Effect of Coir Fiber and Inorganic Filler Hybridization on Innegra Fiber Reinforced Epoxy Polymer Composites: Physical and Mechanical Properties

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Abstract

The present investigation is concentrated on the results of different fillers over the physical, mechanical, and thermal characteristics of the epoxy polymer. These epoxy hybrid composites were produced using mechanical-stirring assisted wet layup method with coir micro-particles, fly ash, Titanium Carbide (TiC) nanoparticles, and Innegra fabrics contains simultaneous considering mechanical stirring with a stirring rod. The tensile, flexural, and inter-laminar shear characteristics of fabricated epoxy hybrid composites were determined using a universal testing machine. After the reinforcement of fly ash and TiC nanoparticles offers the most remarkable improvement in tensile, flexural, and impact strength, about 2.84, 1.65, and 9.19 times compared with pure epoxy polymer. The differential scanning calorimeter (DSC) and thermogravimetric analysis (TGA) showed the epoxy hybrid composites' enhanced thermal stability. The homogeneity of fillers dispersion in the epoxy polymer was observed from a scanning electron microscope (SEM). Keywords: Contact angle; Fly ash; Inter-laminar shear strength; Innegra Fabric; Tensile Strength; TiC nanoparticle

1. Introduction

The epoxy-based resins are broadly utilized in polymer laminates, automobiles, aerospace, adhesives, paints fabrication, surface coating methodology, and other engineering materials (May and Tanaka, 1973; Rs, 1979). The performance of epoxy composites in sectors is because of their higher corrosion and chemical resistance, greater mechanical and thermal characteristics, lower shrinkage (Adams and Gannon, 1986). Epoxy composites can study in specific environmental situations. The cured epoxy polymer exhibited a 3D (three-dimensional) extremely cross-linked chain scheme. The characteristics of epoxy polymers are strained in different applications because of a higher range of cross-linking density which is influenced by intrinsic brittle behavior of components and lower toughness. Various investigations have been performed to enhance the toughness and other characteristics like thermal stability, dynamic mechanical and crack resistance of epoxy polymer by adding different kinds and quantities of nanoparticles in a polymer (Duraibabu et al., 2014). The breaking and uniform dispersion of nanoclusters inside the polymer are two primary issues for nanolaminates fabrication (Sun et al., 2015). Dispersion and cluster size of nanoparticles in polymer resin shows the confident development in pure polymer's fundamental characteristics with stronger interfacial bonding within the molecular networks and fillers (Radoman et al., 2014; Bal, 2010). The particle accumulation trend because of non-uniform dispersion and van der Waals forces in epoxy polymer has strengthened investigators for deciding the excellent processing methods to lower or decrease the accumulated filler size and provide its homogeneous distribution. Many ways are already in the tendency for polymer nanolaminates fabrication from shear comprising using manual stirring, melt mixing, solution mixing (Becker et al., 1996; Rong et al., 2001), direct addition with chemical route techniques (Kumar et al., 2017; Kang et al., 2001), acoustic cavitation (Eskin, 2001), ultrasound vibrations (Wang et al., 2001), ultrasonic waves (Kumar et al., 2016; Xu et al., 2004), and ultrasonic irradiation (Xia and Wang, 2003). The acoustic cavitation
method is an efficient process for distributing nanoparticles in polymer resins (Zunjarrao and Singh, 2016; Halder et al., 2012; Tsekmes et al., 2015).

Tsekmes et al. (2015) have declared that epoxy-based laminates benefit from greater breakdown strength with enhanced thermal stability. Generally, utilization of 60–70 wt.% of silica microparticles in the epoxy laminate insulators can be observed in the industrial sector. Imai et al. (2006) have signified that the silica particle leads to show the lower coefficient of thermal expansion. Ramanujam et al. (2019) have investigated the corona treatment inaugurated because of water contents on IXEPLAS reinforced epoxy nanolaminates. IXEPLAS is a hydrotalcite component customized with zirconium phosphate. Inorganic ion exchangers such as zirconium phosphate lead to better resistance from oxidation and more significant ion entrapping capability. Thus, it can be suitable in the electrical schemes for enhancing their performance.

One of the considerable interests in epoxy laminate samples is water absorption in humid environmental situations. Nogueira et al. (2001) have signified that moisture in epoxy laminates is present in two dissimilar systems, such as bound water in interaction with hydrogen bonding of the laminate and moisture present in the accessible quantity micro-gaps in the resin. In composites' condition, this principle is profound compared to neat epoxy due to the interface between filler and matrix will act as a possible region for the moisture molecule to cooperate (Zou et al., 2008). This principle may potentially tend to considerable irreversible material modifications like final degradation, plasticization, and swelling of insulating material (Wang et al., 2006). Exposing polymeric samples to hot water is one of the usual techniques to affect the long-term process or accelerated aging of moisture uptake principle (Pitarresi et al., 2018). Ramanujam et al. (2019) have performed moisture absorption investigations with pure epoxy and their respective nanolaminates. They have signified that the percentage of the weight of the pristine epoxy polymer is around 0.5%. The epoxy polymer laminate sample applied in the current investigation has a gain in weight % of about 0.2. The pure epoxy polymer has a freer amount applicable in the bulk of the sample for the moisture to penetrate. The fillers reinforcement played as protection for water diffusion in the bulk amount of laminate sample, decreasing the moisture absorption compared to neat epoxy. When the moisture absorption of pure epoxy is higher than two times of moisture absorption in a laminate sample, the current investigation is concentrated more on an epoxy hybrid laminate sample to interpret the influence of water absorption and mechanical characteristics.

Nonetheless, it also has many limitations like thermal stability, softness, and flexural modulus (Wu et al., 2019). Incorporating inorganic particles like carbon-based fillers, calcium carbonate, silica, and calcium sulphate is an efficient technique for enhancing the overall characteristics of polymers (Abdel-Gawad et al., 2018; Yuan et al., 2016; Cui et al., 2017; Khaleghi et al., 2017). Among these fillers, fly ash is a reinforcing agent that can adequately enhance the thermal resistance, impact, and modulus of polymers, owing to their other advantageous properties like chemical corrosion resistance, lower cost, higher mechanical properties, availability, higher mechanical strength, and more necessarily usage of industrialization (Dahalan et al., 2018; Liu and Zhong, 2014). Fly ash is the primary solid waste released from pulverized coal's combusting process in thermal power plants. It is now the most excellent
individual resource of solid waste management (Ding et al., 2017; Temuujin et al., 2019; Wang et al., 2014). The constituent, surface chemical reaction, and mineralogy of fly ash have essential influences on its utilization (Yao et al., 2015; Hower et al., 2017) like wastewater treatment in the industry, catalyst carriers for the modification of poisonous gases, raw materials for plastics, adsorbent, silica aerogels, soil conditioners in agriculture, and construction materials as admixture (Asl et al., 2018; Asl et al., 2019; Cai et al., 2020). When fly ash is made of adaptable active oxides, it can also be applied as a filler for polymer laminates. It can enhance polymer laminates' thermal and mechanical characteristics, whereas fabrication cost is furthermore decreased.

Furthermore, only a restricted quantity of hydroxyl functional group on the surface of fly ash may affect the poor compatibility between polymer and fly ash. Hence, avoid the movement of mechanical stress used to the laminate and subsequently control the laminates' damage (Civancik-Uslu et al., 2018). To enhance the bonding between organic matrix and fly ash, surface treatment of fly ash is essential. Patil et al. (2016) fabricated an optimized planetary ball milling technique from the Taguchi experimental design to manufacturing nano-sized fly ash. Van der Merweet et al. (2014) observed that the treatment of fly ash with sodium lauryl sulphate tends to no considerable modifications in the chemical constituents, and modified fly ash could be applied as reinforcement in polymers under lower content. Furthermore, previous investigations on the activation of fly ash by mechanical alloying have focussed mainly on the fabrication of nano-sized fly ash (Thongsang et al., 2012; Yuan et al., 2019), which generally lacks a long mechanical alloying period higher consumption of energy (Sundum et al., 2018; Li et al., 2017). An easy and efficient treatment technique would share fly ash utilization in the production of advanced industrial components.

Among the different ceramic nanofillers, the titanium carbide (TiC) nanoparticles are known for their higher thermal stability, larger surface area, superior electrical insulation characteristics, and excellent mechanical characteristics that produce it more prominent an inferior aspirant filler (Mallakpour and Khadem, 2015; Omrani et al., 2009). In the present study, the fly ash and TiC are selected as potential reinforcement. The fly ash and TiC are well-known for their higher mechanical characteristics and thermal stability. From the literature study, it has been observed that limited investigations have been conducted to manufacture hybrid composite with coir and Innegra fiber incorporated in the epoxy resin with fly ash and TiC. In this study, an experiment has been appointed to fabricate composites combined with Innegra fiber, fly ash, TiC filler comprised with coir fiber in the epoxy polymer. The effect of fly ash and TiC particles incorporation in Coir fiber-Innegra fiber-epoxy composite on morphological, mechanical, and thermal characteristics were measured from tensile, impact, inter-laminar shear strength, flexural, hardness, scanning electron microscope (SEM), differential scanning calorimeter (DSC), and thermogravimetric analysis (TGA) is considered.

2. Materials And Methods

2.1. Materials
In the present investigation, synthetic epoxy Epotec YD-535 LV and hardener of Epotec TH-725 were used as polymer matrix and procured from Aditya Birla Chemicals Ltd., Thailand. The coconut fibers purchased from Tongmongkol Coconut fiber, Prachuapkhirikhan, Thailand was applied as reinforcement material for the manufacturing of synthetic epoxy laminate. Two types of inorganic fillers were utilized in this investigation as TiC (10 nm of particle size) was supplied from Zhuzhou Sanyinghe International Trade Co. Ltd., China, and fly ash was collected from industrial chimney, Thermal Power Plant, Chennai applied as another filler reinforcement to the matrix. A novel Innegra fiber has been purchased from Innegra Technologies, South Carolina, United States in the form of woven mat and also applied as a reinforcement material to the synthetic epoxy polymer.

2.2. Fabrication of synthetic epoxy hybrid composite

The purchased coconut fibers were cleared up with fresh water to eliminate the dirt particles and dried in sunlight for three days to remove the moisture content. The cleaned coconut fibers were sectioned into small pieces and transferred to the Universal milling machine to produce the micro-fillers from coconut fibers subjected to sieving and separated from a 250 µm filter. The synthetic hardener, epoxy polymer, and reinforcement fillers were weighed accurately in different ratios with the help of weighing balance. The details for the fabrication of the epoxy hybrid composite are shown in Table 1. The mechanical stirring was applied to achieve a homogeneous distribution of fillers in the epoxy polymer. The combined mixture was rapidly transferred to the die mold to avoid premature curing and discharge for a slower curing process. The fabricated sample was removed from the die mold and shifted to the furnace for post-curing at 80 ± 2°C to decrease the sample's residual moisture.
Table 1
Detailed information about the fabrication of coir fiber/ Innegra fiber/ TiC/ fly ash synthetic epoxy hybrid composites

| Hybrid Composite type | Description                                           | Coir micro-particles | Innegra fabric | TiC nanoparticles | Fly Ash |
|-----------------------|-------------------------------------------------------|----------------------|----------------|------------------|---------|
| PE                    | Pure synthetic epoxy (Epotec YD-535 LV and hardener Epotec TH-725) | -                    | -              | -                | -       |
| CIPE                  | Epotec YD-535 LV reinforced with coir micro-particles, and Innegra fabric | 10 wt.%              | 3 layers       | -                | -       |
|                       |                                                       |                      | (116 g/m²)     |                  |         |
| CFIPE                 | Epotec YD-535 LV reinforced with coir micro-particles, fly ash, and Innegra fabric | 5 wt.%               | 3 layers       | -                | 5 wt.%  |
|                       |                                                       |                      | (116 g/m²)     |                  |         |
| CTIPE                 | Epotec YD-535 LV reinforced with coir micro-particles, TiC nanoparticles, and Innegra fabric | 5 wt.%               | 3 layers       | 5 wt.%           | -       |
|                       |                                                       |                      | (116 g/m²)     |                  |         |
| CFTIPE                | Epotec YD-535 LV reinforced with coir micro-particles, fly ash, TiC nanoparticles, and Innegra fabric | 4 wt.%               | 3 layers       | 3 wt.%           | 3 wt.%  |
|                       |                                                       |                      | (116 g/m²)     |                  |         |

2.3. Characterization of fabricated epoxy hybrid composite

2.3.1. Fourier Transform Infra-red (FTIR) spectrum

FTIR spectra analysis was conducted with the sample comprised in a KBr (Potassium Bromide) filler from Perkin Elmer Spectrum 2, FTIR spectrometer. The epoxy hybrid composite's wavenumber was ranging between 4000 and 400 cm⁻¹ with a resolution of 1 cm⁻¹ under 32 scans per minute.

2.3.2. Water absorption capacity

According to the ASTM D 570 standard, the water absorption capacity was performed for epoxy hybrid laminates. The specimen was dried in a heating furnace for a specific period at 80 ± 2°C, and the initial mass of the specimen was determined. The fabricated epoxy hybrid laminates were then immersed in fresh water at 24 ± 2°C for three months. After immersing, specimens were taken out, excess water has been cleaned with a cotton cloth, and the final weight was measured. The variation between the final and initial mass has been determined, and the amount of water absorption has been estimated as per the standard.
2.3.3. Contact angle

The contact angle test experimented in 15 LJ data physics, Optical contact angle (OCA) under 26 ± 2°C and 1 µL of deionized water were poured on the surface of epoxy hybrid laminate with the help of a syringe. The left and right contact angles were determined, and for each specimen, the mean value of five parts was recorded as an outcome.

2.3.4. Mechanical characteristics

The mechanical characteristics are essential for the utilization of composite materials. The specimens (width x length x thickness: 15 mm x 100 mm x 3 mm) for the tensile experiments were conducted as per the ASTM D 3039 standard. The specimens were fixed on the Comotech Universal Testing Machine and expanded longitudinally at 1 mm/min until they break.

Short beam shear and three-point bending experiments were performed in Comotech Universal Testing Machine of 10 kN load cell in three-point flexural loading arrangement. The crosshead speed and preload of 5 mm/min and 0.1 MPa, respectively, were employed. The inter-laminar shear strength of the laminate was investigated by controlling the thickness to span proportion 4 to 5 according to the ASTM D 2344-16 standard. The smaller span length increases the cracking and shear failure along the horizontal plane between the layers. The flexural modulus and strength of the manufactured epoxy hybrid laminates were measured according to the ASTM D 790 standard with a span to a thickness of 32. The bending specimen's dimension encompasses a width of 15 mm and a span length of 80 mm. The flexural strength at the exterior fiber at the center point and the flexural modulus was calculated as per the standard. The Izod impact test of the epoxy hybrid composites was determined from Zwick/ Roell Impact testing machine furnished with an Izod 5.5 J hammer constructed as per the ASTM D 256 standard and accommodating to the technique based on the standard (un-notched).

The Shore D hardness of the fabricated synthetic epoxy hybrid composites was determined with the indenter-based hardness measurement machine. The indenter was pressured on the sample's surface up to the bottom portion of the indenter is in full strike with the composite. The dial of the hardness indenter signifies the Shore D hardness of the epoxy hybrid composites. The experiments were carried out under ten different composite positions, and mean hardness outcomes were determined. All the experiments were performed under 60% of relative humidity and 24 ± 2°C of room temperature.

2.3.5. Fractographic evaluation

The microstructures were observed from a scanning electron microscope, FEI 450 Quanta, with 15 kV of accelerating voltage to examine how different reinforcements influence a polymer's structure.

2.3.6. Thermal characteristics

DSC (Differential scanning calorimeter) graphs from 20 to 300°C were reported from Mettler Toledo, DSC 3+/700/2167 machine at nitrogen atmosphere under the heating rate of 10°C/min. The TGA (thermogravimetric analysis) was performed on Mettler Toledo, TGA 2, TGA/DSC 3 + HT/1600 machine to
examine the epoxy hybrid's thermal stability composites. The specimens were heated between 30 and 700°C under a nitrogen atmosphere with a heating rate of 15°C/min. The mass loss was determined, and residual char content was measured from 600°C with the help of DTG (derivative thermograms).

3. Results And Discussion

3.1. Effect on FTIR spectra

Figure 1 exhibits the FTIR spectrum of pure epoxy polymer and their respective laminates based on the filler and Innegra fabric reinforcement to examine the effect of inorganic fillers and epoxy functional groups. The pure epoxy polymer shows a clear carboxyl peak at 1654 cm⁻¹ existing to C—O stretch functional groups. Other peaks at 1490 and 1596 cm⁻¹ are ascribed to a benzene ring. Whereas two constrain intensity peaks between 930 and 770 cm⁻¹ and the peak at 1226 cm⁻¹ relates to the asymmetric and symmetric stretching peaks of R-O-R, respectively (Niu et al., 2014; Niu et al., 2021). The effect of various inorganic filler constituents on the epoxy polymer's molecular chain has primarily followed the peak at 630 cm⁻¹. The vibration kinds in this region are obscured and imbricating, and the features were terrible. The FTIR spectrum signifies that epoxy polymer and laminate specimens exhibit the evident characteristics peaks of epoxy polymer. The chemical scheme of epoxy polymer is not considerably modified with the incorporation of different fillers. In the FTIR spectrum of CFIPE and CFTIPE laminates, two other characteristic peaks can be examined at 1430 and 880 cm⁻¹ that can be assigned to the asymmetric and out of plane ben vibration the CO32- functional groups in fly ash (Jena and Sahoo, 2019). Hence, it can be proved that calcium carbonate is the principal constituent of fly ash. Hydrogen bonds produce efficiently between the hydroxyl regions. When this happens, the hydroxyl stretch vibration shows red movement in the infra-red spectra. The production of hydrogen bonds between the hydroxyl functional groups affects this principle known as the effect of hydrogen bonding (Finch, 1970).

From the infra-red spectrum, the hydroxyl stretch vibrations of CIPE, CFIPE, CTIPE, and CFTIPE composites were 3430, 3431, 3440, and 3445 cm⁻¹, respectively. As shown in CIPE, CFIPE, CTIPE, and CFTIPE, the incorporation of fillers concluded in producing more hydrogen bonds between coir fiber, epoxy, fly ash, TiC, and Innegra fabric, signifying an increment in crosslinks between various constituents (Zhang et al., 2018). This outcome is the production of a close compact system; hence, these laminates' enhanced mechanical characteristics can be described (Deshmukh et al., 2021). The hydroxyl-stretch vibration of the CISE composite was lower than that of the CFIPE, CTIPE, and CFTIPE due to inorganic fillers' presence (Wu et al., 2020). Further, it can produce a new hydrogen bond with the polymer as an essential constituent of fly ash, TiC, and Innegra fabric may also offer some hydroxyl functional groups to create hydrogen bonds with the polymer. Furthermore, the crystallinity index more actively influenced the mechanical characteristics than the hydroxyl-stretch vibration, signifying that crystallinity was the primary parameter that impacted the polymer composites' mechanical aspects (Ji et al., 2021; Sun et al., 2019).
3.2. Effect on physical properties

Figure 2 (a) exhibits that after 90 days, water absorption attains 2.48% (CTIPE), 2.01 (PE), 1.83% (CFIPE), 1.44% (CIPE), and 1.42% (CFTIPE). The CTIPE composite consumed more water while comparing with other combinations due to the greater pore structure. A similar outcome has been observed by Ajdary et al. (2020), considered that greater pores tend to higher absorption water under capillary action. Furthermore, the particular kind of filler did not noticeably modify the quantity of water consumed. This can be assigned to the principle that all the specimens were porous and consume more water under the capillary action. However, polymer reinforced most hydrophilic functional groups, and CFTIPE, CIPE, and CFIPE are more hydrophobic than polymers (Deshmukh et al., 2021; Ajdary et al., 2019; Zhang et al., 2017). The particles were uniformly distributed in the laminates, improving their durability with water. The hydroxyl functional groups of CIPE and CFTIPE influenced them to combine strongly with the laminate, resulting in this marginally decreasing the quantity of water consumed from the CIPE and CFTIPE laminates.

The water contact angle is the important factor that signifies the water's wettability on the surface of the fabricated laminate. Figure 2 (b) presents the water contact angle after 5 min; the outcomes are PE (81.4°) > CFIPE (76.63°) > CFTIPE (74.73°) > CIPE (70.9°) > CTIPE (69.83°). In 5 min, the PE, CFIPE, CFTIPE, and CIPE was nearing the hydrophobic condition. The CTIPE laminate was previously hydrophilic. The fillers enhanced the water-resistance and can almost attain the influence of waterproofing TiC and fly ash (Sun et al., 2019). The surface energy of the laminates improved with the incorporation of fillers; furthermore, the appearance of certain hydrophilic inorganic components in epoxy polymer reduced the surface energy (Wu et al., 2020). In this context, the dense architecture affected by the increment of crosslinking and the capillary action of various diameters of pores also outcome in distinct water contact angles (Deshmukh et al., 2021; Ajdary et al., 2020). Hence, the water contact angle improved to fluctuating degrees upon the reinforcement of coir fiber, TiC filler, and fly ash.

3.3. Effect on mechanical properties

3.3.1. Tensile properties

Figure 3 (a) presents the stress-strain graphs of the epoxy hybrid composites reinforced with different fillers. The influence of other filler addition on the tensile strength and modulus of epoxy hybrid composites were measured and presented in Fig. 3 (b). Figure 3 (b) exhibits that the tensile modulus and strength of CIPE, CFIPE, CTFIPE, and CFTIPE composite improved by 2.89, 1.71, 2.52, and 2.84 times respectively. The strength of the epoxy hybrid composites based on the dispersion of fillers in the polymer and bonding between the matrix and particles. The uniform distribution of particles in complete polymer acted as a specific character to enhance the tensile characteristics of laminate. The testing process efficiently damages the cluster size of filler and disperses the filler uniformly incomplete polymer. CFIPE laminate presented lower tensile attributes because of the improvement in cluster size and density of the filler. The tensile characteristics of CFIPE and CFTIPE composite showed reduced when compared with
The reduction in tensile characteristics under this reinforcement can be explained by an incrementing susceptibility of accumulation, which tends to lower energy dissipation in the scheme under the deformation of viscoelasticity. It was observed that with the increment in weight and accumulation size, the movement of molecule and chains become simple, which leads to a reduction in tensile characteristics. The greater cluster size of filler also decreases the bonding of filler-polymer and offers a more straightforward method to increase the initial damage during the tensile load.

Other investigators have observed similar results that the improvement in strength is assigned to the principle that the Zirconia filler has much better stability than the epoxy polymer and also to the stronger bonding between the polymer and filler that allows the suitable load dispersion in hybrid laminate regions (Ma et al., 2018; Kumar et al., 2021). The decrement in strength is ascribed to the appearance of the higher amount of clusters. Furthermore, homogeneous cluster-free distribution and strong bond interaction of optimum fillers reinforcement with polymer permits part of the load transfer, leading to the epoxy composites' considerable tensile characteristics (Haldar et al., 2012).

### 3.3.2. Tensile fracture

The SEM micrograph of the tensile experimented samples observed in Fig. 4. The epoxy polymer sample's fracture microstructure examines the crack and brittle behavior, as observed from previous published literature (Mohit et al., 2021). Figure 4 (a) shows the CIPE sample affirms the portion of inadequate filler dispersed from the epoxy polymer and the matrix fragmenting. The extensive matrix damage and failure of a filler-matrix interface are also apparent. SEM micrographs of CFIPE signifies the portion of Innegra fiber deceived and constrained micro-cracks (Fig. 4b). The addition of fly ash decreases the bulging of Innegra fiber bundles and attempts of damage at an initial point of reinforcing by prohibiting the crack generation. The SEM micrograph of CTIPE shows the isolated epoxy/ titanium carbide/ coir fiber cluster and the Innegra fibers' forecast to the outer part. The debonding of the Innegra fiber-epoxy/titanium carbide/ coir fiber is found in certain portions because of the failure of interfacial bonding (Fig. 4c).

Furthermore, compared with the neat epoxy and CIPE, in this condition, almost all the Innegra fibers imply to be coated with the epoxy/ titanium carbide/ coir fiber even after the disintegration. This signifies a better bonding between the Innegra fabric and epoxy/ titanium carbide/ coir micro-particles. SEM micrographs of CFTIPE composite observed at the fractured part represents disintegration, whereas the Innegra fabric-fly ash-TiC-epoxy bonding does not isolate in certain parts even after the damage, which signifies the higher adhesive load between them. Filler accumulation, Innegra fabric damage, debonding, and inadequate filler-polymer adhesion portion are also observed from SEM micrograph of CFTIPE tensile experimented sample (Fig. 4d) (Rao et al., 2021).

### 3.3.3. Flexural properties

The flexural stress-strain graphs are shown in Fig. 3 (c). The flexural experiment's outcome in Fig. 3 (d) shows that the incorporation of TiC or fly ash particles affected an increment in the flexural modulus and strength. The epoxy polymer resin reinforced with coir micro-particle, fly ash, and Innegra fiber shows the
highest flexural strength showing 127.22% enhancement over the neat polymer. The fly ash filler possesses a greater specific surface area compared with the TiC nanoparticles. Thus, when comprised with the polymeric networks, the fly ash particles can produce stronger bonding, a flexural load can be shifted through these inorganic filler networks to other enslave. Even though one network damages under external force, others can still hold the laminate structures, and the whole sample is not intimidated. It can also be concluded from the outcome that TiC incorporation is considered in enhancing the modulus potentially because of around uniform fillers in the epoxy polymer that rigidly support the nearby Innegra fabric layers. The versatility of the inorganic filler in improving the flexural strength reduces beyond the crucial addition of fly ash because of the accumulation of inorganic fillers (Shivamurthy et al., 2013; Taurozzi et al., 2012). This filler accumulation is affected by an increment of filler-filler interactions during the mechanical stirring process (Sudheer et al., 2013). The fillers accumulation would potentially increment polymer-filler interruption viscosity and affect bad wetting reaction between them (Sudheer et al., 2013). Hence, the lower enhancement in the flexural characteristics was endorsed for CFTIPE that signifies 65% improvement compared to a neat epoxy polymer. The CFIPE has a greater flexural modulus when compared with the CTIPE composite (Zhang et al., 2017; Chaurasia et al., 2019). Hence, the highest flexural modulus was endorsed for CTIPE composite that signifies 102.88% improvement compared to a pure epoxy polymer. The flexural modulus also enhanced TiC nanoparticles’ report and the epoxy polymer physical networking (Chaurasia et al., 2019). Furthermore, CFTIPE and CIPE exhibited lower flexural modulus compared with CFIPE and CTIPE since the fillers accumulation may have considered, which reduced the reinforcement efficacy and decreased the efficient contact area within the polymer and filler particles (Rao et al., 2021).

3.3.4. Interlaminar shear strength

The general shear stress vs. strain variation is presented in Fig. 3 (e) is remarkably distinct for every laminate. From the curve, it can be concluded that the CIPE composite can withstand a force 1.98 times higher than the pure epoxy polymer, whereas CFIPE bears an influence of 28% larger than the pure polymer. Also, CFIPE shows a greater shear strength than pure epoxy polymer, as observed from Fig. 3 (e). Five specimens were experimented with in every combination for the calculation of shear strength value. The mean and standard errors of every combination have also been shown in Fig. 3 (f). The higher value of inter-laminar shear strength for CIPE composites is obtained due to the addition of Innegra fabric which improves the interlayer roughness of the surface compared with CFIPE, CTIPE, and CFTIPE composites. As the expansion of inorganic filler expands the crucial value, the epoxy resin network’s movement may get harmed. This can influence bad-wetting with the fiber and potentially reduce the shear strength value (Rao et al., 2021). The resin networks around the surface of the particle are entirely dissimilar from the extent polymer.

3.3.5. Impact strength and Hardness

The impact strength of the epoxy polymer composites can be examined from Fig. 5. Compared with CFIPE or CTIPE composite, the impact strength of the CFTIPE composite was reduced. It has been already discussed in different kinds of literature that incorporating hybrid inorganic fillers in the polymer resin
decrements their impact strength (Zhou et al., 2019; Zhang and Qi, 2014). A higher concentration of incorporating fillers was more significant than the decrement in the impact strength. Furthermore, reinforcing TiC or fly ash in the coir-Innegra fiber-epoxy composite enhanced the impact strength considerably for CFIPE and CTIPE specimens compared to CFTIPE by 7.52% and 18.4%, respectively. This probable increment in the impact strength was because of the epoxy matrix's toughening with fly ash and TiC particles. Also, CTIPE exhibited a higher impact strength of 46.59 kJ/m², slightly higher than CFIPE. Hence, the positive influence of TiC nanoparticles addition on impact strength is apparent. The hybridization exhibited a possible effect on impact strength assignable to the enhanced interfacial adhesion and good toughening features transmitted by the appearance of TiC nanoparticles, coir micro-particles, and Innegra fabric the epoxy matrix. Similarly, Khandelwal and Rhee (2020) also examined and expressed that inorganic filler influence on basalt fabric polymer laminate interface. Within the hybrid laminates, the effect of reducing filler concentration was more uttered than the incrementing filler to filler interactions (Satapathy and Kothapali, 2018).

The shore-D hardness of coir-Innegra fiber epoxy hybrid composites as presented in Fig. 5. The outcome shows that the reinforcement of fillers enhances the fabricated CIPE, CFIPE, CTIPE, and CFTIPE composites’ hardness. As a result of the addition of retrieved filler, the enhancement of hardness was reported as high for CFTIPE laminate than the CIPE, CFIPE, and CTIPE laminate. Furthermore, a considerable enhancement in hardness was examined for CFTIPE laminates. Remarkably, the enhancement was up to 4% for the CFTIPE composite when compared with other combinations. The enhancement in the hardness of a laminate is based on the fly ash and TiC nanoparticles (Agarwal et al., 2014). The inter-particle void of the fillers in the epoxy polymer is considerably reduced with fly ash and TiC nanoparticles. This outcome in a higher quantity of load-bearing regions being obtained in the laminates. This conclusion was also reported in the glass fabric epoxy laminates reinforced with fired alumina (Sabarinathan et al., 2020).

3.4. Thermal stability

The synergy between fillers, Innegra fabric, and epoxy resin is investigated from DSC examination. To examine the glass transition temperature, DSC investigations are conducted with an optimum heating rate of 10°C/min. In Fig. 6 (a), heat flow with incrementing temperature in epoxy hybrid laminates is exhibited. The glass transition temperature of fabricated epoxy hybrid composites is estimated from the extrapolation of the endothermic peak. It is examined that the glass transition temperature measured from DSC curves is highest for CTIPE composite, and it entails the better interaction within the fillers, Innegra fabric, and epoxy matrix. The CFTIPE composite has observed a slight decrement with the addition of fly ash because of the lower stability of hydrogen bonding under elevated temperature (Lei et al., 2016; Wang et al., 2016).

The thermal stability curve of pure and different fillers and Innegra fabrics reinforced epoxy hybrid composites was examined from TGA and DTG (Fig. 6b, c). The enduring weight percentage of pure epoxy and their respective laminates as a function of temperature with the heating rate of 15°C/min has been explained for different constituents' thermal stability examination. The epoxy hybrid composites that
withstand under a higher temperature range at a given weight percentage are more thermally stable when compared with other combinations. The thermal characteristics of pure epoxy and their respective hybrid laminates were determined between the ranges of temperature 160–600°C. This temperature range was considered due to the deviation of weight exhibited additional influence because of appreciable oxidation and water absorption after 600°C and before 160°C (Ogasawara et al., 2011). CFIPE epoxy hybrid laminate exhibited the highest thermal stability (onset and end-set temperature of 344.66 and 416.72°C) compared with other laminates (Table 2). The initial weight loss of epoxy hybrid composites is critical to interpreting their thermal stability.

Furthermore, the epoxy resins have a comparably higher cross-linking focus which establishes comparably higher degradation temperature (Niu et al., 2014; Chatterjee and Islam, 2008). The improvement in thermal stability of CFIPE composite was because of the strong interaction of polymeric network, fillers, and Innegra fabrics. The reduction in thermal stability with TiC fillers in the polymer maybe because of the higher cluster size deviation of fillers in an epoxy polymer. The greater cluster size with non-homogeneous dispersion in the epoxy polymer may negatively impact the interfacial-bonding of filler-polymer and decreases the thermal stability of laminates. The carbon residue yield of epoxy hybrid composite is considerably enhanced in the CFIPE composite (Table 2). The endorsed carbon residue could be assigned to the further cross-linking within the polymer, Innegra fabric, and fillers (Ma et al., 2018).

| Sample designation | Glass transition temperature in °C | Onset temperature in °C | Endset temperature in °C | Carbon residue in % |
|-------------------|-----------------------------------|-------------------------|--------------------------|---------------------|
| PE                | 77.21                             | 334.04                  | 403.67                   | 7.32                |
| CIPE              | 171.17                            | 336.46                  | 405.85                   | 10.93%              |
| CFIPE             | 170.5                             | 344.66                  | 416.72                   | 10.16%              |
| CTIPE             | 177.33                            | 338.84                  | 410.83                   | 8.63%               |
| CFTIPE            | 176.17                            | 344.26                  | 412.23                   | 8.79%               |

4. Conclusion

The developed epoxy hybrid composites were fabricated from the mechanical-stirring assisted wet layup method. Tensile, flexural, inter-laminar shear, impact strength, resistance from water absorption, glass transition temperature, and the epoxy hybrid composites’ thermal stability were improved compared with neat epoxy polymer. The impact strength has the lowest and highest impact strength of 46.59 kJ/m2 and 32.03 kJ/m2 by CTIPE and CIPE composites. This signified a possible hybridization influence transmitted from the TiC nanoparticles on the epoxy polymer laminate. The previous characteristics of the hybrid...
laminates were examined to enhance the addition of fly ash and TiC fillers. Apart from all the laminate fabrication, hybrid laminate with TiC nanoparticles showed an enhancement of 10.88 times in the impact strength compared with epoxy polymer. The chosen optimum resin blend of epoxy, fly ash, TiC nanofillers, and Innegra fabrics as the polymer matrix indeed promoted this enhancement. The stronger interfacial adhesion of fly ash, TiC nanofiller, and Innegra fabrics with the epoxy matrix, as observed from the FTIR examination, permitted an efficient load transfer using the interphases. Even though there was a slight decrement in the tensile strength of the CFIPE composites compared to laminate possessing coir micro-particles, fly ash, and Innegra fabrics alone, the flexural and tensile characteristics were in the proximity of the epoxy polymer. Apart from all the fabricated laminates, the lowest and highest strength of 66.09 MPa and 111.91 MPa were reported from CFIPE and CIPE composite.

Additionally, a similar trend has been examined for tensile modulus, where the remarkable and lower tensile modulus of 1044.7 MPa and 731.71 MPa for CFTIPE and CIPE composite, respectively. The flexural characteristics of the laminates also pursued the same trend as tensile. The laminates flexural modulus and strength investigate exhibit that CFIPE occurs the maximum flexural modulus and strength as 153.01 MPa and 2556.3 MPa. The distribution of coir micro-particles, fly ash, TiC nanoparticles, and Innegra fabrics in the epoxy polymer were proved from SEM micrographs. CFIPE composite showed the most remarkable improvement in onset and end-set temperature up to 344.66°C and 416.72°C, respectively. The glass transition temperature also exhibited the most significant improvement for CTISE composite around 100°C when compared with the pure epoxy polymer because of the better interactions between the fillers, Innegra fabric, and epoxy polymer. Cross-linking density and thermal stability of CFIPE, CTIPE, and CFTIPE composite also exhibited considerable improvement compared with pure epoxy polymer.

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**Statement of ethical approval:** Not applicable

**Conflicts of interest:** None

**Informed consent:** Not applicable

**Ethical standards:** Not applicable

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Figures

Figure 1

FTIR Spectra of coir fiber/ Innegra fiber/ TiC/ fly ash synthetic epoxy hybrid composites
Figure 2

Physical properties of coir fiber/ Innegra fiber/ TiC/ fly ash synthetic epoxy hybrid composites (a) water absorption, and (b) Contact angle
Figure 3

Mechanical properties of coir fiber/ Innegra fiber/ TiC/ fly ash synthetic epoxy hybrid composites (a) Tensile stress-strain, (b) tensile properties, (c) Flexural stress-strain, (d) flexural properties, (e) ILSS stress-strain, and (f) ILSS
Figure 4

Tensile fracture of coir fiber/ Innegra fiber/ TiC/ fly ash synthetic epoxy hybrid composites (a) PE, (b) CIPE, (c) CF1PE, (d) CTIPE, and (e) CFTIPE

Figure 5

Impact strength and shore hardness of coir fiber/ Innegra fiber/ TiC/ fly ash synthetic epoxy hybrid composites
Figure 6

Thermal characteristics of coir fiber/ Innegra fiber/ TiC/ fly ash synthetic epoxy hybrid composites (a) DSC, (b) TGA, and (c) DTG curves