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1.1 Introduction

Hybrid composite is fabricated by adding two or more fibers into a single polymer system [1]. The resulting material has a unique feature that combines the advantages of each fiber. Since different fibers are added together, the benefits of one particular type of fiber property could be compensated with the other fiber lacking a specific property. The performance of hybrid composites could be influenced by many factors [2–7]:

- i. Fiber length
- ii. Fiber loading
- iii. Fiber orientation
- iv. Fiber layer sequence
- v. Fiber/matrix interfacial bonding
- vi. Failure strain of fiber

The hybrid effect is termed as an apparent synergistic improvement of properties due to different fibers in a single matrix system. The selection of fibers and their properties is of main importance to achieve the enhanced properties for the hybrid composites. Besides the physical, chemical, and mechanical stabilities of fiber, the

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matrix system also defines the strength of the hybrid composites. The different types of hybrid composites are characterized as follows [8–12]:

- i. Tow by tow: the fibers are mixed up randomly or regularly.
- ii. Sandwich hybrid composites: one material is sandwiched between two different layers.
- iii. Inter-ply or laminated: two or more fiber layers are alternatively stacked regularly.
- iv. Intimately mixed fibers: various types of fibers are mixed up randomly.

Though the hybrid composites have many advantages, the prime challenges are replacing the synthetic fiber-reinforced composites using biocomposites. Biocomposites exhibit functional and structural stability during storage and degrade upon disposal into the environment. "Engineered natural fiber" is one of the exciting concepts to obtain the enhanced strength in the biocomposites, which involves the blending of the leaf and stem fibers. The correct blending of these two fibers exhibits optimum balance in mechanical properties, resulting in balanced stiffness-toughness properties [13–15].

The mechanical and physical characteristics of the natural fiber are influenced by many factors: (i) maturity of the plant fiber, (ii) harvesting time and region, (iii) soil condition, (iv) rain, (v) sun, etc. Since the natural fibers are nonabrasive and hypoallergenic, they could be processed efficiently. Amongst the various properties of natural fibers, the low density and the cellular structure allow them to exhibit better thermal properties. However, the amorphous hemicellulose on the fiber surface can be a potential threat to the better interfacial bonding between the matrix and the fiber, thereby reducing the properties. Hence, the mechanical and thermal properties of the biocomposites could be further enhanced through chemical treatments [16]. Natural fiber has cellulose, hemicellulose, and lignin susceptible to degradation on exposure to elevated temperature [17–19]. Thus, many studies exploring the thermal properties of the biocomposites have been published over the years [20–22]. By botanical type, the natural fibers are classified into six major types (Table 1.1).

Seed	Bast	Leaf	Core	Grass and seed	Others
Kapok	Jute	Banana	Flax	Canary	Roots
Coir	Ramie	Pineapple	Kenaf	Barley	wood
Cotton	Flax	Curaua	Hemp	Wheat	
Oil palm	Hemp	Sisal	Jute	Grass	
Rice	Kenaf	Abaca		Corn	

 Table 1.1
 Classification of the natural fibers.

1.2 Thermal Characterization

The thermal analyses encompass a family of techniques that would share a common feature, whereby any material's response could be measured through heating or cooling. Thus, a significant connection is held between the temperature and the physical property of the materials. The most common thermal techniques that have been used by researchers and by industrial organizations for thermal characterization are thermomechanical analysis (TMA), thermogravimetric analysis (TGA), differential scanning calorimetry (DSC), and dynamic mechanical analysis (DMA). These techniques are not only used for measuring the physical properties with respect to the temperature changes but also used in the following areas: (i) to substantiate mechanical properties and thermal history of the biocomposites, (ii) to estimate the service life of composites in different environments, and (iii) as one of the quality control approaches in polymers and their manufacturing industries. Figure 1.1 shows some of the essential thermal analysis techniques and the characteristics measured [23-25]. In terms of research, thermal behavior of the biocomposites has been investigated by varying fiber volume fractions [26-28], varying fiber layering patterns [29, 30], using different types of chemical treatments [31, 32], adding different kinds of fillers [19, 33, 34], and using polymer blends [35, 36].

For instance, Table 1.2 presents some of the experimental works carried out on thermal properties using different natural fibers.

1.2.1 DMA

Figure 1.2 presents the step-by-step process involved in the DMA of the polymers and polymer-based composites. Output parameters such as storage modulus (E' or G'), loss modulus (E'' or G''), and damping factor (tan δ) obtained as the function of temperature are shown in Figure 1.3a. As the polymer or composite is heated in the temperature range with the simultaneous application of oscillatory load, it undergoes displacement or strain where some energy gets stored in the material,



Figure 1.1 Various thermal analysis techniques and their applications. DFA, dielectric analysis.

Hybrid composites	Details of study	References				
Thermoset polymers						
Flax/sugar palm/epoxy	DMA	[6]				
Flax/woven aloe vera/epoxy	TGA, DMA	[20]				
Sisal/cattail/polyester	Thermal conductivity	[37]				
Date palm/coir fiber/epoxy	TGA	[38]				
Sisal/jute/sorghum/polyester	TGA	[39]				
Coir/Luffa cylindrica/epoxy	DMA	[40]				
Bamboo/kenaf/epoxy	TGA, DMA, DSC	[41]				
Ramie/sisal/epoxy Sisal/curaua/epoxy	TGA, DSC	[42]				
Flax/aloe vera/hemp/epoxy	TGA, DMA	[43]				
Kenaf/pineapple leaf fiber/phenolic	TGA	[44]				
Thermoplastic polymers						
Sugar palm/roselle/polyurethane	TGA	[45]				
Jute/bamboo/polyethylene	DSC, TGA	[46]				
Sugar palm/roselle/polyurethane	TGA	[47]				
Seaweed/sugar palm fiber/thermoplastic sugar palm starch agar	TGA	[48]				
Coir/pineapple leaf fiber/polylactic acid (PLA)	TGA	[49]				
Coir/pineapple leaf fiber/ PLA	TGA, TMA	[50]				
Biodegradable polymers						
Sisal/hemp/bioepoxy	DMA, TGA	[29]				
Kenaf/sisal/bioepoxy	TGA, DSC, DMA	[51]				
Sisal/hemp/bioepoxy	TGA	[52]				

 Table 1.2
 Reported thermal based works of natural fiber-reinforced hybrid composites.

while some energy is dissipated as heat due to the internal friction. The resultant strain measured by applying the oscillatory load is represented as loss modulus, storage modulus, and phase angle or damping factor. The ability of the tested material to store the energy is termed as the storage modulus while the tendency of the material to dissipate heat energy is termed as the loss modulus. Storage modulus represents the stiffness of a polymer or composite and is often related to Young's modulus. Loss modulus is related to the molecular chain motions such as transition and relaxation within the polymer during the heating process and applied load. Tan δ is a dimensionless number obtained through the ratio of loss modulus to the storage modulus. Lower tan δ indicates higher stiffness and better interfacial bonding between fiber and polymer matrix, which restricts the molecular mobility within the polymeric chains.



Figure 1.2 Thermal characterizations of the polymer and polymer-based composite through DMA, step-by-step process.

Polymers are viscoelastic and can be classified into crystalline, amorphous, and semicrystalline (has both crystalline and amorphous characteristics) depending upon the composition. It is because of this characteristic that polymers or polymer-based composite undergoes phase change during the simultaneous application of the load and heating process (Figure 1.3b). Figure 1.3a shows the typical data obtained from DMA. Glass transition temperature (T_g) is the tangent obtained in the phase change region between glassy state and rubbery state. T_g can be below the melting temperature for a polymer, which has both crystalline and amorphous characteristics. The material tends to get softer rather than melting at T_g . DMA is particularly useful in identifying the cross-linking density of the polymer, as shown in Figure 1.3b. It can be noticed that polymers with a high cross-linking density have higher T_g and greater loss modulus and storage modulus, while it is vice versa for polymers with low cross-linking density [53].

1.2.2 TMA

TMA is a common technique used for investigating the dimensional change of material under the combination of temperature and a fixed load. Figure 1.4 presents the step-by-step process involved in the TMA of the polymers and polymer-based composites. Dimensional change of material (at nanoscale) under the influence of temperature and load can be measured in various testing modes shown in Figure 1.5. Changes in the free volume of material depending upon the heat absorption or heat release with respect to the temperature can also be determined with this technique.

Figure 1.6a–c shows that the T_g measurement for a polymer or a polymer composite can be derived from the TMA, DSC, and DMA.



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Figure 1.3 Thermal characterization of polymer and polymer-based composite with DMA. (a) Typical curve. (b) Viscoelastic characteristics of the polymer. Source: Saba et al. [53].

1.2.3 DSC

Figure 1.7 presents the step-by-step process involved in the DSC of the polymers and polymer-based composites. In DSC, the sample is heated around 30 to the elevated temperature beyond 300 °C with the constant supply of liquid nitrogen in a controlled chamber. Heat flow from the sample is measured as a function of the temperature shown in Figure 1.8. The changes in crystalline properties (T_g), melting temperature (T_m), and cold crystallization temperature (T_c) due to the introduction of two or more natural fibers in the hybrid composite can be evaluated.

1.2.4 TGA

Figure 1.9 presents the step-by-step process involved in the TGA of the polymers and polymer-based composites. It is an effective technique for evaluating thermal







Figure 1.5 Test modes in TMA. Source: Saba and Jawaid [54].

decomposition characteristics of the polymers and polymer composite reinforced with the natural fibers or the synthetic fibers. It provides the quantitative mass change of the sample due to the heating under the controlled atmosphere. A natural fiber obtained from the plants and trees is made up of the constituents such as cellulose, hemicellulose, lignin, pectin, wax, moisture, and ash. The percentage of constituents can vary from one fiber to another, which has a significant influence on the thermal decomposition characteristics of natural fiber and their composites. Also, these fiber constituents are volatile and can decompose at elevated temperatures.



Figure 1.6 T_{g} measured from the various thermal characterization techniques: (a) TMA, (b) DSC, and (c) DMA. Source: Saba and Jawaid [54].





A few milligram of sample is placed in the TGA chamber and heated from room temperature to as high as 700 °C at a defined ramp rate in the presence of nitrogen to prevent oxidation inside the chamber. The thermal stability of a polymer-based composite is usually assessed from the thermogram (TG curve) and the derivative thermogram (DTG curve) obtained from the TGA, as shown in Figure 1.10. Parameters



Figure 1.8 Thermogram from the DSC. Source: Chandrasekar et al. [55].



Figure 1.9 Thermal characterizations of polymer and polymer-based composites through TGA, step-by-step process.

such as the onset, endset, inflection temperature, and residue percentage at the end of the heating process in the TGA chamber are usually compared to identify changes due to the reinforcement percentage and type of fiber. Degradation temperature at 5%, 10%, 20%, 40%, and 80% weight loss along with the residue can also be discussed.

In the case of a polymer, thermal decomposition usually occurs in single stage, whereas, for the natural fiber, thermal decomposition occurs in two or three stages



Figure 1.10 A typical TGA curve of the hybrid composite with kenaf and bamboo fibers. (a) Thermogram. (b) Derivative thermogram. Source: Chee et al. [41].

depending on the fiber constituents. Initial mass loss between 50 and 150 °C is due to the evaporation of moisture in the fiber. The weight loss at a temperature range between 150 and 300 °C is associated with the decomposition of hemicellulose and lignin. The final weight loss between 300 and 700 °C is attributed to the decomposition of cellulose. Since the fiber constituents vary from one fiber to another, TGA has proved to be an excellent tool for determining the changes in thermal decomposition characteristics of the hybrid polymer composite reinforced with two or more natural fibers. Thermal stability is also evaluated by residue percentage at the end of the heating process. The higher the residues, the better the thermal stability of the composite.

1.3 Conclusion

Thermal characterization of the hybrid composites using various commercially available techniques such as DMA, TMA, DSC, and TGA has been discussed. The following are the conclusions:

- DMA is useful in determining the creep properties and interfacial interactions of the composites and measuring their stiffness, material behavior with respect to the phase transitions, damping, and relaxation processes in a range of frequencies and temperatures.
- TMA helps in defining the material structure with respect to the dimensional and volumetric change, surface roughness, molecular structure, cure, and cross-linking polymerization under both static and dynamic loads.
- DSC is considered as one of the primary tools for thermodynamic analysis and cure kinetics. It gives useful information on the phase transitions upon heating and quantifies the glass transition temperature, melting temperature, and crystal-lization temperature related to the polymers and polymer-based biocomposites.

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• TGA has been widely used to illustrate the thermal stability of the composites, which provides the quantitative mass change of the sample due to heating. It also provides vital information on the decomposition characteristics of the constituents of the composites at elevated temperatures.

The forthcoming chapters of this book would give extensive information on the above-discussed thermal characterization techniques with respect to different natural fibers and polymers targeted for various applications.

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