional to stiffness and strength of composites. This finding was further emphasized when Qi et al [62], went on further to test multi-layer composites with different volume fractions. The finding here was consistent with Brunbauer et al [61], where it was concluded that as the fibre volume fraction increased so did the tensile strength, even with multilayers. Hence during processing the fibre volume fraction was maintained at 55%. This goes in line with what has been discussed above as well. The higher the fibre volume fraction, the higher the stiffness and strength is. While strength is desirable quality, the stiffness is not always desired as it makes the material brittle and easy to snap. Hence the optimum fibre void fraction desired for this work is 55%.

### 2.4 FAILURE OF HYBRID STRUCTURES

These are processing techniques and surface additions that create a stronger and more durable composite. Borrego et al [63] and Wang et al [64] have studied the addition of nanoclays to the composite during processing. The theory is that
these nano particles will create stronger hybrid structures. Below are other studies carried out that demonstrated how nano particles are responsible for creating a stronger composite.

The percentage increase for the modulus and for the strength with the fibres in the matrix was even greater, as indicated by Zhou and Lee [65]. They indicated that the tensile modulus and strength of high-performance thermoset polymer composites synthesized by using both long carbon fibres and nanoclays, improved by 80% and 64%, respectively with 5% MMT nanoclay. Ratna et al [47] showed that with clay re-inforced epoxy the impact strength of nanocomposite was increased from 0.7kJ/m to 1.1 kJ/m (57% increase) with a clay content of 5%.

Borrego et al [59] pointed out that during the fatigue testing of glass fibre reinforced epoxy composites enhanced with nanoparticles, the nanoparticles act as a barrier for the fatigue crack propagation.

From the studies that have been conducted so far, it can be seen that the addition of MMT nanoclays increase the tensile static properties (UTS, Young’s Modulus and Impact Strength) of the resin alone and with the MMT nanoclay, the resin and carbon long fibre as well. It also shows that the optimum MMT Nanoclay concentration can vary between 3-5%, however higher concentrations above this point, results in product degradation [46].

2.4.1 Fatigue Failure Theory

In materials science, fatigue [64] is defined as the weakening of a material caused by repeatedly applied cyclic loading. The nominal maximum stress value that can cause such damage is usually less than the ultimate tensile strength of the material.

Fatigue occurs when a material is subjected to cyclic repeated loading and unloading. If the loads are above a certain threshold, microscopic cracks will begin to form at the surface. Eventually a crack will reach a critical size, the crack will propagate suddenly, and the structure will fracture.
The basic method of presenting engineering fatigue data is by means of the S-N curve, a plot of stress, \( S \) against the number of cycles to failure, \( N_f \). Usually, millions of cycles might be required at lower loading levels, to cause failure, so the abscissa in usually plotted logarithmically.

Three basic factors are necessary in order for fatigue cracks to initiate [52]. First, the loading pattern must contain minimum and maximum peak values with large enough variation or fluctuation. Secondly, the peak stress levels must be of sufficiently high value. Thirdly, the material must experience a sufficiently large number of cycles of the applied stress. The number of cycles required to initiate and grow a crack is largely dependent on the first two factors.

In addition to these three basic factors, there are a host of other variables, such as stress concentration, corrosion, temperature, overload, metallurgical structure, and residual stresses which can affect the propensity for fatigue. Since fatigue cracks generally initiate at a surface, the surface condition of the component being loaded will have an effect on its fatigue life.

Surface roughness is important as it is directly related to number and the level of stress concentrations on the surface. When the stress concentration is high, the more likely it is for a crack to nucleate. Smooth surfaces increase the time to nucleation. Notches, scratches, and other stress risers decrease fatigue life. Surface residual stress will also have a significant effect on fatigue life.

Figure 2.2a shows the cyclic behavior for tension and compression loading that is the specimen is pulled apart and then pushed together again and thus creating the compression effect. Figure 2.2b shows the cyclic behavior for tension and tension loading, that is the sample is pulled and then relaxed.
Over the years, fatigue failure investigations have led to the observation that the fatigue process actually embraces two domains of cyclic stressing or straining that are distinctly different in character. In each of these domains, failure occurs by apparently different physical mechanisms: one where significant plastic straining occurs and the other where stresses and strains are largely confined to the elastic region [53]. The first domain involves some large cycles, relatively short lives and is usually referred to as low-cycle fatigue. The other domain is associated with low loads and long lives and is commonly referred to as high-cycle fatigue. Low-cycle fatigue is typically associated with fatigue lives between about 10 to 100,000 cycles and high-cycle fatigue with lives greater than 100,000 cycles.

In summary, fatigue analysis may be thought of as a process of initiating and then growing a crack which finally causes the structure to break into two or more pieces. This process can be represented by the following equation:
**Equation 2.1: Total Life Calculation**

Total Life = Crack Initiation + Crack Growth, or

\[ N_f = N_i + N_p \]  

Where: \( N_f \) = Total fatigue cycles to failure  
\( N_i \) = Number of cycles to initiate a crack  
\( N_p \) = Number of cycles to propagate a crack to final fracture

Ho et al [66] points out that using nanoclays in composites, proved that 5 wt% nanoclay gave the highest UTS and Vickers Hardness value amongst all the nanoclay compositions from 0wt% through to 8wt%. The ductility however decreased drastically with the addition of the higher nanoclay content inside the epoxy sample, which would cause catastrophic failure. The nanoclay content should be kept below the 5wt% in order to get good mechanical properties.

### 2.4.2 Damage Characteristics

Khan et al [67] studied unidirectional CFRP containing nanoclays and found that the incorporation of nanoclay into CFRP composites not only improves the mechanical properties but also the fatigue life for a given cyclic load level. The corresponding fatigue damage area is significantly reduced due to the nanoclay.

When a stress is applied to the sample, dimples appear on the surface of the sample. Dimples are referred to small microscopic cavities within the structure of the sample. The size of these dimples increased in both width and depth with decreasing the stress level applied during fatigue or alternatively with an increase in fatigue life. The mechanisms of dimple growth and formation can be explained as follows: micro cracks are formed during the early stages of fatigue loading at weak locations where there are high stress concentrations. Localized plastic deformation occurs due to the strain concentration around these micro cracks,
which in turn promotes the formation of dimples. Khan et al further stated that, the lower the applied stress level, the longer this stage lasts. Dimples grow in a stable fashion with increasing number of fatigue cycles before they coalescence into major cracks and final failure. They tend to coalescence rapidly at high stress levels, leading to premature fracture without extensive growth of deep and wide dimples. The elongation or stretching of the localized deformation zones or dimples is highly dependent on the applied stress.

These dimples occur during crack initiation, however as the crack progresses and the number of cycle's increases, the dimples become larger and the fracture mechanism changes. After this matrix cracking, fibre breaking and delamination takes place and this determines the final fracture.

2.4.3 Fatigue Parameters

While the fatigue testing is being carried out, there are certain factors that need to be considered. One is the fatigue loading percentage and the other is the fatigue frequency.

2.4.3.1 Fatigue Loading Percentage

This percentage determines at which rate the sample will be fatigue tested. The ultimate tensile stress is required which is then referred to as the maximum stress. This is obtained from the static tensile test that is conducted. Khan et al [67] studied unidirectional CFRP containing nanoclays, and used fatigue loading percentages of 80%, 60% and 45%. In order to be comparable with the results and be able to fill in the gaps for the other percentages, 90%, 75% and 60% was used in this study.
2.4.3.2 Fatigue Testing Frequency

The frequency of the fatigue testing plays a major role during testing. Some of the samples can have fatigue cycles of more than 5 Million cycles, this will generate a lot of heat. So the frequency needs to be kept to a minimum in order to minimize the temperature generation. The loading frequency is considered to have a significant influence on the sensitivity to fatigue for carbon fibre composites for the effect of self-generated heating and intrinsic rate dependence, as stated by Curtis et al [68]. The surface temperature of specimens increase rapidly in the intermediate number of cycle range of 10,000 cycles and the increase in surface temperature was larger than 20°C, as observed by Pandita et al [69]. The magnitude of the change of the surface temperature depended on the fibre orientation as well as maximum fatigue stress. It was found that 2Hz offers the most consistent temperature profile, while 10Hz offered the highest temperature profile range. Based on these findings, 3 Hz was utilized. This offered the minimum heat generation however, still at a level, where the testing can take place.

2.5 MOTIVATION AND SCOPE OF RESEARCH WORK

Research has been conducted on fibre-reinforced hybrid structures with addition of nanoclays with favorable improvements in properties. Materials, in general, may be categorized as being either ductile or brittle [70]. This depends on the characteristics of failure. In the case of polymers, there are two primary deformation mechanisms, namely, crazing and shear band formation [71]. Shear band formation is believed to be the precursor to ductile failure [71]. Brittle failure occurs when crazing is the dominant deformation mechanism [72]. Thermoset polymers exhibit brittle fracture at low temperatures and ductile fracture at high temperatures [73]. The failure mechanism in thermoplastics can be both brittle and ductile. The failure mechanisms for tensile, flexural and other common mechanical testing are well documented. However, literature on failure due to fatigue mechanisms are not extensive, especially with the inclusion of nanoclays.
Therefore this study involves the processing and analysis of woven CFRP with the addition of nanoclays. Specimens were subjected to fatigue tests in order to determine the life cycle at different loading rates which were 90%, 75% and 60% at a frequency of 3Hz. The crack initiation, crack propagation and crack paths that were observed during the fatigue testing of these composite structures are discussed. The temperature and frequency effect of the test samples during testing and crack density and rate are also analysed and discussed.
CHAPTER 3

3.1 DESIGN OF RESEARCH

Products, aircraft, spacecraft, satellite, sporting goods and high performance motor vehicles, made with carbon fibre re-enforcement plastic (CFRP), undergo strenuous fatigue cycles on a daily basis. This research undertakes to identify and understand these fatigue cycles and study factors that can improve these cycles and hence extend the lifetime of the product.

The strategy adopted for this research was purely experimental, where specimens were created and tests were carried out which are similar to the operating conditions which the end products are subjected to. This research approach was both qualitative and quantitative. Qualitative in that ASTM standards which have already been devised and approved by a regulatory board were adhered to when preparing the specimens and the running tests. Quantitative in that experimental work carried out were done using five specimens for each test and the data obtained was analysed using approved methods and conclusions drawn.

Figure 3.1, outlines the design method that was adopted for this research work. Carbon fibre re-inforced plastic (CFRP) composites were prepared using the vacuum assisted resin infusion method (VARIM) during which time Cloisite® Nanoclays were added to the composite in the concentration of 0wt%, 1wt%, 3wt%, and 5wt%. The VARIM process was selected due to it being able to ensure a uniform thickness throughout the laminate during the infusion process due to the constant pressure applied to the entire laminate. Transmission electron microscope (TEM) and X-ray Diffraction (XRD) were the characterisation and the morphological methods that were carried out. Static (tensile, Impact, hardness, double cantilever beam and end notch flexure) and dynamic test (fatigue) were carried out on the prepared specimens to obtain the life cycle for the sample. The static tests and dynamic tests were carried out on 0wt%, 1wt%, 3wt% and 5wt% nanoclay samples.
### 3.1.1 Flow Diagram of Research Design

**Figure 3.1:** Flow Diagram highlighting the experimental design

[Diagram showing the experimental design with steps and tests involved, including Flow, Vacuum Infusion Process, Carbon Fibre Reinforced Polymer (CFRP), Tensile Test, Impact, Hardness, Fatigue Test, XRD, TEM, Woven Carbon fibre, Epoxy Resin + Hardener, Nanoclays 0, 1, 3 & 5wt%, Static Testing, Mechanical testing, Dynamic Testing, Failure Mechanism, Scanning Electron Microscope (SEM).]
3.2 RESEARCH METHODOLOGY

Research can be stimulated by various methods such as deductive logic, inductive logic, scientific methods, critical thinking and collaborative work. The method that best describes this research, as shown in Figure 3.2, is the scientific method for the following reasons.

Step 1: Observation: It is at this step where the idea or concept for the research is born. Carbon fibre is known as the strongest fibre, however it is brittle and under fatigue conditions it can break easily. Hence the question was raised as to what can be done to the material to make the final product stronger and hence last longer, yet still maintain its lightweight structure.

Step 2: Hypothesis: Through researching the subject, it was postulated to add nanoclays into the material during the manufacturing process to create a hybrid structure. During this stage, it was established what quantities of nanoclays would be used, that is 0wt% (reference sample), 1wt%, 3wt% and 5wt%. The tests that were to be carried out were highlighted at this stage. Tensile test to determine the fatigue loading, hardness and impact testing to determine the effect of the nanoclays on the brittleness of the material, fracture tests to determine the fracture mechanics of the material and then finally fatigue testing. The procedure to follow in order to manufacture the specimens for all the testing were also identified.

Step 3: Testing: This step is purely experimental. Specimens were cut and tests were carried out as per regulated standards. This step was a continuous and repetitive process. This process continued until the hypothesis was proven.

Step 4: Predictions: this is the last and final step, where the hypothesis raised earlier was proved using the relevant results from the experimental data performed.

Due to the process followed in this research study being very similar to the scientific approach highlighted in Figure 3.2, it can now be stated that this research is indeed a scientific method.
Figure 3.2:  *Demonstrating the key aspects of a Scientific Method*

Figure 3.2 shows the cyclic process that is followed in a scientific method procedure. As the diagram shows this procedure can be repetitive, which was the case in this research study.

3.2.1 Processing

3.2.1.1 Materials to manufacture the Composites

Woven carbon fibre with a density of 200 g/m² was acquired from Advanced Materials Technology (AMT) Composites, South Africa. Woven fibre was selected based on the fact that it offers excellent multi-axis properties and it lends great toughness and durability to a composite structure. The carbon fibre was purchased in 20mx1m rolls and thereafter cut into 300mx300mm sheets. The resin, LR20 and the hardener LH281 (Unmodified Cyclicaliphatic Amine) epoxy resin system was obtained from AMT Composites. The peel ply, perforated release film, distribution mesh and vacuum bagging were all obtained from AMT Composites as well and they all were procured in rolls and thereafter cut into the
desired lengths. The 8 mm tubing and the spiral distribution tubing were obtained from a local electrical supplier, Industrial Rubber and Engineering Supplies cc. The wax that was used as a release film for the hardened composite laminate was procured from a local hardware store. The nanoclay utilized was Cloisite®15A which was obtained from Southern Clay Products Inc., USA. Cloisite 15A is a quaternary ammonium salt modified natural montmorillonite (MMT) polymer additive, which improves various plastic physical properties, such as reinforcement, heat deflection temperature (HDT), coefficient of thermal expansion (CTE) and thermal barrier [74]. The pump that was utilised was N022AN 18 Vacuum Pump IP20 which was purchased from Labotec (PTY) Ltd, South Africa. A resin trap was utilised to prevent damage of the pump.

3.2.1.2 Processing of Composites
The laminate was produced using a vacuum assisted resin infusion method (VARIM). The advantages of this process as compared to the others are that it provides better fibre-to-resin ratio, there is less resin wasted, there is very consistent resin usage, there is unlimited set-up time and the process is much cleaner. Figure 3.3 highlights the key points of the VARIM infusion process.

![Figure 3.3: The VARIM Process [75]](image)
The infusion process was carried out on a clean, dry, scratch-free glass surface. Acetone was used to do a final clean on the surface, before applying a layer of wax. This wax was allowed to dry for 5 minutes before it was polished and buffed. The area that was to be used for the resin infusion was then marked off using the sealant tape. 10x300mmx300mm carbon fibre layers, which were cut from rolls of carbon fibre were weighed and then carefully placed one sheet at a time directly onto the waxed glass surface, in a 90°/0° orientation. Any loose threads were removed.

The peel ply layer was then placed directly over fibres, ensuring to cover all the fibres completely. The distribution mesh was then placed directly over the peel ply layer to allow for even distribution of the resin through the fibres. Breather fabric strips were placed at the suction end to trap excess resin and prevent it from getting into the resin trap. The vacuum plastic bag was then placed directly over, covering the entire setup. This was the crucial step during the infusion process. The vacuum was then applied to the setup and left to run for 5 minutes, then the pump was stopped and the setup was allowed to rest for 10 mins to check for any air leaks. Any leaks present could be detected via a drop in pressure which can be checked on the pressure gauge situated on the vacuum pump. If there were no leaks then all was set for the infusion. If there were any leaks present then the seals along the perimeter of the setup would have to be secured again and all piping checked as well to ensure sufficient clamping. Thereafter the pump was left running, while preparing the resin, to compress the fibres further and extract any excess air trapped in the fibres.

The LR 20 resin was weighed out in a glass beaker with a 1:1 ratio to the weight of the carbon fibre sheets and heated on a hot plate (Heidolph MR Hei-STD) to 70°C. The viscosity of the resin is high at 1500 mPa.s and processing of the resin becomes very difficult, hence it is heated to reduce the viscosity. The nanoclay Cloisite® 15A was weighed on a precision electronic scale (Boeco Germany, 4 decimal place) at the required concentration of 0wt%, 1wt%, 3wt% and 5wt% of the epoxy resin plus hardener mixture weight. This was added to the heated resin.
and homogenized at a rate of 500rpm using the magnetic stirrer hot plate. The heating of the resin allowed the nanoclays to enter into the resin structure and bond easily with it. Thermogravimetric analysis has shown that the organic compound of organoclay deteriorate and breakdown at temperatures of 180°C [76]. The mixture was maintained at 70°C for 1 hour.

During processing, difficulties infusing were encountered when the nanoclay concentrations were higher. It was discovered that above 5wt% nanoclay concentration, the resin and nanoclay mixture became very viscous and this created problems for the resin flow during vacuum infusion. The resin took too long to flow through the dense fibres and had difficulty filtering through the layers. The resin also gelled before reaching the end of the laminate to complete the infusion process.

The rate of infusion was calculated and it was found that the rate decreased by 26% from 0wt% to 5wt%. Previous research conducted showed best results at 2wt% [48] and 3wt% [46]nanoclay concentrations without fibre and at 3wt% and 5wt% nanoclay concentrations with fibre, hence it was decided to stop at 5wt% nanoclay concentration. The increment of nanoclay concentration was chosen at 0wt%, 1wt%, 3wt%, and 5wt% as these intervals would provide an adequate range to work with.

The epoxy resin was removed and allowed to cool to room temperature before infusion could take place. The translucent colour indicated uniform distribution of the nanoclays within the epoxy resin mixture. The hardener, LH281 was added to the epoxy resin nanoclay mixture at a ratio of 70:30 and mechanically stirred for 5mins until the mixture was completely homogenised. During this time the vacuum pump was still running. The inlet tubing, while still clamped, was slowly immersed into the beaker with the resin mixture. The clamp was slowly released and the resin was quickly sucked up into the tubing and into the fibres. This process for the resin to be completely absorbed by the system was 35mins. The gel time of the resin mixture is 45mins, as per the data sheets as supplied by the manufacturer, so this does give the user some flexibility with time. Once all the
resin has been absorbed, the inlet tubing was clamped again before any air ingress into the system could occur. The vacuum pump was left running for 2 hours to extract any excess resin in the system. The next day, 24hrs later, the vacuum bag was removed from the sealant tape and this released any vacuum pressure that was still left. The laminate was easily removed from the table due to the waxed surface. The peel ply, perforated film and distribution mesh were all removed at one time as they all stuck to the peel ply. The laminate was then left for 5 days to fully cure before use.

Figure 3.4, shows the complete setup for the resin infusion process as performed in the composite processing workshop. There is a resin trap between the vacuum pump and the pipe outlet, which collected any excess resin that got sucked through the pipe, and hence prevented damage to the pump. The spiral binder tubing at the inlet and the outlet pipes distributes the flow evenly through the process.

The cured composite laminates were cut by a CNC machine to the desired specifications as per ASTM standards.
Figure 3.4: Showing the actual Infusion setup used for the CFRP preparation in the composite processing lab at DUT.

3.2.2 Morphology

3.2.2.1 Transmission Electron Microscope (TEM)

The Transmission Electron Microscope (JEOL 1010 TEM) was used to investigate the morphology of the organoclay, as shown in figure 3.5.
The TEM gives the researcher an idea about the distribution and the arrangement of the nanoclays within the matrix. The ultra-thin specimens were produced via a process called microtoming. Microtoming was carried out using the Leica EM UC 7 microtome, where very thin samples between 60-150nm thick were cut. These microtomed specimens were collected hygienically using a flame and then placed onto filter paper until when required. TEM provided an image showing the dispersion of the nanoclays within the resin mixture. TEM could only be carried out on resin samples without the fibre, as the blades were unable to cut through the fibre during the microtoming process.
3.2.2.2 X-Ray Diffraction (XRD)

Another characterizing technique used in this study was XRD. X-ray diffraction has been in use in two main areas, for the fingerprint characterization of crystalline materials and the determination of their structure. Each crystalline solid has its unique characteristic X-ray pattern which may be used as a "fingerprint" for its identification. Figure 3.6, demonstrates how the XRD process works. A beam is passed through a material and this beam will be transmitted and/or diffracted. The angle between these two beams is measured and this gives the diffracted angle. Every crystalline material has its own unique angle and hence the material can be identified.

![Figure 3.6: Schematic of how XRD testing is performed [77]](image)

Once the material has been identified, X-ray crystallography may be used to determine its structure that is how the atoms pack together in the crystalline state and what the interatomic distance and angle are. X-ray diffraction is one of the most important characterization tools used in solid state chemistry and materials science. The size, shape and distribution of the unit cell for any compound can
easily be determined using XRD. The sample is crushed into a fine powder state and then placed into the crucible for XRD characterization.

The basal spacing or d-spacing may be determined using Bragg’s equation. This is the distance between basal layers of the MMT molecule.

**Equation 3.1: Bragg’s equation**

\[ d = \frac{\lambda}{2 \sin \theta} \]  

(3.1)

XRD is used to calculate this interlayer d-spacing [11].

### 3.2.3 Mechanical Testing

Before the experimental work could begin, the specimens needed to be prepared in accordance with the relevant ASTM standard. The following section describes the manufacturing process for the specimens for each of the tests, the test methods and the equipment that was utilized.

#### 3.2.3.1 Hardness Testing

The laminate created above was cut as per ASTM D 2583-07 [78] to produce specimens to be used for the hardness tests. The sample size here was 250x25x2mm. The hardness test was performed using Barcol Impessor GYZJ 934-1.

Verification was required with predetermined discs of values 45-48 and 87-89 before commencing the testing. The hardness check was done using the Barcol meter and if the value falls within this range then the unit is calibrated and testing can commence. The verification process needed to be done each time before using the equipment. Testing was then performed on the composite specimens of 0wt% to 5wt%. As per the ASTM standard, up to 10 checks needed to be performed on various points along the sample in order to get a true estimate of the hardness value of the entire sample.
Hardness measurements quantify the resistance of a material to plastic deformation. Indentation hardness tests compose the majority of processes used to determine material hardness, and can be divided into two classes: microindentation and macroindentation tests. Microindentation tests typically have forces less than 2 N (0.45 lbf). Hardness, however, cannot be considered to be a fundamental material property. Instead, it represents as pointed out by Meyers et al [79], an arbitrary quantity used to provide a relative idea of material properties.

3.2.3.2 Impact Testing:
The Impact test was performed using the Charpy Impact tester. Testing was done here also on a predetermined sample and if the value fell within the acceptable range, then testing commenced.

Impact testing laminate was created as per the ASTM standard D6110-04 [80]. The sample size here was 127x12.7x2mm. Specimens were cut and prepared using the circular saw and notches with an angle of 22° were inserted in the middle of the sample, using a precision scale, as per Figure 3.7.

Figure 3.8 shows the actual test equipment that was used to perform the Charpy test. The two arms at the top were released and the sample was broken. The force required to break the sample was recorded.

This is a standardized high strain-rate test which determines the amount of energy absorbed by a material during fracture. Tough materials absorb a lot of energy, whilst brittle materials tend to absorb very little energy prior to fracture [81].
Figure 3.7: Charpy test specimen, showing the “V” notch in the middle of the sample. [82]

Figure 3.8: The Charpy test equipment
3.2.3.3 Fibre Volume Fraction

Fibre volume fraction is the percentage of fibre volume in the entire volume of a fibre-reinforced composite material. Manufacturing processes for components made from thermoset composites usually require the optimal achievement of two physical properties. The first is the fibre volume fraction which determines the composites strength. The second property is the void content found within the matrix resin. During the lay-up process the presence of voids is inevitable. A high void content indicates that the resin has not thoroughly embedded the fibres, resulting in the deterioration of properties.

Test methods determine the constituent content of the composite materials by removing the matrix by digestion or ignition leaving the reinforcement essentially unaffected. This allows for an accurate determination of the fibre volume fraction.

Typical fibre volume fractions are shown in Table 1.1 below, which highlights that the optimum fibre volume fraction for woven carbon fibre is 45 %. However with the VARIM system, this value can get higher, up to 55%.

<table>
<thead>
<tr>
<th>Reinforcement form</th>
<th>Possible range of fibre volume fractions (%)</th>
<th>Typical value for fibre volume fraction (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unidirectional</td>
<td>50 – 70</td>
<td>60</td>
</tr>
<tr>
<td>Woven</td>
<td>35 – 55</td>
<td>45</td>
</tr>
<tr>
<td>Random Mat</td>
<td>10 - 30</td>
<td>20</td>
</tr>
</tbody>
</table>

Once the laminates have been processed, they are analyzed for fibre volume fraction. This can be carried out in two methods, one using the direct calculation using the actual volumes of the fibre and the resin volume utilized to create the laminate during processing.
Equation 3.2: Fibre volume fraction

\[
V_f = \frac{\text{Fiber weight}}{\text{Fiber + resin weight}}
\]  

(3.2)

Using Equation 3.2, the \( V_f \) was calculated to be 55.6%. The resin weight in the above equation incorporates the nanoclay and the hardener volumes as well.

The second method is doing an evaluation of the cured laminate and identifying the actual amount of the fibre and resin present. ASTM D3171 \([84]\) was followed for this procedure. Procedure A was followed which was the use of Nitric acid, HNO₃, to dissolve the resin and then calculate the amount of fibres left behind. Five samples were used for each nanoclay percentage to get a representative average value. The samples were dried in an oven at 50°C for 4 days, which was how long it took for the samples to reach an equilibrium weight. 100 ml beakers were filled with 30 ml Nitric Acid and the samples were immersed into this liquid. The beakers were then placed into a temperature bath at 80°C for 8 hours. At this point the resin had disintegrated and the fibres remained in the Acid. The fibres were then filtered using a vacuum suction system and washed with distilled water to remove any acid that may be present. These fibres were further dried to remove the water and then weighed. This was done for each nanoclay percentage from 0-5wt%.

Equation 3.2 was then used to calculate the fibre volume fraction. With this method the average values for each nanoclay percentage was between 49 – 53%.

The Void content was calculated using Equation 3.3:

Equation 3.3: Void content

\[
V_v = 100 - (V_f + V_m)
\]  

(3.3)

The void content value was negligible which implied that there were little to no voids present in the laminate itself.
3.2.3.4 Tensile Testing:
The equipment utilized for the tensile testing was the universal mechanical testing machine (MTS), Servo Hydraulic MTS 100 kN at the test rate of 5mm/min. Calibration is done biannually by an external company, IMP Calibration services and no further verification was required before utilising the machine for testing.

Tensile testing specimens were prepared using 10 sheets of carbon fibre. The 10 sheets were chosen as the optimum number of sheets after a trial and error process in order to obtain a thickness of 2 mm, as required by the ASTM standard 3039 [85]. Specimens were cut using a Computer Numerical Control (CNC) machine with utmost precision. Specimen sizes were 250x25x2mm. Tabs used were made of glass fibre. The test rate used was 5mm/min. 5 samples of each 0wt%, 1wt%, 3wt%, and 5wt% were tested and the averages were obtained. The sample was loaded between the jaws of the MTS machine and the load was continuously increased at a constant rate, until the sample stretched and reached its breaking point.

The data obtained was the extension and load values, from which the stress and strain were calculated. The modulus was then calculated from the stress and strain values.

The extension value, which was the distance that the sample was stretched was also recorded. The calculated tensile strength and modulus values were used to calculate the loading rate percentages for the fatigue tests.

3.2.4 Dynamic Testing

3.2.4.1 Fatigue testing specimen
Fatigue test specimens were prepared in a similar way as to the tensile test specimens and the test method used here was ASTM 3479 [86]. Emery paper, 50x50mm, rather than glass fibre tabs was used as tabs at the end of the specimens to ensure a firmer grip between the testing equipment jaws and hence
prevent slippage. The universal mechanical testing machine (MTS), Servo Hydraulic MTS 100kN was used to perform the tensile test.

Test specimens were cut using CNC precision cutting equipment, and the edges were sanded down a little to create rounded edges. Three specimens were tested for each fatigue cycle carried out. The tests were conducted at room temperature and at a stress ratio, \( R = \sigma_{\text{min}}/\sigma_{\text{max}} = 0.1 \). A test frequency of 3 Hz was used which was low enough to minimize the adiabatic heating effect [87]. To determine the S-N graphs, the maximum stress levels were kept at 90%, 75% and 60% of the corresponding ultimate tensile strength of the composite.

Table 3:2: Duration for each of the Fatigue loading percentages for each Nanoclay Percentage

<table>
<thead>
<tr>
<th>Nanoclay %</th>
<th>90% testing Duration, days</th>
<th>75% testing Duration, days</th>
<th>60% testing Duration, days</th>
</tr>
</thead>
<tbody>
<tr>
<td>0wt%</td>
<td>0.25</td>
<td>6</td>
<td>10</td>
</tr>
<tr>
<td>1wt%</td>
<td>0.25</td>
<td>10</td>
<td>15</td>
</tr>
<tr>
<td>3wt%</td>
<td>0.25</td>
<td>15</td>
<td>23</td>
</tr>
<tr>
<td>5wt%</td>
<td>0.14</td>
<td>12</td>
<td>30</td>
</tr>
</tbody>
</table>

It was decided to stop at 60% and not go to a lower fatigue loading percentage, due to the fact that the fatigue testing at lower loading takes too long and at lower loading percentages, the sample may plateau out and run for months at a time. The testing will have to be done in triplicate, so for one loading rate for one nanoclay percentage, the testing may span 6 months. Hence the fatigue percentage loading rates used here were 90%, 75% and 60%.

The same load rates were not used for each of the nanoclays percentages, since the fatigue loading percentages would not be consistent. It would in fact be lower and would hence be incorporated in the final results for the 90%, 75% and 60%.
This is an alternative test method that can be used for comparison purposes. However the test method utilised in this research work incorporates these values.

3.2.5 Failure Mechanism Analysis

3.2.5.1 Microscopy (Optical and Scanning)

The optical microscope, OM which is often referred to as the "light microscope", is a type of microscope which uses visible light and a system of lenses to magnify images of small samples. The image from an optical microscope can be captured by normal light-sensitive cameras to generate a micrograph. Originally images were captured by photographic film but modern developments in CMOS and later charge-coupled device (CCD) cameras allow the capture of digital images. Purely Digital microscopes are now available which just use a CCD camera to examine a sample, and the image is shown directly on a computer screen without the need for eye-pieces [88]. The OM was used on the fractured samples prior to the scanning electron microscope (SEM) work being undertaken. The OM gave a good visual of the damaged surface, and allowed the researcher to accurately pinpoint the areas to be observed under the SEM.

The Scanning Electron Microscope, SEM is an instrument that produces a largely magnified image by using electrons instead of light to form an image. The SEM has many advantages over traditional microscopes. The SEM has a large depth of field, which allows more of a specimen to be in focus at one time. The SEM also has much higher resolution, so closely spaced specimens can be magnified at much higher levels. Since the SEM uses electromagnets rather than lenses, the researcher has much more control in the degree of magnification. All of these advantages, as well as the actual strikingly clear images, make the scanning electron microscope one of the most useful instruments in research today [89]. The equipment used here was the JEOL JEM-7500F (Tokyo, Japan) operated at an accelerating voltage of 2 kV. The SEM was able to accurately provide details images of the fibre breakages, hairline crack propagation and of the delamination that occurred.
3.2.5.2 Effect of Temperature on Fatigue testing

The temperature was recorded at 3 different points on the sample itself while the fatigue test was being conducted. Temperature 1 was recorded at the top of the sample towards the top jaw, temperature 2 was recorded at the middle of the sample and temperature 3 was recorded at the bottom of the sample towards the movable lower jaw. Refer to Figure 3.9 for the schematic depicting the location of the different temperatures. An infrared thermometer was utilized to obtain the temperature of the sample. For the 90% fatigue loading the temperature was recorded at intervals of 5000 cycles and for the 75% and the 60% loading it was recorded at intervals of 50000 cycles. The initial temperature was when the fatigue testing was started and the final temperature was when the sample fractured. During testing, the bottom jaw of the MTS machine moved while the top jaw stayed fixed.

![Figure 3.9](image)

*Figure 3.9: Showing the location of the temperature check points along the sample during the fatigue testing*
3.2.5.3 Fatigue Crack Density Growth
The crack density of the fatigue specimens was investigated during testing. A 25 mm section of the test sample was monitored by visual inspection during the 60% fatigue loading and the crack initiation and growth was monitored on a daily basis.

3.3 SUMMARY OF CHAPTER
This chapter focuses on the preparation of the CFRP composites with the addition of nanoclays in the concentration of 0wt%, 1wt%, 3wt% and 5wt% using the vacuum assisted resin infusion method (VARIM). Static (tensile, Impact, hardness, double cantilever beam and end notch flexure) and dynamic test (fatigue) were carried out on the prepared specimens to obtain the life cycle for the sample. Characterisation and morphological studies were done via XRD, TEM and SEM.
CHAPTER 4

4.1 BACKGROUND

Carbon fibre is used in many applications, mainly because of its low weight to high strength ratio. The use of carbon fibre panels on aircrafts in particular have been steadily increasing throughout the years [3]. These aircraft structures are required to carry sufficient fuel for a single trip. If the composite panels are made lighter but with the same strength, then the aircraft can increase the number of passengers that it carries or the fuel consumption significantly decreases. Numerous studies show the benefit of creating a hybrid structure by adding nanoparticles into their composite panels, as they provide added strength. In this study the nanoparticle introduced into the composite is Cloisite® 15A, a Montmorillonite, MMT. This research study focuses on the fatigue of CFRP and examines the effect that the nanoclays have on the fatigue life of a composite structure.

Carbon fibre re-inforced plastic (CFRP) specimen were produced using the Vacuum Assisted Resin Infusion Method (VARIM). During infusion, a virgin sample with no nanoclay was produced and Cloisite® 15A nanoclays were added into the resin mixture, in 1wt%, 3wt%, and 5wt% concentrations to create different specimens of varying wt% of nanoclay concentration samples. Test samples were prepared and tests were according to ASTM standards. In this section the results and discussion of the research study are presented. Morphological analysis look at the X-Ray Diffraction and TEM analysis of the prepared test specimens. Static testing look at the tensile test, Barcol Hardness and Charpy tests conducted. Dynamic testing look at the tension-tension fatigue testing at room temperature.
4.2 MORPHOLOGICAL ANALYSIS

4.2.1 X-Ray Diffraction

The XRD curve of Cloisite 15A only and Carbon fibre composite infused with 1wt%, 3wt%, and 5wt% Cloisite®15A nanoclay is shown in figure 4.1. The graph for Cloisite®15A as per figure 4.1 shows a distinct peak at a 2θ value of 5.75° and a corresponding initial interlayer spacing of 18.069 Å. The properties of nanoclay dispersed polymer composites depend on the basal spacing of the nanoclay. As nanoclays were added into the composite it was seen that this peak decreased considerably, the polymer is able to enter into the basal spacing of the nanoclay and hence this spacing becomes larger.

![XRD Curves for Cloisite 15A and 0-5% Nanoclay % Resin](image)

*Figure 4.1: XRD spectra for Cloisite 15A, 1% - 5% Nanoclays*

The XRD analysis was used to determine the interlayer d-spacing as described in chapter 3, section 3.2.2.2. The d-spacing calculated for the 1wt%, 3wt% and 5wt% samples were 24.86, 25.72 and 24.86, respectively. Generally an increased spacing between the basal layers and a hydrophobic, organophilic surface make
it more likely for the polymer to enter between the layers of the clay [90]. This happens when an exchange between the organic ions and the natural clay particles takes place.

The peak that was present in the Cloisite 15A curve is however absent in the curves for the nanoclay samples. This indicated that the polymer has entered into the basal spacing of the nanoclay. The calculation of the basal spacing for the nanoclay samples showed that the value was higher than that for the Cloisite 15A only sample, hence this was an indication that the polymer had entered into the basal spacing.

An exfoliated structure is formed when individual platelets dispersed in the matrix are separated by distances greater than 8.8 nm and the platelets can be orientated, forming short stacks [91]. Since the d-spacing increase was greater than 8.8 nm for all the nanoclay samples, the arrangement can then be described as exfoliated. However, further analysis in the form of TEM needed to be performed to confirm this arrangement.

4.2.2 Transmission Electron Microscopy

Figures 4.2a, 4.2b and 4.2c show the TEM image of the 1wt%, 3wt% and 5wt% nanoclay infused specimens, respectively. Three arrangements are possible when a composite is infused with nanoclays. The sample can be intercalated, where the interlayer spacing between the platelets are 1 nm. The sample can be exfoliated when the individual platelets that are dispersed in the matrix are separated by distances greater than 8.8 nm and the platelets can be orientated, forming short stacks or tactoids [92]. An agglomerated clay structure is formed when no polymer penetrates the interlayer spacing thus forming clustered platelets. These structures are micron sized particles reminiscent of conventional micro composite filler materials.

From the XRD results in the above section 4.1, since the peak was absent in all the nanoclay samples, it can be safely stated that none of the samples have an
agglomerated clay structure. The image shown in Figure 4.2a shows a uniformly distributed 1wt% nanoclay sample. Small concentrated clusters and uniformly distributed groups are seen. The scattered lighter sections are the voids, which were created during the VARIM process. During the VARIM process, the resin is heated to 70°C and the nanoclays are added into this resin mixture. When the hardener was added, the reaction that took place was exothermic and as such, air bubbles were created. Although much care was taken to remove these bubbles, small bubbles were still present and these showed up as voids at a later stage. Since the distance between the platelets here is more than 8.8 nm, the structure can be said to be exfoliated.

![Figure 4.2a: TEM Image of 1wt% Nanoclay infused composite specimen](image)

Figure 4.2b shows the structure for the 3wt% nanoclay sample. Here the clusters are bigger than those of the 1wt% sample and also uniformly distributed groups. The distance here as well between the platelets is greater than 8.8nm, however there is agglomeration here.
Figure 4.3b: TEM Image of 3wt% Nanoclay infused composite specimen

Figure 4.2c show the TEM arrangement for the 5wt% sample. The arrangement in this sample was orderly and there were no voids present. The lines show an ordered stacked arrangement with clearly visible interspaces. This suggests the formation of an intercalated nanoclay structure.

Figure 4.4c: TEM Image of 5wt% Nanoclay infused composite specimen
The basal spacing calculated above during the XRD analysis show that the basal spacings are indeed larger than the Cloisite 15A alone. This showed that the polymer entered the layers and as such, the distribution of the nanoclays were uniform. The nanoclay arrangement for the 1wt% and the 3wt% is exfoliated and that of the 5wt% is intercalated.

4.2.3 Fibre Volume Fraction

Fibre volume fraction of the composite is an important part of determining the strength of the laminate. From literature [93] the optimum fibre volume fraction for woven fibre laminate for high ultimate tensile strength should be between 35 - 55%. As soon as the fibres are presented in a non-unidirectional form, such as woven fabrics or random mats, a proportion of the fibre reinforcement is oriented out of plane that is that the fibres are scattered when compared to plane of reference. The plane of reference is usually determined by the direction of the resin flow and as soon as the fibres move out of this plane, the fibres are regarded as being orientated out of plane.

Close packing is no longer possible, and the maximum fibre volume fraction is reduced. Hence the maximum is 55%. For the unidirectional mats, the fibre volume fraction can go up to 65%.

The fibre volume fraction was calculated in two ways. The first was direct calculation performed during the processing method. The amount of resin and fibre were carefully weighed to get best consistent fibre volume fraction value.

<table>
<thead>
<tr>
<th>Carbon fibre sheet weight (g)</th>
<th>Resin Weight (g)</th>
<th>Hardener (g)</th>
<th>Nanoclay (g) (5wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>363.00</td>
<td>271.50</td>
<td>10.95</td>
<td>7.26</td>
</tr>
</tbody>
</table>

Table 4:1: Data for determining the fibre volume fraction of CFRP
The calculation using the values above, gave a fibre volume fraction of 55.6%. To further confirm the fibre volume fraction a burn off test was carried out on the fully infused and cured laminate. ASTM D 3171 – 99 [84], was utilized for this test. The laminate was placed into nitric acid and the matrix was allowed to dissolve overnight at a temperature of 70°C. The resultant fibres were then vacuum filtered at 17kPa and dried in an oven for one hour. These fibres were then weighed and compared to its original weight. This value was calculated to be 55%. This 1.07% difference was acceptable.

### 4.3 QUASI-STATIC TESTING

#### 4.3.1 Tensile

Tensile testing specimens were prepared and testing were carried out as per ASTM D3039 [85]. Five samples of each nanoclay percentage was tested and analyzed. The ultimate tensile strength was required from this test so that the fatigue testing could be carried out. The stress strain graph was plotted as shown in Figure 4.3 below.

The stress-strain graph for CFRP with 0wt%, 1wt%, 3wt%, and 5wt% nanoclay concentration all displayed a typical viscoelastic stress strain curve behavior. As the nanoclay percentage increased the stress and strain factors that the sample experienced increased as well. The higher nanoclay weight percentage sample, that was the 5wt% sample showed a 36% better result in terms of tensile stress, when compared to the 0wt% sample.

The Figure 4.3 shows similar trends for all four samples and the distinct regions characteristic of tensile testing is not clearly evident here. There is no adjustment period, however the sample continues to be strained and migrates into the elastic region. Once the ultimate stress has been reached, the sample fractures.
At the point of failure for the samples, the only difference in them is the concentration of the nanoclays. At the point of failure, it is these nanoclays that are responsible for the strain being at a higher value.

Figure 4.4(a) and 4.4(b) are the actual tensile tested samples for 0wt% and 5wt% samples, respectively. The sample in Figure 4.4(a) has a clean and neat edge at the point of fracture, as indicated by the circle on the figure. For this to happen, it meant that the sample was uniformly stressed at the same point and that the matrix holding the sample together fractured easily.

This type of fracture was similar to that of brittle materials where cracks spread rapidly, with little or no deformation and fracture may be sudden and catastrophic. This also shows that there was little strain was that effected on the sample and it then broke clean and neatly.
Figure 4.6: a) Tensile fractured sample of virgin nanoclay specimen

In comparison, Figure 4.4(b) showed a different scenario where the fractured edge was not neat and clean as that for the 0wt% sample. It was however jagged and uneven and the fracture was not in a straight line. There was also evidence of fibre separation and delamination that took place. In order for this to happen, it highlighted that the composite was highly stressed, however it did not fracture at the exact point where the stress was, which was in the middle of the sample. The matrix took the strain and deteriorated first, while still keeping the fibres in place. As the stress continued, the fibres still stayed intact and separated as sheets of fibre. It was at this point where ultimately the fibre strength determines the strength of the composite as a whole and the fracture took place.

The strain that the 5wt% sample incurred was twenty percentage higher than that of the 0wt% sample.

Figure 4.7: b) Tensile fractured sample of 5wt% nanoclay specimen
The ultimate tensile strength curve shown in Figure 4.5, showed a similar trend as with that in Figure 4.4 above. There was a continuous increase in the strength from the 0wt% sample to the 5wt% sample. The percentage increase was 34% from 0wt% to 5wt%.

The extension indicated the samples ability to be pulled during the process of the tensile testing. From Figure 4.6 it can be seen that as the nanoclay percentage in the sample increased, the extent to which the sample can be stretched, also increased. The strain to failure that the 5wt% sample was able to withstand enabled it to have a higher strength than the 0wt% sample. This highlights the crack arresting mechanism of the nanoclays and shows the nanoclays enhanced capabilities.

![Ultimate tensile testing graph](image)

*Figure 4.8: Ultimate Tensile Strength graph for CFRP at 0wt%, 1wt%, 3wt%, and 5wt% nanoclay concentration*
Figure 4.9: Extension graph for CFRP at 0wt%, 1wt%, 3wt%, and 5wt% nanoclay concentration

Table 4.2 is a summary of the static tests that were carried out. As the nanoclay percentage increased so too did the values for each of the static testing performed.

Table 4.2: Summary of Average Tensile testing results for each nanoclay concentration

<table>
<thead>
<tr>
<th></th>
<th>Modulus</th>
<th>UTS</th>
<th>Extension</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>GPa</td>
<td>Increase%</td>
<td>MPa</td>
</tr>
<tr>
<td>CF + 0wt% NC</td>
<td>44.68 ± 0.6</td>
<td>594.19 ± 10</td>
<td>3.01</td>
</tr>
<tr>
<td>CF + 1wt% NC</td>
<td>46.53 ± 0.5</td>
<td>648.96 ± 12</td>
<td>2.85</td>
</tr>
<tr>
<td>CF + 3wt% NC</td>
<td>48.30 ± 0.7</td>
<td>671.67 ± 11</td>
<td>2.80</td>
</tr>
<tr>
<td>CF + 5wt% NC</td>
<td>50.76 ± 0.4</td>
<td>795.73 ± 8</td>
<td>2.52</td>
</tr>
</tbody>
</table>

From the parameters examined from the tensile testing conducted it can be established that the 5wt% specimen could withstand the most amount of load and
stress before failing. This failure was catastrophic when compared to the other specimens. These observations can be attributed to the inclusion of the nanoclays.

4.3.2 Barcol Hardness

The Barcol test was carried out to provide the hardness value of the sample which would assist to explain the mechanism with which the sample failed during testing. Three specimens of each the 0wt%, 1wt%, 3wt%, and 5wt% were tested and there were ten test points tested along the entire length of each sample. These results are shown in Figure 4.7 below.

The hardness of the 0% nanoclay sample, which was the virgin sample, was the lowest and hence could be seen as the softer specimen. As the tests progressed through to the 5% sample, these specimens’ values got progressively higher and hence this implies that the samples were harder.

![Barcol Hardness Testing](image)

*Figure 4.10: Barcol Hardness testing values CFRP for 0wt%, 1wt%, 3wt%, and 5wt% nanoclay concentration*
According to Pavlini et al. [94], indentation hardness correlates linearly with the tensile strength. This was very evident in the results shown in Figure 4.7. The Barcol tester tests the surface of the sample and stopped when there was an indentation on the surface of the specimen. The nanoclays are bonded to the fibre and the matrix intrinsically, this was evident in the values obtained for the Barcol test. It takes a higher value for the indenter to create an indent in the specimen as the nanoclay concentration increased.

Meyers et al. [79] stated that the hardness can only offer a comparative idea of the material's resistance to plastic deformation and this was what will determine the brittleness of the specimen. From the tests carried out with the Barcol tester, it could be concluded that as the nanoclay concentration in the specimens increased, so too did the hardness value and hence as the nanoclay percentage in the specimen increased, the specimen becomes brittle.

4.3.3 Charpy Impact

The impact test can be identified as the amount of energy absorbed by a specimen during fracture. As the specimen was hit or “impacted”, it absorbed the energy to the point where it could not absorb anymore and then this impact continued to break the specimen. The sample size for this test tended to be higher than the five recommended, as per Mills [95], due to the possible inconsistencies in the notch angle, notch size and the sample size that can greatly affect the results. The number of samples used here was six samples, as recommended by Mills.

The graph for the Impact testing as shown in Figure 4.8 showed a downward trend from 0wt% to 5wt% nanoclay concentration. This trend was opposite to that of the Tensile and the Barcol testing. The 5wt% specimen test had the lowest value. The nanoclays present in the specimens in the matrix and have left little room for the absorption of any energy. The 5wt% specimens fractured off neatly.
at the 22° angle notch, while the other nanoclay percentage specimens broke in several pieces and flew off into different directions.

The qualitative results of the impact test can be used to determine the ductility of a material, according to Mathurt et al [96]. If the material broke on a flat plane, the fracture was brittle, and if the material broke with jagged edges or shear lips, then the fracture was ductile. Usually a material does not break in just one way or the other, and thus comparing the jagged to flat surface areas of the fracture would give an estimate of the percentage of ductile and brittle fracture [94].

![Impact testing chart](image)

**Figure 4.11:** Impact testing of CFRP for 0wt%, 1wt%, 3wt%, and 5wt% nanoclay concentration

Each of the six specimens tested for the 5wt% samples, had neat, clean breaks and as per Mathurt et al [96], this is synonymous of a brittle sample. Figure 4.9 shows the 5wt% Impact sample with the V-Notch before and after the testing was carried out. Figure 4.10 shows the 0wt% and 1wt% fractured sample after testing.
The fractured edge was straight, which supports Mathurt et al. theory about the higher nanoclay samples, in this case the 5wt% sample, being brittle. When compared to Figure 4.9, the fractured edges in Figure 4.10 can be seen to be jagged and have rough fractured edges. The impact tests further confirm that as the nanoclay percentage increased, so too did the brittleness of the composite. The results for the Barcol hardness and the Impact Testing results are tabulated below. There was a percentage increase between each nanoclay percentage for all the results.
Table 4.3: Average Barcol Hardness and Impact Results

<table>
<thead>
<tr>
<th>Nanoclay %</th>
<th>Barcol Hardness</th>
<th>% Increase</th>
<th>Impact (J)</th>
<th>% Increase</th>
</tr>
</thead>
<tbody>
<tr>
<td>0wt%</td>
<td>58.80</td>
<td></td>
<td>4.75</td>
<td></td>
</tr>
<tr>
<td>1wt%</td>
<td>59.23</td>
<td>1%</td>
<td>4.52</td>
<td>5%</td>
</tr>
<tr>
<td>3wt%</td>
<td>61.05</td>
<td>3%</td>
<td>4.23</td>
<td>6%</td>
</tr>
<tr>
<td>5wt%</td>
<td>67.20</td>
<td>10%</td>
<td>3.84</td>
<td>9%</td>
</tr>
</tbody>
</table>

4.4 DYNAMIC TESTING: FATIGUE

With only very low fractions of clay, including the enhanced thermal stability [97, 98], clay-epoxy nanocomposites showed a wide array of property improvements, reduced moisture and gas permittivity [99] and superior flame retardancy [100]. The nanoclay, in particular, which was Cloisite15A, exhibited ameliorating effects on fracture and fatigue resistance of CFRPs.

4.4.1 Tension-Tension Fatigue Testing

The S-N graphs for each of the fatigue loading that was the 90%, 75% and the 60% are combined and are represented below in Figure 4.11.

For each of the fatigue loading percentages, the mean of the five samples tested, were plotted and error bars are included to show the scatter. A trend line was passed through the points, in order to obtain an accurate as possible trend for each fatigue loading. The 0wt% sample fractured at a lower load when compared to the 1wt%, 3wt% and 5wt% samples. The 1wt% and 3wt% samples fractured at higher cycles when compared to the 5wt% sample. The 5wt % specimen became progressively better as the fatigue loading percentage decreased and this resulted in the 5wt% sample having a highest fatigue life cycle. This is a desirable quality in all products.
Figure 4.14: S-N graph for 90%, 75% and 60% fatigue loading for 0wt%, 1wt%, 3wt% and 5wt%

Table 4.4 shows the mean cycles for each fatigue percentage loading and for each nanoclay percentage with their deviations.

Table 4.4: Fatigue Results showing the number of cycles for 90%, 75% and 60% fatigue loading

<table>
<thead>
<tr>
<th>Nanoclay %</th>
<th>90% Fatigue Loading</th>
<th>75% Fatigue Loading</th>
<th>60% Fatigue Loading</th>
</tr>
</thead>
<tbody>
<tr>
<td>0wt%</td>
<td>35110 ± 1018</td>
<td>931518 ± 58951</td>
<td>1830476 ± 17929</td>
</tr>
<tr>
<td>1wt%</td>
<td>46764 ± 586</td>
<td>989518 ± 22206</td>
<td>2640130 ± 155033</td>
</tr>
<tr>
<td>3wt%</td>
<td>58904 ± 717</td>
<td>1850969 ± 74517</td>
<td>3537201 ± 171714</td>
</tr>
<tr>
<td>5wt%</td>
<td>2286 ± 70</td>
<td>961580 ± 59963</td>
<td>5373984 ± 142676</td>
</tr>
</tbody>
</table>
At the 90% fatigue loading, the number of cycles for each nanoclay percentage sample increased until the 3wt% sample, which had a maximum life cycle of 58904 cycles. The 5wt% sample fractured early at 2286 cycles. This was attributed to the brittle quality of the CFRP as the nanoclay concentration increased. Note that five samples were tested for the 5wt% sample and all five samples behaved in a similar manner.

At the 75% loading, a similar trend was observed where the 3wt% sample again performed better than the 0wt% and 1wt% nanoclay percentages. At the 5wt% sample, the number of cycles to failure dropped again as with the 90% fatigue loading. The number of cycles to failure and the fracture mechanism for the 5wt% sample was not sufficient enough to determine that the sample exhibited brittle fracture.

At the 60% fatigue loading, the 5wt% sample ran for the most amount of cycles, performing 52% better than the 3wt% sample and 194% better than the 0wt%. A graphical representation of the time of fracture within the fatigue life cycle of the sample was depicted in the diagram in figure 4.12 below.

All the 90% loading samples fractured less than the 100K life cycle mark. For the 75% loading, it could be noticed that the number of fatigue cycles start in the 1M life cycle region. It should be noted that the 3wt% sample fatigue life was the best in the 75% loading. In the 60% loading the samples follow a trend as per the tensile testing, where they become progressively better as the nanoclay percentage increased.
**Figure 4.15:** Fatigue Cycle Failure at 90%, 75% and 60% loading at a constant frequency of 3Hz for 0, 1, 3 and 5 wt% nanoclay specimens

Taking out the effect of the specimen area, it could be seen that Figure 4.13 looks very similar to that in Figure 4.11.

The graph for the 0wt% and the 1wt% are almost parallel in the normalized graph. And a similar trend is observed for the 3wt% and the 5wt% when compared to figure 4.11. It could therefore be stated that the above graph can be useful for any sized CFRP sample. The results presented above are the actual number of cycles to fracture for each of the fatigue loading rates and for each of the nanoclay percentages.
A discussion will follow which will discuss the mechanisms at which these samples fracture. Degrieck et al. [101] stated that fatigue failure in conventional composites did not occur in a localized fashion but throughout the volume of the material. Damage in such composites took the form of fibre breakage, matrix cracking, debonding, transverse ply cracking, and delamination, and largely depended on the method used in manufacturing the composite. These processes occurred sometimes interactively or independently. For a detailed analysis of this, the crack growth, propagation and arrest will be examined further.

4.4.2 The effect of Frequency and Temperature
The loading frequency used for the fatigue testing was 3Hz. The loading frequency is considered to have a significant influence on the sensitivity to fatigue for carbon fibre composites for the effect of self-generated heating and intrinsic rate dependence, as stated by Curtis et al. [68]. The surface temperature of
specimens increase rapidly in the intermediate number of cycle range of 10,000 cycles and the increase in surface temperature was larger than 20°C, as observed by Pandita et al [69]. The magnitude of the change of the surface temperature depended on the fibre orientation as well as maximum fatigue stress. It was found that 2Hz offered the most consistent temperature profile, while 10Hz offered the highest temperature profile range.

The temperature was recorded at 3 different points on the sample itself while the fatigue test was being conducted. Temperature 1 was recorded at the top of the sample towards the top jaw, temperature 2 was recorded at the middle of the sample and temperature 3 was recorded at the bottom of the sample towards the movable lower jaw. Table 4.5, 4.6 and 4.7 show the surface temperature at points 1 (Top Jaw), 2 (Middle Jaw) and 3 (Bottom Jaw) for each fatigue percentage loading rate and for each nanoclay percentage.

Table 4.5: Temperature increase of the test sample during Fatigue Testing – Temperature 1

<table>
<thead>
<tr>
<th>Sample Temperature Increase (Temp 1)</th>
<th>90%</th>
<th>75%</th>
<th>60%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature, °C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Final</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>% ↑</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0wt%</td>
<td>25.2</td>
<td>25.8</td>
<td>2%</td>
</tr>
<tr>
<td>1wt%</td>
<td>24.5</td>
<td>25.1</td>
<td>2%</td>
</tr>
<tr>
<td>3wt%</td>
<td>24.7</td>
<td>25.7</td>
<td>4%</td>
</tr>
<tr>
<td>5wt%</td>
<td>25.3</td>
<td>25.9</td>
<td>2%</td>
</tr>
</tbody>
</table>
Table 4.6: Temperature increase of the test sample during Fatigue Testing – Temperature 2

<table>
<thead>
<tr>
<th>Sample Temperature Increase (Temp 2)</th>
<th>90%</th>
<th>75%</th>
<th>60%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature, °C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial</td>
<td>Final</td>
<td>% ↑</td>
<td>Initial</td>
</tr>
<tr>
<td>0wt%</td>
<td>25.2</td>
<td>27.8</td>
<td>10%</td>
</tr>
<tr>
<td>1wt%</td>
<td>24.5</td>
<td>27.0</td>
<td>10%</td>
</tr>
<tr>
<td>3wt%</td>
<td>24.7</td>
<td>27.6</td>
<td>12%</td>
</tr>
<tr>
<td>5wt%</td>
<td>25.3</td>
<td>26.0</td>
<td>3%</td>
</tr>
</tbody>
</table>

Table 4.7: Temperature increase of the test sample during Fatigue Testing – Temperature 3

<table>
<thead>
<tr>
<th>Sample Temperature Increase (Temp 3)</th>
<th>90%</th>
<th>75%</th>
<th>60%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature, °C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial</td>
<td>Final</td>
<td>% ↑</td>
<td>Initial</td>
</tr>
<tr>
<td>0wt%</td>
<td>25.2</td>
<td>28.3</td>
<td>12%</td>
</tr>
<tr>
<td>1wt%</td>
<td>24.5</td>
<td>27.3</td>
<td>11%</td>
</tr>
<tr>
<td>3wt%</td>
<td>24.7</td>
<td>28.0</td>
<td>13%</td>
</tr>
<tr>
<td>5wt%</td>
<td>25.3</td>
<td>26.1</td>
<td>3%</td>
</tr>
</tbody>
</table>

As the fatigue loading rate decreased the surface temperature of the sample increased and it can be seen that this trend occurred for all the tests carried out. The increase in surface temperature at point 3, which was the stationary jaw was the highest. The maximum temperature increase was noted to be 12°C. As observed by Pandita et al [69], the surface temperature increase of 20°C was sufficient enough in order to effect the test sample and hence the results. Since the surface temperature increase for the samples tested in this research work
was only 12°C, it can be stated that the temperature effect did not affect the fatigue results.

The surface temperature rate increase was 2.6°C/hr for 90% fatigue loading, 0.135°C/hr for 75% fatigue loading and 0.046°C/hr for the 60% fatigue loading. This low rate of temperature increase, was negligible to effect any changes to the fatigue results.

Figure 4.14, below show the temperature increase at each fatigue loading rate at the three different temperature test points on the jaw of the MTS Machine. The figure show the increase in the surface temperature of the test sample being the highest at the lowest fatigue percentage loading rate and at the bottom stationary jaw. The 60% fatigue loading has the total highest temperature increase due to the test running for an average duration of 21 days.

![Temperature Increase Chart](image.png)

**Figure 4.17:** Showing temperature increase during fatigue testing

The nanoclay arrangement of the 1wt% and the 3wt% is exfoliated and that of the 5wt% sample is intercalated, as discussed in Chapter 4, section 4.2.2. The
intercalated nanoclay arrangement, and in this case, a brittle material, performs better during 60% fatigue loading, than the 75% and 90% fatigue loading. It can be inferred that a slow rate of heat generation and the high fatigue loading - does not favour brittle materials hence the short fatigue cycle at the 90% fatigue loading and that a slow rate of heat generation and low fatigue loading of 60% - favours brittle materials.

4.4.3 Fatigue Crack Growth

Crack propagation during fatigue loading is one of the primary causes of failure in engineering materials. While this was inevitable, it was helpful to ascertain a trend that will determine the long term effects and life span of materials that undergo fatigue loading. Mandel [102] postulated a way to calculate the degradation rate of the samples, by calculating the slope first, using the fatigue data and then by calculating the percentage degradation from this. In order to calculate the slope the following equation was used:

\[
\text{Equation 4.1: Fatigue Slope Calculation} \quad b = \frac{\sigma_{\text{max}} - \sigma_{\text{UTS}}}{\log N_f} \quad [4.1]
\]

Where \( \sigma_{\text{max}} \) is the maximum stress, \( \sigma_{\text{UTS}} \) is the stress at the various fatigue loading rates, \( b \) is the slope and \( N_f \) is the number of fatigue cycles at the point of fracture.

Table 4.8 shows the values for the slope of the fatigue curves. It can be noted that as the nanoclay percentage increased within each fatigue loading, the slope of the curve also increased. Thereafter the degradation rate was calculated. For the 90% fatigue loading the rate decreased until the 3wt% sample and then it increased at the 5wt% sample. This tied up with the 5wt% sample fracturing earlier than the other samples. For the 75% sample there was a similar trend. For the 60% fatigue testing, it could be seen that as the nanoclay percentage increased, the rate of degradation decreased, even in the 5wt% sample. From the
0wt% sample to the 5wt% samples, there was a 7.44% decrease in the rate of degradation.

Table 4.8: Degradation rates of fatigue samples

<table>
<thead>
<tr>
<th></th>
<th>90%</th>
<th>75%</th>
<th>60%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>b (Slope)</td>
<td>σuts /b</td>
<td>b (Slope)</td>
</tr>
<tr>
<td>0 wt%</td>
<td>13</td>
<td>2.44%</td>
<td>25</td>
</tr>
<tr>
<td>1 wt%</td>
<td>14</td>
<td>2.36%</td>
<td>27</td>
</tr>
<tr>
<td>3 wt%</td>
<td>14</td>
<td>2.33%</td>
<td>27</td>
</tr>
<tr>
<td>5 wt%</td>
<td>21</td>
<td>3.00%</td>
<td>34</td>
</tr>
</tbody>
</table>

If the curves for each of the nanoclay percentages were extrapolated to include lower fatigue loading percentages, it could be seen that the lower percentages, 0wt%, 1wt% and 3wt% will reach a finite life cycle value. However the 5wt% sample would continue and eventually plateau, giving the 5wt% sample at the 60% fatigue loading an infinite life cycle.

4.4.4 Fatigue Crack Density Growth

The crack density of the fatigue specimens were investigated during testing. A 25 mm section of the test sample was monitored by visual inspection during the 60% fatigue loading and the crack initiation and growth was monitored on a daily basis. The results are presented below in Figure 4.15.

The rate of crack growth in the 0wt% sample was faster than that in the 1wt% and 5wt% samples. Although the crack density of the 1wt% and 3wt% are lower at lower cycle levels, they slowly pick up as the number of cycles continue. The slow rate of the crack density can be attributed to the inclusion of the nanoclays into the samples. All samples were prepared in the same way the only difference between them was the percentage of nanoclays within the samples.
At the time of fatigue fracture, which was at 2.5 million cycles, the 1wt% sample was observed to have 55 cracks within the sample size selected for investigation. These cracks propagated at a higher rate through the sample and caused the fracture. The 3wt% sample was observed to have 52 cracks at the same number of cycles that the 1wt% sample fractured. This was 6% less cracks at the same time. However, the cracks continued to increase and grow until the fracture point at 3.2 million cycles. At the same time, the 5wt% sample had 50 cracks observed, this corresponded to a 4% drop when compared to the 3wt% sample and 12% drop when compared to the 1wt% samples. The cracks in the 5wt% sample still continued to grow and propagate till it fractured at 5.5 million cycles. The calculation for the number of cracks observed here were average values for each of the respective nanoclay percentages.

*Figure 4.18: Fatigue Crack Density Growth for 0wt%, 1wt%, 3wt% and 5wt% at 60% fatigue loading*
4.4.5 Crack Initiation and Propagation

Before a crack can initiate there would be prior toughening mechanisms that will take place, like crack deflection, micro crack toughening, transformation toughening, crack bridging and particle toughening. These phenomena halt the crack initiation, until such time that the stress becomes too large for these phenomena to overcome, and a crack is initiated.

See Figure 4.16 for the tension-tension fatigue specimen for 1wt% at 60% fatigue loading. After 1.6 Million cycles and after ~seven days, visible cracks could be seen, towards one edge only. The white marks are representative of micro-cracks that form beneath the surface and that has travelled through the fibre layers and the matrix and has surfaced. After each day, the amount of white lines increased until after eleven days, the sample eventually fractured. The micro cracks that occurred coalesced with each other to form larger cracks and these propagated through the sample. The 1wt% sample was not as densely packed with nanoclays as the 5wt% specimen, so the cracks were able to initiate and propagate to the surface, with relative ease.

![Figure 4.19: Crack Initiation and propagation of 1wt% specimen at 60% fatigue loading](image-url)
Figure 4.17 displays the specimens for the 5wt% sample at various intervals of fatigue cycles. The micro-cracks here appear a lot more abundantly as when compared to the 1wt% sample. These micro cracks delay in the coalescing of larger cracks and this delay can be attributed to the nanoclays present in the matrix. The 5wt% sample was densely packed with the nanoclays and when the cracks propagate, they make contact with a nanoclay and the crack was arrested. As the number of cycles increased, the cracks were able to propagate through or around the nanoclay and reach the surface where they formed larger cracks. Here as well the nanoclays served to delay the process and eventually the cracks propagated and the sample fractured.

![Image of specimens at various intervals of fatigue cycles](image)

**Figure 4.20: Crack Initiation and propagation of 5wt% specimen at 60% fatigue loading**

Fatigue occurs when a material is subjected to repeated cyclic loading and unloading. If the loads are above a certain threshold, microscopic cracks will begin to form at the stress concentrators such as the surface, persistent slip bands (PSBs), and grain interfaces. Kim and Laird [103] stated that, a crack will
initiate, the crack will grow and eventually the crack will reach a critical size, the crack will propagate, and the structure will fracture.

The 1wt% sample when compared to the 5wt% sample, shows that the cracks initiate and propagate quickly. The cracks as shown in the SEM image in Figure 4.18a also propagated in straight lines, this showed that the sparsely distributed nanoclays did not hinder the crack path. And that these cracks occurred even through the woven structure, in a straight line.

The cracks in the 5wt% sample in Figure 4.18b, deflected in different directions and the cracks also caused delamination through the fibre layers. Ritchie [104] stated that crack propagation was a mutual competition between intrinsic microstructural damage mechanisms, which promoted crack extension ahead of the tip, and extrinsic crack-tip shielding mechanisms, which act primarily behind the tip to retard crack growth. In light of this phenomena, in Figure 4.18b, the crack can be seen to be arrested and moved into different directions and even into the fibre layers below the surface.

*Figure 4.21: a) 1wt% and b) 5wt% sample showing crack propagation through the matrix*
Figure 4.19a highlighted the same phenomena but this time there are multiple cracks. The crack highlighted by the red circle can be seen to be arrested, however due to the ongoing stresses created by the cyclic fatigue motion, a new crack appeared next to it.

![Figure 4](image)

**Figure 4.22: 5wt% sample showing a crack being arrested and a new crack being created**

Figure 4.20 shows micro crack sites and in the middle there was an example of the crack being arrested and deviated. Since the 5wt% sample was densely packed with nanoclays, this could be a site where a nanoclay was present which arrested the crack and then the crack had to move around the nanoclay. SEM evidence of this phenomena could not be obtained, however, an EDX was performed at this point, as showed in Figure 4.20 which showed a high concentration of silica. Silica is a bulk component of nanoclays, so it can be inferred that the nanoclay was at the point that the crack deviated and it is due to this nanoclay that the crack re-routed. The high concentration of Titanium, Ti is from the Tipex which was used to mark the point on the sample.
Figure 4.23: EDX of the 5wt% sample showing the composition of elements at the site where the crack was arrested.

Through the discussion and images presented above, it can be seen that cracks form at a faster rate through the lower percentage nanoclay samples and that the cracks propagate in a straight line. However at higher percentages, the crack propagation was arrested and at times new cracks were formed in that same vicinity or the crack re-routed around that point of obstruction. The point at where this phenomena occurs is the point where there were nanoclays present.
4.4.6 Fatigue Failure Mechanism

Figure 4.21, represents a failure envelope for the different types of fracture that occurred during the fatigue testing. Only three samples for each fatigue loading and each nanoclay percentage are reported here. The final fracture mode only was categorized here. The quadrants were broken up into three areas, Fibre Breakage, Matrix Failure and Delamination.

The 90% fatigue loading failure was characterized mostly by fibre breakage and matrix failure and as the nanoclay percentage increased, the extent of fibre breakage and matrix cracking decreased and the trend moved towards delamination, where the fibre sheets separated from each other. The repeated cyclic stresses imposed on the samples by fatigue testing causes the layers to separate, forming separate layers of the fibre.

The 75% fatigue loading was characterized mostly with matrix failure at the lower nanoclay percentages of 0wt% and 1wt% and some delamination at the 3wt%
and 5wt%. At the 60% loading, all the nanoclay samples were characterized by delamination.

Consequently, it can be summarized that as the fatigue loading decreased, from 90% to 60%, the failure mechanism moved from matrix cracking to delamination. As the nanoclay percentage increased to 5wt%, the failure mechanism was predominantly delamination. This is important to note, since delamination is an indication of the strength of the material.

4.4.7 Statistical Analysis
For the fatigue testing, three samples each for each fatigue loading percentages were tested, for each nanoclay percentage. Each test result was different from each other. In order to determine if this variation was adequate, a statistical analysis ANOVA was performed. In this analysis, the mean of each of the results
within each nanoclay concentration and each fatigue loading rate were analyzed. The results are tabulated in table 4.9 below.

Table 4.9: Summary of ANOVA analysis for fatigue results

<table>
<thead>
<tr>
<th>Fatigue Loading rate</th>
<th>Significant Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>90%</td>
<td>0.000</td>
</tr>
<tr>
<td>75%</td>
<td>0.000</td>
</tr>
<tr>
<td>60%</td>
<td>0.000</td>
</tr>
</tbody>
</table>

The significance value for each of the fatigue loading rate, was 0.000, this implies that there was no significant difference in the mean values.

Table 4.10 below shows the ANOVA analysis between each nanoclay percentage and fatigue loading. The mean values, the standard deviation and the standard errors are shown for each of the tests performed.

Table 4.10: ANOVA Analysis between the each fatigue loading and nanoclay percentage:

<table>
<thead>
<tr>
<th>NC %</th>
<th>90% Fatigue Loading</th>
<th>75% Fatigue Loading</th>
<th>60% Fatigue Loading</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>Std Dev</td>
<td>Std Error</td>
</tr>
<tr>
<td>0wt%</td>
<td>35070</td>
<td>3123</td>
<td>1803</td>
</tr>
<tr>
<td>1wt%</td>
<td>45827</td>
<td>2012</td>
<td>1162</td>
</tr>
<tr>
<td>3wt%</td>
<td>58501</td>
<td>1996</td>
<td>691</td>
</tr>
<tr>
<td>5wt%</td>
<td>2140</td>
<td>261</td>
<td>152</td>
</tr>
</tbody>
</table>

Figure 4.22 shows graphs for each of the fatigue loading, for each of the nanoclay percentages. For the 90% fatigue loading, the 3wt% had the highest fatigue cycles and the 5wt% the lowest, for the 75% fatigue loading, the 3wt% had the highest fatigue cycles and the 5wt% the lowest and for the 60% fatigue loading,
the 5wt% had the highest fatigue cycles and the 0wt% the lowest fatigue cycles. It can be inferred that at the lower fatigue loading rates, the 3wt% performs the best and for the high fatigue loading rates, the 5wt% performs the best.

Figure 4.26: ANOVA Analysis results for each Fatigue loading and Nanoclay Percentage

The breakdown of the values for each fatigue loading rate, was as shown in Table 4.11. As a rule, if the significant difference is greater than 0.05, this implies that there was a significant difference in the results. From table 4.10, it can be seen that the ANOVA analysis showed that there is no significant difference between the values.
Table 4.111: Multiple Comparisons for each of the fatigue loading

<table>
<thead>
<tr>
<th></th>
<th>90%</th>
<th></th>
<th>75%</th>
<th></th>
<th>60%</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Tukey</td>
<td>LSD</td>
<td>Tukey</td>
<td>LSD</td>
<td>Tukey</td>
<td>LSD</td>
</tr>
<tr>
<td>0% - 1%</td>
<td>0.001</td>
<td>0.000</td>
<td>0.003</td>
<td>0.006</td>
<td>0.023</td>
<td>0.005</td>
</tr>
<tr>
<td>0% - 3%</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>0% - 5%</td>
<td>0.000</td>
<td>0.000</td>
<td>0.005</td>
<td>0.007</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>1% - 3%</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.007</td>
<td>0.002</td>
</tr>
<tr>
<td>1% - 5%</td>
<td>0.000</td>
<td>0.000</td>
<td>0.005</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>3% - 5%</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
</tr>
</tbody>
</table>
CHAPTER 5

Carbon Fibre Re-inforced plastic composite laminates were produced via the VARIM method. During this process, Cloisite 15A nanoclays were added to the resin mixture to be infused into the composite. Nanoclays were added in the concentration of 0wt%, 1wt%, 3wt% and 5wt%.

The objective of this study was to investigate the effect of nanoclay concentration on the fatigue failure properties at fatigue loading rates of 90%, 75% and 60% and investigate the failure mechanisms.

- XRD proved that the distribution of the nanoclays within the matrix was evenly distribution. The 1wt% showed exfoliated structures, the 3wt% showed a mixture of exfoliated, flocculated and intercalated structures and the 5wt% showed intercalated structures.
- Tensile testing showed that the nanoclay percentage played a role in the strength of the composite and that as the nanoclay percentage increased, so too does the strength of the composite.
- It was found that the 5wt% sample became progressively stronger as the fatigue percentage loading rate moved from 90% to 75% to 60% and the specimen was able to withstand longer cycle times as opposed to the 0wt%, 1wt% and 3wt% nanoclay samples. This was attributed to the slow increase in the temperature which favors intercalated structures.
- The lines for the 0wt% and 1wt% on the S-N diagram were parallel to each other and very steep. This indicated a higher rate of fatigue degradation during the cyclic loading. The lines for the 3wt% and the 5wt% were gentle slopes, which indicated low rates of degradation.
- The 5wt% at 90% loading behaved similar to that of a brittle substance and this brittleness was proved by performing the hardness and impact testing.
- However as the fatigue loading percentage became lower and hence the force became lower, the 5wt% sample performed progressively better than the
other samples and had cycles of up to 5.5M. These results were attributed to the inclusion of the nanoclays.

- The damage mechanism examined using the scanning electron microscope showed an array of types of fractures, at the different percentage loading rates and at the different nanoclay percentages.
  - For the 90% fatigue loading, the 0wt% and 5wt% showed similar fracture characteristics. Both fractured along a straight line, with neat lines. However the 1wt% and 3wt% showed delamination, matrix failure and fibre breaking.
  - For the 75% fatigue loading, 0wt% and 3wt% were similar where fracture occurred along a fairly straight line and debonding was evident. The 3wt% and 5wt% both showed signs of extensive delamination with the 5wt% individual fibre strands, fracturing in different directions, whereas the 1wt% sample delaminated and fractured along a single plane, only.
  - For the 60% fatigue loading, the 0wt% was the only sample that displayed debonding and fibre breakage. The 1wt%, 3wt% and 5wt% all fractured in the similar manner here, that being, delamination. Once the fibre sheet layers separated, it was the fracture of the individual fibre strands rather than the failure of the entire bundle or the fibre sheet itself that ultimately brought about the failure of the entire specimen.

In conclusion, the addition of nanoclays into the matrix, strengthened the matrix and allowed the sample to have a higher fatigue cycle life. The damage mechanism was mainly characterized by delamination.
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