Pyrolysis Kinetic Behaviour of Glass Fibre-Reinforced Epoxy Resin Composites Using Linear and Nonlinear Isoconversional Methods

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Abstract: Due to the increasing demand for glass fibre-reinforced epoxy resin composites (GFRC), huge amounts of GFRC waste are produced annually in different sizes and shapes, which may affect its thermal and chemical decomposition using pyrolysis technology. In this context, this research aims to study the effect of mechanical pre-treatment on the pyrolysis behaviour of GFRC and its pyrolysis kinetic. The experiments were started with the fabrication of GFRC panels using the vacuum-assisted resin transfer method followed by crushing the prepared panels using ball milling, thus preparing the milled GFRC with uniform shape and size. The elemental, proximate, and morphology properties of the panels and milled GFRC were studied. The thermal and chemical decomposition of the milled GFRC was studied using thermogravimetric coupled with Fourier-transform infrared spectroscopy (TG-FTIR) at different heating rates. Meanwhile, the volatile products were examined using TG coupled with gas chromatography–mass spectrometry (GC-MS). The TG-FTIR and TG-GC-MS experiments were performed separately. Linear (Kissinger–Akahira–Sunose (KAS), Flynn–Wall–Ozawa (FWO), and Friedman) and nonlinear (Vyazovkin and Cai) isoconversional methods were used to determine the pyrolysis kinetic of the milled GFRC based on thermogravimetry and differential thermal gravimetry (TG/DTG). In addition, the TG/DTG data of the milled GFRC were fitting using the distributed activation energy model and the independent parallel reactions kinetic model. The TG results showed that GFRC can decompose in three stages, and the main decomposition is located in the range 256–500 °C. On the other hand, aromatic benzene and a C-H bond were the major functional groups in the released volatile components in FTIR spectra, while phenol (27%), phenol,4-(1-methylethyl) (40%), and p-isopropenylphenol (34%) were the major compounds in GC-MS analysis. Whereas, the kinetic results showed that both isoconversional methods can be used to determine activation energies, which were estimated 165 KJ/mol (KAS), 193 KJ/mol (FWO), 180 KJ/mol (Friedman), 177 KJ/mol (Vyazovkin), and 174 KJ/mol (Cai).

Keywords: glass fibre-reinforced epoxy resin composites; pyrolysis; TG-FTIR-GC–MS analysis; pyrolysis kinetic; mechanical pre-treatment

1. Introduction

Glass fibre-reinforced epoxy resin composites (GFRC) is a well-established and essential material in the manufacture of aircraft, vehicle, and infrastructure structures [1,2]. It has wide applications in defense, electronics, renewable energy, etc. due to its outstanding...
physical properties including light weight and high chemical, mechanical, and thermal durability [3–5]. According to recent studies, the market for fibre-reinforced composites in the USA has reached $12 billion in 2020 with an expected annual growth rate of 6.6% due to its adaptation in many modern applications such as wind energy [6–8]. This heavy use has lead to producing a huge quantity of GFRC waste on a regular and increasing basis in the world [9]. The aircraft, wind energy, and electronic (waste printed circuit board) sector is the largest contributor to this share of production [9–12]. Usually, GFRC waste is composed of several layers of glass fibre collected together using resin. In addition, this type of waste can be classified based on the type of resins into thermosets (epoxy resin) and thermoplastic (acrylic poly-methyl methacrylate (PMMA)) [6,13]. In some applications, this composition is presented as a mixture with copper and other mineral layers for electrical conductivity [14,15], which are classified as heavy metals and polluting elements for soil and groundwater, as well as resins that are classified as toxic materials [16].

Therefore, it must be disposed of safely and no longer disposed by traditional methods, bad burial, burning, or throwing of waste, which are the most popular disposal methods adopted by composite industries [17] to avoid the accumulation of waste and the resulting obstacles. In addition, it can also help keep up with this strong demand for virgin glass fibres [18]. Despite all these defects, GFRC waste has a high economic value for these compounds if these materials (glass fibre and epoxy resin) are recovered in a cost-effective way without causing negative environmental effects [19]. All these factors have led to an increase in awareness about ways to dispose of such waste, where the mechanical, chemical, and thermal processes were adapted to extract the fibre and epoxy in the form of fillers or petroleum products [20,21]. Usually, mechanical treatment is used for size reduction, and converted GFRC waste into fine particles (fibres mixed with resin) in micro-size can be used as filler material in concrete, asphalt, wood, plastic, etc. [22–24]. In order to remove resin from the milled fibre particles, chemical treatment using organic solvents was used to dissolve epoxy resin [25], while epoxy resin fraction can be recovered from the organic solutions using a rotary evaporator. Although both processes succeeded in achieving the specific goal, some limitations have been appearing such as the need for a lot of chemicals, power consumption, etc., which makes it hard to apply for industrial scale [9].

Therefore, a pyrolysis process was used to decompose GFRC into energy products (e.g., gas, oil, and char mixed with short glass fibres) with a high calorific value in a wire-mesh reactor [26]. In addition, the thermal and chemical decomposition of GFRC was studied using thermogravimetric coupled with Fourier transform infrared spectroscopy (TG-FTIR) [27,28]. In addition, the kinetic parameters of GFRC pyrolysis were determined using different methods, including linear isoconversional methods (Kissinger–Akahira–Sunose (KAS), Flynn–Wall–Ozawa (FWO), and Friedman) and nonlinear isoconversional method (e.g., Vyazovkin). It is worth mentioning that the experiments in these studies were performed on GFRC waste in the form of a by-product collected from acid solution storage tank manufacturing factory without specific composition [27]. In another study the experiments were performed on 67% E-type glass fibre and 6509 epoxy resin without giving any attached to the composition of formulating products using GC analysis [28]. As shown, these studies did not take into account the size of waste and pre-treatment using mechanical processes, even though these practices are a necessary part of such treatments [29,30].

In this context, this research aims to study the effect of pre-treatments on the pyrolysis of GFRC. As shown, the pyrolysis experiments were performed on feedstock that has a different structure and is not similar to the commercial products. In order to put the layer of pyrolysis of GFRC and understand better its thermal and chemical decomposition and its pyrolysis kinetic parameters, this work aims to study the pyrolysis behavior of GFRC using TG-FTIR and TG-GC-MS measurements (separately). In addition, the kinetic pyrolysis of GFRC was studied using linear and nonlinear isoconversional methods. Finally, the TG curves were plotting using the distributed activation energy model (DAEM) and Independent Parallel Reactions Kinetic Model (IPR) [31,32].
2. Experimental

2.1. Materials and Design of the Research Experiments

Glass fabric Panda™ (Weave: Twill 2/2 type and weight: 163 g/m²) was purchased from R&G Faserverbundwerkstoffe GmbH, Germany. Epoxy resin and its hardener (EPIKOTE Resin MGS® RIMR 135 and EPIKURE Curing Agent MGS® RIMH 1366) were supplied by Momentive. Other chemicals used in this research were purchased from Sigma-Aldrich, and the gases used in pyrolysis experiments were supplied by Lithuanian Energy Institute. Figure 1 shows the flowchart of the experiments and analysis in the present research. As shown, the research was designed in five steps: (a) preparation of fibreglass/epoxy laminate panels using vacuum-assisted resin transfer method, (b) characterisation of GFRC, using elemental, proximate, and composition analysis, (c) study of the thermal decomposition of GFRC using thermogravimetric analysis, (d,e) study of the chemical compounds of the obtained volatile compounds using FTIR and GC/MS measurements, (f,g) study of pyrolysis kinetic parameters of GFRC using linear and nonlinear isoconversional methods, and (h,i) fitting the TG-DTG exponential data of GFRC waste using DAEM and IPR models, respectively. All these phases and their optimum conditions are illustrated in details in the following sections.

2.2. Preparation of GFRC Panel

In order to prepare the fibreglass/epoxy laminate panel (with a surface area of 100 cm² and thickness 1 mm), four Glass fabric Panda™ sheets (with nominal size of 10 × 10 cm) were cut from the supplied fabric roller; then, they were adhesively stuck together with epoxy/hardener solution using the vacuum-assisted resin transfer method. After that, the fabricated panels were post-cured using an infrared lamp at 75 °C for 8 h, exposed to the main curing treatment in oven at 85 °C for 6 h for achieving of the cross-linking of polymer structure and the homogeneity, thus preparing the final GFRC panel [1,2]. Finally, the cured panel was cut into small pieces (100 mm²) and then crushed into fine particles to minimise mass and heat-transfer resistances during the thermal decomposition using TGA analysis [33]. The milling process was performed using ball milling (mill model Fritsch P-5) at the frequency of 20 Hz for 30 min grinding, thus preparing a powder having a high degree of fineness from the crushed GFRC [25].

Figure 1. The layout of the present research.
2.3. Characterisation of the Milled GFRP

The morphology of the obtained GFRC panel and the crushed GFRP samples were observed using Scanning Electron Microscopy (SEM) and Metallurgical Microscope, receptivity. The carbon (C), nitrogen (N), hydrogen (H), and sulphur (S) content in the milled GFRP sample was measured with an Elemental Analyser (Perkin Elmer 2400 CHN), while the proximate analysis was used to determine the amount of moisture, volatile matter, and ash in the milled GFRP sample according to ASTM standard methods (E1756-01, E872-82, and E1755-01) [34]. It is worth mentioning that the oxygen (O) and fixed carbon content in elemental and proximate analysis was calculated by difference. In order to improve the overall accuracy of the final results, all measurements were repeated three times, calculating the average values.

2.4. Thermogravimetric Measurements

Thermogravimetric analysis (TGA; model: STA449 F3; NETZSCH, Selb, Germany) was used to evaluate the thermal decomposition of the milled samples in nitrogen ambient. The thermal decomposition experiments using TGA were performed on 7–10 mg from each sample until 900 °C with nitrogen flow rate: 60 mL min\(^{-1}\). In order to study the effect of heating rate on the thermal decomposition and chemical decomposition of GFRC (in the next section), the TGA experiments were carried out at different heating rates: 5, 10, 15, 20, 25, and 30 °C min\(^{-1}\), because several studies showed that the heating rate has a significant effect on the yield of the formulated volatile products. In addition, the intensity of these volatile compounds increased with the increase of the heating rate [34,35]. Meanwhile, the kinetic analysis using linear and nonlinear isolation methods was performed on three different heating rates lower than 20 °C min\(^{-1}\), particularly 5, 10, and 15 °C min\(^{-1}\) [35]. Finally, the weight loss obtained at each heating rate was recorded versus pyrolysis temperatures using Pyrys software. Then, fitting TGA curves was followed by estimating DTG curves through numerical derivation of the obtained TGA data and then fitting the DTG curves and determining the maximum thermal decomposition peaks and their intensity and position. Finally, the pyrolysis parameters of the decomposed GFRP at heating rates (5–30 °C min\(^{-1}\)) and their effect on the devolatilisation index (Di) of volatile matters released during the pyrolysis experiments using TGA were determined using Equation (1), where AT is defined as the changing in temperature in the range equal Rd/Rmax = 0.5 and Rd is defined as a decomposition rate, and all parameters can be extracted from TGA-DTG curves [36]. In addition, the heat-resistance index (THRI) was determined (using Equation (2)) to measure the ability of GFRC to resist a heat flow and confirm the thermal stability trend observed using TGA-DTG measurements [37]. It is worth mentioning that T\(_5\) and T\(_{30}\) are defined as the temperatures at 5% and 30% of weight losses, respectively and can be extracted from TGA data.

\[ D_i = \frac{\text{weight loss rate (R}_\text{max})}{\text{Initial decomposition temperature (T}_i\text{)) × Maximum decomposition temperature (T}_m\text{) × (AT)}} \]  

(1)

\[ \text{THRI} = 0.49 \times [T_5 + 0.6 \times (T_{30} - T_5)] \]  

(2)

2.5. Chemical Analysis of the Formulated Chemical Compounds

The functional groups of the formulated chemical compounds at the maximum decomposition temperatures for each heating rate were examined using TG-FTIR. In order to identify and quantified the chemical compounds of these formulated volatile products correctly, the thermogravimetry–gas chromatography–mass spectrometry (TG–GC–MS, Thermo Scientific ISQ™ single quadrupole GC–MS) was used. The GC–MS measurements were carried out using a laboratory test rig composed of a micro Automation Autoinjector™ unit connected with TGA analyser to collect the formulated gases at the specified temperatures, then analysing their chemical compounds using GC–MS (Shimadzu GC-2010) according to the following conditions: scanning range 30–600 m/s, pumping time (20 s),
Table 1. Linear and nonlinear isoconversional methods used to determine kinetic parameters for the pyrolysis of GFRC [40–45].

| Equation No. | Method                      | Expressions               | Plots                     | Slope Value |
|--------------|-----------------------------|---------------------------|---------------------------|-------------|
| (3)          | Kissinger–Akahira–Sunose    | \[ \ln \left( \frac{\beta}{T^2} \right) = \ln \left( \frac{\text{AR}}{E_{\beta}(y)} \right) - \frac{E_a}{RT} \] | \( \ln(\beta/T^2) \) versus \( 1/T \) | \(-E_a/R\) |
| (4)          | Flynn–Wall–Ozawa            | \[ \ln \beta = \frac{\text{lnAEa}}{RT} - 5.335 - \frac{0.516E_a}{RT} \] | \( \ln\beta \) versus \( 1/T \) | \(-1.0516E_a/R\) |
| (5)          | Friedman                    | \( \ln \left( \frac{\text{dy/dt}}{\beta} \right) = \ln(\text{Af}(y)) \left( \frac{-E_a}{RT} \right) \) | \( \ln(dy/dt) \) versus \( 1/T \) | \(-E_a/R\) |
| (6)          |                               | \( (\alpha) = \int_0^\infty \frac{dy}{\beta \text{AEa}} = A \int_0^\infty \exp(-E/RT)dt \) |                                             | \(-E_a/R\) |
| (7)          | Vyazovkin                   | \( \Phi(E_y) = \sum_{i=1}^{n} \frac{n_i}{\sum_{j=1}^{n} \frac{n_j}{T_i}} \) |                                             | \(-E_a/R\) |
| (8)          |                               | \( I(E,T) = \int_0^T e^{-E/RT}dT = \frac{RT}{E} e^{-E/RT} h(x) \) |                                             | \(-E_a/R\) |
| (9)          | Cai                         | \( h(x) = \frac{x^4 + 118x^2 + 86x^2 + 96x}{x^3 + 20x^2 + 120x + 120} \) |                                             | \(-E_a/R\) |
| (10)         | DAEM                        | \( \ln \left( \frac{\beta}{T^2} \right) = \ln \left( \frac{\text{AR}}{E_a} \right) + 0.6075 \frac{E_a}{RT} \) |                                             | \(-E_a/R\) |
| (11)         | IPR                         | \( \frac{dm_{\text{calc}}}{dt} = -(m_0 - m) \sum_{i=1}^{3} C_i \frac{dx_i}{dt} \) |                                             | \(-E_a/R\) |
| (12)         |                               | \[ \text{Dev.}(\%) = \frac{100 \sqrt{\text{F.O.}_{\text{proc}}(Z-N)}}{\max(|dm/dt|)} \] |                                             | \(-E_a/R\) |

3. Results and Discussion
3.1. Microstructure of the Fabricated GFRC

Figure 2 shows the microstructure cross-section and morphology SEM images of the fabricated panel (after and before crushing). It is clear that the cured panel was composed of many laminates containing destroyed weft (horizontal direction) and warp yarns (vertical direction) resulting from the shear stress produced during cutting the samples. These yarns are joined together using epoxy resin. Figure 2A shows the shape of the warp yarns (vertical direction) that have round shapes with similar size and good distribution and covered with epoxy debris in the form of flakes and bulk particles, which indicate that failure...
was occurring in the form of ductile fracture [46]. In contrast, the surface morphologies of panel surface are very smooth with little debris produced during the cutting process without any cracks or notched, as shown in Figure 2B, which means that the preparation and curing processes were occurring according to the standard methods [47]. Figure 2C shows the metallurgical image of the milled GFRC powder. As shown, the powder has a high degree of fineness in microscale and is composed of two components: fibreglass threads or particles and epoxy resin separated in the form of fibreglass–epoxy composite agglomerations.

Figure 2. (A) SEM micrograph of the fracture cross-section, (B) SEM image of the surface of the fabricated GFRC laminates, and (C) Metallographic image of the milled GFRC sample.

3.2. Basic Properties of GFRC

Table 2 shows the ultimate and proximate analyses of GFRC powder. As shown in the table, the ultimate measurements showed that carbon (C) and oxygen (O) are the main elements in the tested GFRP powder with an average amount of 32.67 wt% and 61.24 wt%, respectively. In addition, a small amount of hydrogen (H) was observed with concentration estimated at 3.94 wt%. The strong presence of these elements (C, O, and H) indicates that GFRP can be considered as a promising source of energy products and carbon precursor. In addition, a small percentage of nitrogen (N) was also noticed during the measurements (2.16 wt%) with a completely absence of sulphur (S), which helps to decrease toxic emissions (e.g., SO₂ and NOₓ) during the pyrolysis process and their potential application at large scale [48]. In contrast, the proximate measurements showed that the GFRP powder is rich with a volatile matter content (42.28 wt%) that contributes to increase the heating value of the formulated fuel during the conversion process. In addition, a high amount of ash content (55.1 wt%) was observed that can act as a catalyst during the thermal decomposition, which leads to change reactivity of the feedstock (GFRC powder) to be more activated and increased yield and quality of the obtained oil [49].

Table 2. Ultimate and proximate analysis of the milled GFRP.

| Elemental Analysis (wt%) | Proximate Analysis (wt%) |
|-------------------------|--------------------------|
| N          | C          | H          | S          | O          | Moisture | Volatile Matter | Fixed Carbon | Ash         |
| 2.16 ± 0.09 | 32.67 ± 0.23 | 3.94 ± 0.06 | <0.01 ± 0.00 | 61.24 ± 0.26 | 0.08 ± 0.00 | 42.28 ± 0.13 | 2.54 ± 0.07 | 55.1 ± 0.18 |

3.3. TG-DTG Data Analysis

Figure 3 shows the TGA-DTG curves of the milled GFRC for all tested heating rates. It seems that the milled samples are decomposed in successive three stages as shown in the TGA analysis curves (Figure 3A) with almost similar features even with changing heating rates, but with different weight loss and position of decomposition zones. As shown, the first weight loss is located in the range of 50–200 °C (Y1) due to evaporation moisture with a very small weight loss <1 wt%. After that, another weak decomposition zone (Y2) was observed due to decomposing some organic components in epoxy and hardener [50], while the main degradation stage (Y3) was located in the range of 317–467 °C with a weight loss of 38 wt%. This degradation zone is related to the full degradation of epoxy and
its compounds, especially the bromine element [51]. In the last decomposition stage, a significant decomposition was observed (42–46%) due to the formulated char mixed with the fibreglass particles, which needs a very high temperature for decomposition them, and that is why it remained a solid residue [52]. Actually, these results agree with the preparation matrix conditions and concentrations, which showed the amount of epoxy estimated at 38%, which means that all the epoxy fraction was decomposed and fibreglass remained mixed with char.

Finally, DTG curves showed that all the feedstock components are decomposed together in a major single peak reaction (since the weight loss of other stages was very small). It seems that all samples had similar weight loss, which indicate that the heating rates did not have an effect on the weight loss and the main changes were happening in the chemical structure of the formulated components as shown in the next sections. In addition, it was observed that the maximum decomposition temperature increased gradually from 400 to 430 °C by increasing the heating rates as a result of increasing the applied temperature on the surface area of the milled samples, which led to increasing the generated heat fluxes, facilitating their transmission inside the internal molecular and accelerating the decomposition reaction in less time until decomposing the feedstock completely [34]. All the pyrolysis characteristic parameters are shown in Table 3.

![Figure 3. (A) TGA and (B) DTG curves of FGE and GFRC at different heating rates.](image)

Table 3. The pyrolysis characteristic parameters for GFRC at different heating rates.

| Pyrolysis Parameters | 5   | 10  | 15  | 20  | 25  | 30  |
|----------------------|-----|-----|-----|-----|-----|-----|
| Onset temperature Ti (°C) | 256 | 260 | 278 | 308 | 286 | 279 |
| Tm (°C)              | 346 | 361 | 367 | 357 | 364 | 355 |
| Tf (°C)              | 477 | 478 | 500 | 462 | 454 | 466 |
| Rmax (%/min)         | 2.9 | 5.8 | 7.8 | 14.6 | 18.7 | 21.9 |
| Di (% min^{-1} °C^{-3}) | 5.2 × 10^{-7} | 8.7 × 10^{-7} | 9.8 × 10^{-7} | 2.2 × 10^{-6} | 2.9 × 10^{-6} | 3.9 × 10^{-6} |
| ΔT                  | 63  | 71  | 78  | 60  | 62  | 57  |
| Mf (%)              | 58.2 | 58.03 | 56.69 | 60.59 | 58.25 | 57.87 |
| T5                  | 301.4 | 310.8 | 314.9 | 319.2 | 317 | 316.6 |
| T30                 | 364.5 | 377.1 | 384.6 | 380 | 394.9 | 393.5 |
| THRI                | 166.24 | 171.78 | 174.79 | 174.28 | 178.23 | 177.74 |
3.4. Chemical Analysis of the Synthesised Chemical Compounds

FTIR coupled with TG was used to examine the obtained chemical compounds at the maximum decomposition temperatures (based on the DTG results) of GFRC for all the specified heating rate ranges. The FTIR measurements were performed in the ranges from 346 °C to 364 °C, and all results are shown in Figure 4. As shown in the 2D FTIR spectra, the tested GFRC samples under the specified heating rates give almost the same features in terms of the generated functional groups, particularly sharp peaks in the range 1200–1250 cm\(^{-1}\) assigned to aromatic benzene compound, 1300 and 1500 cm\(^{-1}\) related to the N-O group, 1700 cm\(^{-1}\) being the typical place for carbonyl (C=O), 2300 cm\(^{-1}\) being the typical place of CO\(_2\), and 3600 cm\(^{-1}\) due to an aromatic and aliphatic C-H group. It seems that the decomposed samples are rich with aromatic compounds and the absorbances of these compounds increased significantly as the heating rate increased as a result of the abundance in the produced heat flux, which led to decomposition of the whole complex of organic molecules into aromatic compounds [53]. Meanwhile, the 3D FTIR spectra showed that the disturbance peaks disappeared as the heating rates increased, which was proof that all the organic components in epoxy resin were decomposed to energy and chemical compounds. GC-MS measurement was used in the next section to determine the main composition and yield of these formulated compounds.

![Figure 4. Two-dimensional (2D)-3D FTIR analysis of the decomposed milled GFRP at different heating rates.](image)

3.5. Chemical Analysis of the Synthesised Chemical Compounds Using GC–MS

Figure 5 shows the GC–MS results of the generated chemical compounds at maximum decomposition zones of GFRP samples under the different heating rates, and the obtained chemical compounds with their respective peak areas are shown in Table 4. According to GC-MS analysis, the synthesised volatile products are composed of many chemical compounds. Phenol (4.25–26.99%), phenol, 4-(1-methylethyl) (10.31–40.08%), and p-Isopropenylphenol (23.64–34.21%) were represented as the major compounds in GC-MS analysis, and these compounds can be separated using membranes or other technologies [54,55]. As shown, the yield of these compounds changed with increase of the heating rate due to the significant increase in the produced heat flux during the process. The strong presence of phenol and p-isopropenylphenol compounds confirms that the formulated products from the pyrolysis of GFRC were typical energy products [56,57]. In addition, these pyrolysis compounds (phenol, p-isopropenylphenol, etc.) can be used in many fields such as chemicals, fuel, pharmaceuticals, etc. Based on these results, the pyrolysis process is recommended to convert GFRC into high added energy products and the recovery of short glass fibres, especially at high heating rates.
3.6. Kinetic Analysis of FEC Pyrolysis
3.6.1. Estimation of Activation Energies Using Isoconversional Methods

Figure 6 shows KAS, FWO, and Friedman fitting curves for conversion rate from 10% to 90% at heating rates: 5, 10, and 15 °C min\(^{-1}\). The activation energies can be calculated from the slope of these plotting curves, which are represented by \(-\frac{E_a}{R}\) (KAS and Friedman) and \(-1.0516\frac{E_a}{R}\) (FWO). As shown from the fitting curves, KAS and FWO plots are composed of parallel straight lines in the main conversion range (20–80%). Meanwhile, the Friedman plot is composed of straight lines distributed randomly with a big variation between the fitting points, which means that FWO and KAS models are valid techniques to model the reaction mechanism of GFRC pyrolysis in the major conversion phase, and the values of the calculated \(E_a\) at all conversion rates are summarised in Figure 7 and Table 5. In addition, the coefficient of determination \(R^2\) was calculated to evaluate the accuracy of the fitted curves as shown in Table 5. As shown, the average \(E_a\) is estimated at 164.5886 kJ mol\(^{-1}\) (KAS), 192.6161 kJ mol\(^{-1}\) (FWO), and 180.3665 kJ mol\(^{-1}\) (Friedman). It is clear that all models gave much less error with \(R^2\) in the ranges 0.9804-0.9918. As mentioned in the Introduction section, several studies have been developed to determine the \(E_a\) of pyrolysis of GFRC and were estimated at 28.17 kJ/mol (GFRP waste was a by-product from the acid solution storage tank manufacturing factory) and 41.4–78.4 kJ/mol (for by-product from reuse plastic fuel) [26,27], which means that the GFRC (fibreglass and epoxy) produced an \(E_a\) higher than other types of GFRC by 78% and 52%, respectively. This is due to the fact that the GFRC used in the current research was composed of two mainly components, including glass fibre and epoxy, and each component has a different \(E_a\). However, the glass fibre did not decompose alongside epoxy in a single reaction (based on the DTG results), and the obtained \(E_a\) represents the summation of \(E_a\) of all the decomposition of the epoxy component only. This is why the pyrolysis of GFRC gave a higher \(E_a\) value compared to other feedstocks, and it is supposed to compare with pure epoxy, which was estimated at 130 to 230, 78–262, and 76.2–327 kJ/mol [58]. This difference in results is due to the different chemical composition, size, crystallinity, and testing ambient, which led to the digestibility of GFRC and conversion into energy products [59,60]. In order to improve the accuracy of these results, nonlinear isoconversional methods were used in the next section.
| Compounds | 5 °C/min | 10 °C/min | 15 °C/min | 20 °C/min | 25 °C/min | 30 °C/min |
|-----------|----------|-----------|-----------|-----------|-----------|-----------|
| Silane, [4-[1,2-bis(trimethylsilyloxy)ethyl]-1,2-phenylene]bis(oxy)bis[trimethylsilyl] | 27.64 | 27.64 | 27.64 | 27.64 | 27.64 | 27.64 |
| Silane, [4-[1,2-bis(trimethylsilyloxy)ethyl]-1,2-phenylene]bis(oxy)bis[trimethylsilyl] | 3.61 | 3.61 | 3.61 | 3.61 | 3.61 | 3.61 |
| N-[5-(3-Hydroxy-2-methylpropenyl)-1,3,4,5-tetrahydrobenzo[c]d][indol-3-yl]-N-methylacetamide | 40.08 | 40.08 | 40.08 | 40.08 | 40.08 | 40.08 |
| 2-Ethylacridine | 3.13 | 3.13 | 3.13 | 3.13 | 3.13 | 3.13 |
| Cyclotrisiloxane, hexamethyl | 5.18 | 5.18 | 5.18 | 5.18 | 5.18 | 5.18 |
| Octasiloxane, 1,1,3,3,5,5,7,7,9,9,11,11,13,15,15-hexadecamethyl | 29.06 | 29.06 | 29.06 | 29.06 | 29.06 | 29.06 |
| 1,2-Benzisothiazole-3-amine | 2.42 | 2.42 | 2.42 | 2.42 | 2.42 | 2.42 |

### Table 4. GC-MS compounds generated at 5–30 °C/min.
Table 4. Cont.

| 5 °C/min | 10 °C/min | 15 °C/min | 20 °C/min | 25 °C/min | 30 °C/min |
|----------|-----------|-----------|-----------|-----------|-----------|
| Time (min.) | GC Compounds | Area (%) | Time (min.) | GC Compounds | Area (%) | Time (min.) | GC Compounds | Area (%) | Time (min.) | GC Compounds | Area (%) | Time (min.) | GC Compounds | Area (%) |
| 23.997 | 1,1′-Biphenyl, 4-phenoxy- | 2.22 | 26.130 | Cyclohexane, 1-ethyl-2-propyl- | 3.56 | 26.136 | Cyclooctane, 1-methyl-3-propyl- | 3.74 | 26.138 | 1,1′-Biphenyl, 4-phenoxy- | 2.69 |
| 24.358 | Silane, 1,4-phenylenebis(trimethyl | 2.36 | 26.130 | Cyclohexane, 1-ethyl-2-propyl- | 3.56 | 26.136 | Cyclooctane, 1-methyl-3-propyl- | 3.74 | 26.138 | 1,1′-Biphenyl, 4-phenoxy- | 2.69 | 26.136 | Cyclosiloxane, hexamethyl- | 3.69 |
Figure 6. Plots of isoconvensional methods at different values of conversion.

Figure 7. The relationship between apparent activation energy and conversion rates.

Table 5. The estimated activation energy at different conversion rates.

| y  | KAS (KJ/mol) | R²  | FWO (KJ/mol) | R²  | Friedman (KJ/mol) | R²  | Vyazovkin (KJ/mol) | R²  | Cai (KJ/mol)  | R²  |
|----|--------------|-----|---------------|-----|-------------------|-----|--------------------|-----|---------------|-----|
| 0.1| 175          | 0.9355 | 203          | 0.9381 | 221               | 0.9916 | 186               | 0.9368 | 184          | 0.9325 |
| 0.2| 205          | 0.9871 | 235          | 0.9891 | 240               | 0.9992 | 217               | 0.9902 | 215          | 0.9882 |
| 0.3| 203          | 0.9835 | 233          | 0.9861 | 222               | 0.999 | 215               | 0.9857 | 213          | 0.985  |
| 0.4| 165          | 0.9945 | 193          | 0.9955 | 154               | 0.9992 | 177               | 0.995  | 174          | 0.9951 |
| 0.5| 169          | 0.9946 | 198          | 0.9956 | 165               | 0.999 | 181               | 0.9951 | 179          | 0.9951 |
| 0.6| 139          | 0.9822 | 165          | 0.9861 | 142               | 0.9895 | 152               | 0.986  | 148          | 0.9844 |
| 0.7| 140          | 0.9822 | 167          | 0.986  | 151               | 0.987 | 153               | 0.9856 | 149          | 0.9844 |
| 0.8| 140          | 0.9821 | 168          | 0.986  | 165               | 0.9996 | 154               | 0.9854 | 150          | 0.9843 |
| 0.9| 141          | 0.9821 | 168          | 0.986  | 159               | 0.9613 | 156               | 0.9853 | 151          | 0.9843 |
| Avg.| 164          | 0.9804 | 192          | 0.9831 | 180               | 0.9918 | 177               | 0.9827 | 174          | 0.9814 |

3.6.2. Estimation of Activation Energies Using Nonlinear Isoconvensional Methods

The activation energy at each conversion rate was determined numerically using nonlinear isoconvensional methods (Vyazovkin and Cai approaches) after several iterations at 5, 10, and 15 °C min⁻¹. The solution started with Ea = 200 kJ/mol (as an initial condition) until the values of Ea became constant; then, it took these values as optimal values of Ea and
all iterations, as shown in Tables 6 and 7. As shown in the tables, Ea values became constant
in both cases at the third iteration. These values were used to plot Vyazovkin and Cai curves
at all conversion rates as shown in Figure 8, where the Y-axis in the case of Vyazovkin
is represented by Equation (14), while Equation (15) represents the Y-axis in case of Cai
methods [44,45]. It is clear that the determined Ea values using nonlinear isoconversional
methods were fully matched with those calculated by linear isoconversional methods
with $R^2 > 98$, which means that linear and nonlinear isoconversional methods are valid for
simulating the pyrolysis kinetics of GFRC.

\[
\ln \left\{ \frac{\beta_i}{T_{y,i}^2} h(x_{y,i}) - \frac{x_{y,i}^2 e^{x_{y,i}}}{x_{y,i-0.1}^2 e^{y_{y,i-0.1}}} h(x_{y,i-0.1}) \right\} \tag{14}
\]

\[
\ln \left\{ \frac{\beta_i}{T_{\alpha,j}^2} h(x_{\alpha,j}) - \frac{x_{\alpha,j}^2 e^{x_{\alpha,j}}}{x_{\alpha,j-\Delta\alpha,j}^2 e^{x_{\alpha,j-\Delta\alpha,j}}} h(x_{\alpha-\Delta\alpha,j}) \right\} \tag{15}
\]

Table 6. The determined activation energy using the Vyazovkin method at different number of iterations.

| Conversion (y) | Initial Value | First Iteration | Second Iteration | Third Iteration | Fourth Iteration |
|---------------|---------------|-----------------|------------------|-----------------|------------------|
| 0.1           | 200           | 185.732984      | 186.312627       | 186.665928      | 186.665928       |
| 0.2           | 200           | 216.605307      | 217.280157       | 217.693776      | 217.693776       |
| 0.3           | 200           | 213.9912192     | 214.657926       | 215.066552      | 215.066552       |
| 0.4           | 200           | 176.9886399     | 177.5400617      | 177.87803       | 177.87803        |
| 0.5           | 200           | 180.934589      | 181.4983048      | 181.843808      | 181.843808       |
| 0.6           | 200           | 151.9479942     | 152.4214001      | 152.711552      | 152.711552       |
| 0.7           | 200           | 152.4360676     | 152.9109941      | 153.202078      | 153.202078       |
| 0.8           | 200           | 153.867198      | 154.3465832      | 154.6404        | 154.6404         |
| 0.9           | 200           | 155.3727803     | 155.8568563      | 156.153548      | 156.153548       |
| Average       | 200           | 176.4307104     | 176.980394       | 177.317297      | 177.317297       |

Table 7. The calculated activation energy using the Cai method at different number of iterations.

| Conversion (y) | Initial Value | First Iteration | Second Iteration | Third Iteration | Fourth Iteration |
|---------------|---------------|-----------------|------------------|-----------------|-----------------|
| 0.1           | 200           | 184.720452      | 184.712138       | 184.712138      | 184.712138       |
| 0.2           | 200           | 215.557078      | 214.675974       | 215.62359       | 215.62359       |
| 0.3           | 200           | 213.154332      | 212.286976       | 213.220844      | 213.220844       |
| 0.4           | 200           | 174.751966      | 173.945508       | 174.80185       | 174.80185       |
| 0.5           | 200           | 179.4768122     | 178.751          | 179.49926       | 179.49926       |
| 0.6           | 200           | 148.90374       | 148.895426       | 148.895426      | 148.895426       |
| 0.7           | 200           | 149.851356      | 149.252928       | 149.843222      | 149.843222       |
| 0.8           | 200           | 150.64968       | 150.059386       | 150.641366      | 150.641366       |
| 0.9           | 200           | 151.506022      | 150.94067        | 151.497708      | 151.497708       |
| Average       | 200           | 174.2857354     | 173.656356       | 174.3039338     | 174.3039338     |
3.6.3. Fitting of TGA-DTG Data Using DAEM and IPR

Figures 9 and 10 shows the TGA-DTG experimental data (red dotted lines) and fitting (blue dotted lines) curves of TGA-DTG at 5–30 °C/min for GFRC sample using Equations (11) and (12). It seems that the plotting TGA-DTG data fully matched with the experimental curves with a deviation less than <1 for all heating rates in both cases DAEM (TGA curves) and IPR (DTG curves), indicating that the DAEM and IPR models succeeded in the fitting of thermal decomposition curves of GFRC pyrolysis. Based on the DAEM and IPR models, activation energy and pre-exponential factors were determined, and all values are summarised in Table 8. According to the TG-FTIR-GC-MS analysis of GFRC, the pyrolysis process can be considered as a promising approach to dispose of GFRC and convert them into energy components and fibre powder. In addition, nonlinear isoconversional methods were successful to determine kinetic parameters and to plot TGA-DTG curves of GFRC.
4. Conclusions

In this research, the basic fundamentals of pyrolysis treatment fibreglass/epoxy composites (GFRC) using the TG-FTIR-GC-MS measurements were studied. The pyrolysis kinetics of GFRC were also investigated using linear and nonlinear isoconversional methods, including KAS, FWO, Friedman, Vyazovkin, and Cai models. The research started with a production of the GFRC panel; then, it analysed the element and ultimate properties, which was followed by studying the thermal and chemical decomposition properties of GFRC and determining the chemical compounds using the TG-FTIR-GC-MS system. Based on the obtained TGA-DTG curves, the activation energies for all the processes and for each conversion rate was calculated using linear and nonlinear isoconversional methods, followed by modeling TGA and DTG curves using DAEM and IPR models. The measurements and results of pyrolysis GFRC revealed the following:

(A) The TGA-DTG results showed that GFRC decomposed thermally in three phases with a total mass loss of 43% and the major decomposition region was located in the range of 256–500 °C.

(B) TG-FTIR spectra showed that the aromatic benzene and C-H bond were the main volatile compounds in the decomposed samples, and its abundance increased with the increase of heating rate.

(C) GC-MS results showed that phenol (4.25–26.99%), phenol, 4-(1-methylethyl)- (10.31–40.08), and p-isopropenylphenol (23.64–34.21%) were the main volatile and flammable compounds, and their yield was affected significantly by the heating rate.
The kinetic models using linear and nonlinear isoconversional methods revealed that the average activation energies can be estimated at 165 KJ/mol (KAS), 193 KJ/mol (FWO), 180 KJ/mol (Friedman), 177 KJ/mol (Vyazovkin), and 174 KJ/mol (Cai) with $R^2 > 98$.

DAEM and IPR models showed a high performance for plotting of the TGA-DTG experimental data of GFRC samples for all heating rates with deviation lower than <1 for TGA and DTG curves.

Based on the advertised results, the pyrolysis treatment can be employed as a cleaner and sustainable technology for converting epoxy to energy products. In addition, the linear and nonlinear isoconversional methods can be used to model the pyrolysis kinetics of GFRC under applying any heating rate in the main conversion zone (20–80%). The DAEM and IPR models are also highly recommended for plotting the TGA-DTG curves of GFRC with high predictability. In addition, GFRC can be used as a promising new source of renewable energy.

Author Contributions: S.Y.: Conceptualization, Data curation, Formal analysis, Writing—original draft, Writing—review & editing. J.E.: Conceptualization, Data curation, Formal analysis. N.S.: Conceptualization, Data curation, Formal analysis. M.P.: Formal analysis. M.A.A.: Conceptualization, Data curation, Formal analysis, Software. All authors have read and agreed to the published version of the manuscript.

Funding: This research received no external funding.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Not applicable.

Conflicts of Interest: The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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