**Dynamic Model-Free and Model-Fitting Kinetic Analysis during Torrefaction of Oil Palm Frond Pellets**

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**Abstract**

Torrefaction is a thermal conversion method extensively used for improving the properties of biomass. Usually this process is conducted within a temperature range of 200-300 °C under an inert atmosphere with residence time up to 60 minutes. This work aimed to study the kinetic of thermal degradation of oil palm frond pellet (OPFP) as solid biofuel for bioenergy production. The kinetics of OPFP during torrefaction was studied using frequently used iso-conversional model fitting (Coats-Redfern (CR)) and integral model-free (Kissinger-Akahira-Sunose (KAS)) methods in order to provide effective apparent activation energy as a function of conversion. The thermal degradation experiments were conducted at four heating rates of 5, 10, 15, and 20 °C/min in a thermogravimetric analyzer (TGA) under non-oxidative atmosphere. The results revealed that thermal decomposition kinetics of OPFP during torrefaction is significantly influenced by the severity of torrefaction temperature. Via Coats-Redfern method, torrefaction degradation reaction mechanism follows that of reaction order with \( n = 1 \). The activation energy values were 239.03 kJ/mol and 109.28 kJ/mol based on KAS and CR models, respectively.

**Keywords**: Torrefaction; Oil palm frond; Kinetic parameters; Biomass; Bioenergy

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1. **Introduction**

Malaysia has been committed to diversify its fuel source where policies on fuel diversification were first introduced in 1999 to promote renewable energy (RE) usage. Malaysia is blessed with indigenous generation mix of renewable energy resources such as biomass and biomass materials (oil palm mill/plantation residues, forestry biomass, agro-based and farming industries biomass waste), mini-hydro power, solar power and wind energy [1]. In relation to future energy demand and fossil fuel depleting reserves, biomass has been identified as one of the viable renewable energy sources and is gaining importance for possible utilization as co-fuel, particularly in coal-fired power plant. According to Ozturk et al. [2], Malaysia produces approximately 168 million tons of biomass yearly, which includes timber, oil palm waste, rice husks, coconut trunk fibers, municipal, and sugarcane wastes. As for electricity generation from renewable sources, based on Peninsular Malaysia Electricity Supply Outlook 2017 report [3], Peninsular Malaysia has a total licensed generated capacity of 392 MW, with 89 MW (23%) of generation capacity from locally sourced biomass, second to solar photovoltaic (PV) of 235 MW (60%), followed by biogas of 34 MW (9%) and mini hydro of 34 MW (8%). Renewable ener-

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Biomass in raw form has its limitations such as high moisture content, low bulk and energy densities, hygroscopic behavior and poor grindability, which makes it substantial for bioenergy feedstock. Torrefaction, also known as slow pyrolysis, is able to upgrade biomass properties where torrefied solid products are mainly characterized on its upgraded quality, i.e. increased energy density, hydrophobicity, increased brittleness thus improved grindability, microbial and biodegradation resistance, and low bulk density. Understanding of kinetic parameters such as activation energy, pre-exponential factor and reaction mechanism of biomass thermal degradation, where in this case, during limited torrefaction temperature range (200–300 °C), is also in need for torrefaction reactor design and optimization of thermochemical process conditions [8]. To date, oil palm frond characteristics when subjected to torrefaction has not been documented intensively, particularly in densified form. Therefore, the objectives of this study focus on oil palm fronds properties in terms of fuel characteristics, thermal degradation behavior and kinetic analysis during torrefaction. The findings from this study is deemed important in order to produce compatible and viable solid bioenergy feedstock.

2. Materials and Methods

2.1 Torrefaction Experiments

Oil palm frond (OPF) samples were collected from private-owned plantation in Klang province, Selangor, Malaysia. Bulky OPF were chipped and crushed into small particle sizes of approximately 5 mm. Prior to densification, the samples were then dried to achieve moisture content less than 12 wt%. Pelletization was done in Forest Research Institute of Malaysia (FRIM) using pellet machine with a cylindrical die opening of 6 mm in diameter and the length of produced pellets ranged from 1 to 6 cm. Torrefaction experiments were performed by heating 20 g of raw OPF pellets placed in a crucible and positioned in the middle of each zone of horizontal tube furnace OTF-1200X (MTI Corporation, USA) as shown in Figure 1. To ensure inert condition was achieved in the furnace, nitrogen gas was flowed continuously for 15 minutes at 1 L/min. The torrefaction process was carried out at fixed slow heating rate of 10 °C/min from ambient to target temperatures of 200 °C, 250 °C, and 300 °C and upon reaching these values, the heating was held for duration of 60 minutes. Pellet chars collected were weighed and stored in airtight containers and for further use, the samples would again be dried at 105 °C for 24 hours.

2.2 Characterization of Raw and Torrefied OPFP Samples

Proximate and elemental analyses were carried out using Thermo Finnigan Flash 1112 analyzer and thermo balance TGA/SDRA51 (Mettler Toledo), respectively, and followed the procedures of ASTM International E1131-08(2014) and ASTM International D5373-16. High heating values (HHV) of raw and torrefied OPF pellets were determined using a bomb calorimeter (IKA-works C5000) in accordance to ASTM International D5865-13. All results collected were based on duplicate tests and repetitions were done if inconsistency data was observed.

Thermal decomposition during torrefaction of pellet samples were achieved using a thermogravimetric analyzer, TGA/SDRA51e (Mettler Toledo, USA) using nitrogen gas at atmospheric pressure with constant flow-rate of 50 mL/min. To provide inert environment and
preventing secondary volatiles release, nitrogen gas was purged for 15 minutes. For each test, pellet sample size of 5 mg was placed in a 70 µL alumina crucible. TGA's furnace temperature was raised from 25 to 105 °C and heating was held at this temperature for 5 minutes to ensure complete removal of surface moisture and this temperature was also selected to be the basis of kinetic analysis [9]. To achieve complete thermal degradation, the heating was continued up to temperature 800 °C.

2.3 Torrefaction Kinetics of Raw OPFP

For torrefaction kinetic analysis, samples were heated at four dynamic heating rates of 5, 10, 15 and 20 °C/min. In this study, single-step reaction scheme was implemented according to Equation (1) [10]:

\[
\text{Solid Biomass} \rightarrow \text{Char} + \text{Volatiles}
\]

where \( k \) is the global apparent rate constant and dependent on temperature as described in Arrhenius Law and is shown in Equation (2):

\[
k(T) = A \exp \left( \frac{-E_a}{RT} \right)
\]

where \( A \) is the pre-exponential factor (min\(^{-1}\)), \( E_a \) is the apparent activation energy (kJ/mol), \( R \) is the universal gas constant (8.314 J/mol K), and \( T \) is the temperature (in K). For single step solid-state reaction, the differential equation under isothermal conditions can be expressed as Equation (3):

\[
\frac{d\alpha}{dt} = k(T).f(\alpha)
\]

where \( d\alpha/dt \) is the rate of conversion (\( \alpha \)) with time (\( t \)), and \( f(\alpha) \) is the reaction model. Conversion, \( \alpha \) can be defined as in Equation (4):

\[
\alpha = \frac{m_i - m_f}{m_0 - m_f}
\]

where \( m_0 \) is the initial sample weight, \( m_i \) is the sample weight at time \( t \), and \( m_f \) is the final sample weight. For non-isothermal linear heating, \( \beta = dT/dt \) and by substituting Equation (2) into Equation (3), the expression becomes Equation (5):

\[
\frac{d\alpha}{dT} = \frac{A}{\beta} \exp \left( \frac{-E_a}{RT} \right) f(\alpha)
\]

Rearranging Equation (5) and integrating within the limits of 0 to \( T \) for temperature and 0 to \( \alpha \) for conversion gives Equation (6):

\[
g(\alpha) = \int_0^\alpha \frac{d\alpha}{f(\alpha)} = \frac{A}{\beta} \int_0^\infty \exp \left( \frac{-E_a}{RT} \right) dT
\]

The temperature integral term of right side has no methodical solution and in order to simplify Equation (6), the term \( -E_a/RT \) is replaced with \( p(x) \), which gives Equation (7):

\[
g(\alpha) = \left( \frac{AE}{BR} \right) p(x)
\]

Thus, in the current study, the term \( p(x) \) was proposed to be approximated using widely-used isoconversional model-free method, i.e. Kissinger-Akahira-Sunose (KAS) and model-fitting method, i.e. Coats-Redfern. For model-free method, it is known to be efficient enough to handle the complexity of biomass decomposition reaction. Estimation on the temperature integral was based on linearization by Doyle’s empirical approximation as shown in Equation (8) [11]:

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**Figure 1. Torrefaction experimental set-up**

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The KAS equation which relates heating rate and inverse temperature can be expressed as the following Equation (9) [12,13]:

$$\ln \left( \frac{\beta}{T^2} \right) = \ln \left( \frac{AE_a}{Rg(a)} \right) - \frac{E_a}{RT}$$

(9)

At the progressing values of conversion, the slope of straight-line plot of \(\ln \left( \frac{\beta}{T^2} \right)\) versus \(1/T\) allows activation energy to be determined. As for model-fitting kinetic method, Coats-Redfern (CF) is a widely-accepted method which uses the integral form of the non-isothermal rate law where approximation of the temperature integral produces the expression shown in Equation 10 [14]:

$$\ln \left[ \frac{g(\alpha)}{T^2} \right] = -\frac{E_a}{RT} + \ln \left( \frac{AR}{\beta E_a} \right) \left( 1 - \frac{2RT}{E_a} \right)$$

(10)

The above equation can be further simplified in accordance to customary values of \(E_a\) (80–260 kJ/mol) and by assuming \(2RT/E_a << 1\), the simplified form is expressed as in Equation 11 [15]:

$$\ln \left[ \frac{g(\alpha)}{T^2} \right] = -\frac{E_a}{RT} + \ln \left( \frac{AR}{\beta E_a} \right)$$

(11)

A straight-line plot will enable apparent activation energy, reaction mechanism and pre-exponential factor to be determined from the slope and the intercept, respectively. Integral reaction mechanism \(g(\alpha)\) indicating the reaction limiting step is chosen based on the best fit data with the highest correlation coefficient, \(R^2\).

3. Results and Discussion

3.1 Physico-chemical Characteristics and Thermal Decomposition Behaviors of Raw and Torrefied Biomass Pellets

Generally, for a raw biomass to be converted into fuel, the physico-chemical properties need to be studied. The aim is to determine its compatibility and initial properties in its original form, which compared to upgraded biomass fuel when subjected to torrefaction treatment. The characteristics of raw biomass helps in simplifying parameters selection in torrefaction process and thus, determines the best parameters critical in producing good quality biochar compatible for coal substitute. As shown in Table 1, both raw (ROPFP) and torrefied OPF (TOPFP) biomass pellets at temperatures 200 °C (TOPFP200), 250 °C (TOPFP250), 300 °C (TOPFP300), were examined to determine their potential as solid biochar. Moisture contents of both raw biomasses showed low values (<3 wt%) due to open-drying activities prior to pelletization in addition to moisture being driven off during pelletization (die temperature approximately 70 °C).

Figure 2(a) and (b) demonstrate the weight loss curves (TG) and differential thermogravimetric profiles, respectively, for both raw and torrefied pellets of OPF, respectively. For the purpose of clarity and decomposition evalu-
tion between raw and torrefied products, only torrefied biomass samples with holding time 60 minutes were selected for discussion. At high torrefaction temperature, thermal decomposition between raw and torrefied samples displayed a clear difference, as more evidently shown in derivative thermogravimetric (DTG) profiles (Figure 2(b)).

For biomass samples torrefied at temperature 200 °C, hemicellulose content still exists as shown by the existence of small/less apparent shoulder peak, which subsequently caused their thermal decomposition profiles followed that of raw samples. During light torrefaction temperature regime (200–250 °C), thermal degradation was mainly caused by moisture liberation and limited devolatilization of hemicellulose due to minor depolymerization reactions [16]. While for samples torrefied at temperature at 250 °C, characteristic peak / shoulder of hemicellulose disappeared indicating hemicellulose and light volatiles were successfully removed during torrefaction pretreatment. Hemicellulose degrades extensively at temperature ≥ 250 °C and reported to end at temperature approximately 280 °C [17].

In terms of reactivity, based on the derivative mass loss profiles shown in Figure 2, biomass samples torrefied at 200 °C showed greater reactivity than its raw forms. This occurrence was further confirmed via proximate analyses data (Table 1) where volatile matter was found to increase as \( T_{tor} \) of 200 °C then decrease as \( T_{tor} \) further increased from 250 °C to 300 °C. Uneven distribution of biomass macromolecules and possible interactions among the constituents that resulted in formation of new volatiles and possible overlapping decomposition of extractives might be the reasons for this increase of volatile matter and then, released again as heating continues [18]. Halpern et al. [19] have also reported that between these temperatures, thermally unstable hemicellulose degradation occurred due to dehydration reactions which involved bond scissions that released \( \text{H}_2\text{O} \), formation reactions of carbonyl and carboxyl groups accompanied by elimination of CO and \( \text{CO}_2 \) and finally, limited devolatilization and carbonization which produces tars and chars.

DTG peak heights were observed to decrease as torrefaction temperature increased, and shifted to the right, indicating reduction in reactivity as more volatiles were driven off during torrefaction process. This also implies that cellulose and lignin started to decompose with limited carbonization, thus significantly reduced the volatiles of torrefied samples resulting in less reactive product. Peaks representing these two components were more noticeable at temperature above 300 °C, in which according to Yang et al. [20], thermal degradation temperature ranges for cellulose and lignin were 315–400 °C and 160–900 °C, respectively. Trends of TG and DTG profiles in this study are in agreement with previous researches [21–23]. In order to further verify the proposition that torrefied biomass products having similar properties with sub-bituminous coals, DTG profiles of Malaysian sub-bituminous coal (Silantek) were also added in Figure 2(b). Interestingly, torrefied biomass at highest \( T_{tor} \) demonstrates similar profiles as Silantek coal, albeit their maximum peak temperatures were earlier than coal’s. According to Chen and Kuo [24], during torrefaction temperature range, significant decomposition of hemicellulose and partial cellulose occurred at 290 °C, resulting

![Figure 2](image-url)

**Figure 2.** (a) TG and (b) DTG curves of raw and torrefied OPFP at holding time 60 min

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in carbon-rich solid char with coal-like properties.

3.2 Raw OPFP Torrefaction Kinetics

3.2.1 Model-free kinetic evaluation via KAS method

Model-free method is based on the assumption that when activation energy value is constant, the rate of reaction is a function of only temperature [25]. Experimental data with four heating rates (5, 10, 15, and 20 °C/min) for temperature range 105 °C up to 400 °C obtained from thermogravimetric analysis were fitted according to the chosen model-free method, i.e. KAS, in order to calculate the apparent activation energy ($E_a$) and their corresponding correlation coefficients, $R^2$. It should be noted that, as aforementioned, the drying stage was not included in the torrefaction kinetic analysis as at temperatures less than 105 °C only moisture and physically absorbed water was removed [26]. Thus, the analysis of KAS model applied was intentionally limited to torrefaction process with lower temperature set at 105 °C and extended upper temperature limit set at 400 °C. For ease of kinetic analysis interpretation, degradation region of raw biomass under study has been based on temperature range proposed by Yang et al. [27], which are; 220–315 °C (dominant hemicellulose degradation stage) and 315–400 °C (cellulose degradation stage). Thus, the decompositions involved during torrefaction has been henceforth referred to as Stage 1 and Stage 2, respectively, rather than by lignocellulosic composition. The reason is with regard to the selected temperature range (105–400 °C) for the torrefaction kinetic analysis, and taking into account the complexities involved in biomass decomposition, cellulose, hemicellulose and lignin decompositions that might occur simultaneously.

For the current study, integral equation of KAS method (Equation (9)) was applied where the slope of straight line of $\ln(\beta/T^2)$ against $1/T$ was plotted in Figure 3(a), which then allows $E_a$ to be determined. Tabulated values of $E_a$ at different conversions and corresponding correlation coefficients ($R^2$) are shown in Table 2 and displayed in Figure 3(b) of intervals 0.05. It should be noted that the end value of conversion degree ($\alpha$) in these analyses ended at 0.90 where calculated $E_a$ values for all raw samples using each method displayed sudden increase at $\alpha = 0.95$. This rise is possibly due to the sole lignin decomposition where lignin needs higher chemical bond-breaking energy to decompose due to its aromatic chemical structures. This is further confirmed by observing the DTG thermal profile (Figure 2(b)) where cellulose decompositions in raw OPFP ended at around 356–382 °C, depending on individual heating rates. Thus, beyond these temperatures, lignin decomposition is believed to be significant and since lignin is mainly consists of aliphatic and aromatic structures, higher thermal energy is needed to overcome its minimum energy barrier (higher $E_a$), thus the sudden surge in activation energy values at $\alpha = 0.95$.

Average $E_a$ values are noticeably high in ROPFP (239–251 kJ/mol) in which, based on the literatures on various agricultural residues kinetic studies, $E_a$ calculated for various types of biomass are as follows; rice straw (140–267 kJ/mol) [26], rice husk (168 kJ/mol) [28], olive pomace (162–602 kJ/mol) [29], bagasse (169–

![Figure 3. (a) KAS plots and (b) activation energy for ROPFP at different values of conversion](image-url)
kJ/mol) [28], and olive tree pruning (151–209 kJ/mol) [30]. Whereas, from previous studies on woody biomass of different categories (softwood/hardwood) showed lower $E_a$ values via various non-isothermal kinetic methods, for example; ashwood-hardwood (133–167 kJ/mol) [31], poplar wood (159 kJ/mol) [32], Ahun wood (127 kJ/mol) and Araba wood (125 kJ/mol) [33]. Therefore, from this comparison, it can be inferred that agricultural biomass, such as: raw OPFP, showed higher $E_a$ values of 239.03 kJ/mol (Table 3), as compared to woody biomass, which is agreeable to the current results obtained. An increase of $E_a$ values during Stage 1 was also expected where according to Grigiant et al. [31], this is due to the random scission of linear chain in hemicellulose which led to the rise in $E_a$ values as the thermal heating progressed. The crosslinking behavior and its effects on the numerously branched polymeric structure of hemicellulose and extractives constituents that yet to be degraded, gave rise to the $E_a$ values upon thermal heating. In the case of ROPFP, as $\alpha$ reached 0.35 and above, a decreasing trend of $E_a$ values was observed in which the earlier stage of decomposition might not follow the abovementioned explanations. This suggests almost complete degradation of hemicellulose occurred at average temperature of 297 °C, typically described as the end temperature of hemicellulose degradation from literatures [34]. The efficacy of linear fitting for the experimental data obtained via TGA into KAS methods were assessed via correlation coefficients ($R^2$) as listed in Table 2 with obtained good fitting of data where $R^2$ obtained was more than 98%.

3.2.2 Model-fitting kinetic evaluation via Coats-Redfern (CR) method

Model-fitting kinetic method applied in this study, Coats-Redfern (CR) enabled the determination of the complete kinetic triplets i.e. activation energy, reaction mechanism ($g(\alpha)$) and pre-exponential factor ($A$). Based on the highest correlation coefficients obtained from these plots, Table 4 lists the summary of the kinetic parameters attained via CR method for raw OPFP at respective heating rates as well as decomposition stages. Figure 4 displays the plot of $[g(\alpha)/T^2]$ against $1/T$ for the overall torrefac-

Table 2. Kinetic parameters determination using KAS method

| Average T (°C) | Decomposition stage | Conversion, $\alpha$ | $E_a$ (kJ/mol) | $R^2$ |
|---------------|---------------------|---------------------|----------------|-------|
| 253.39        |                     | 0.05                | 251.00         | 0.955 |
| 264.83        |                     | 0.10                | 250.09         | 0.983 |
| 272.83        |                     | 0.15                | 256.07         | 0.974 |
| 279.41        | Stage 1 (dominated by hemicellulose degradation) | 0.20                | 258.17         | 0.979 |
| 285.43        |                     | 0.25                | 261.44         | 0.977 |
| 291.16        |                     | 0.30                | 268.66         | 0.970 |
| 296.97        |                     | 0.35                | 268.00         | 0.976 |
| 302.69        |                     | 0.40                | 255.00         | 0.989 |
| 308.22        |                     | 0.45                | 248.50         | 0.994 |
| 313.36        |                     | 0.50                | 239.49         | 0.999 |
| 318.21        |                     | 0.55                | 221.56         | 1.000 |
| 322.13        |                     | 0.60                | 216.59         | 1.000 |
| 326.16        |                     | 0.65                | 209.73         | 1.000 |
| 329.63        | Stage 2 (dominated by cellulose degradation) | 0.70                | 206.04         | 0.999 |
| 333.14        |                     | 0.75                | 204.83         | 0.999 |
| 336.98        |                     | 0.80                | 211.57         | 0.998 |
| 341.68        |                     | 0.85                | 211.66         | 0.996 |
| 348.94        |                     | 0.90                | 264.24         | 0.983 |
| 367.13        |                     | 0.95                | 403.68         | 0.749 |

Table 3. Summary of ROPFP torrefaction kinetics via KAS method

| Summary of ROPFP decomposition | Conversion range | $E_a$ (kJ/mol) | $R^2$ |
|-------------------------------|-----------------|---------------|-------|
| Overall                       | 0.05< $\alpha$<0.90 | 239.03        | 0.987 |
| Stage 1                       | 0.05< $\alpha$<0.50 | 255.64        | 0.980 |
| Stage 2                       | 0.55< $\alpha$<0.90 | 218.28        | 0.997 |
tion temperature range at heating rate 10 °C/min for various reaction mechanisms (reaction order (F1–F7), diffusion (D1–D4), geometrical contraction (R1–R3), power law (P2–P4) and nucleation models (A2–A4)) along with the stages involved during torrefaction for raw OPFP. As shown in Table 4, Stage 1, which is mainly associated with light volatiles decompo-

### Table 4. Kinetic parameters determination using Coats-Redfern method

| Heating rate (°C/min) | α range   | Mechanistic model | $E_a$ (kJ/mol) | $A$ (min$^{-1}$) | $R^2$  |
|-----------------------|-----------|-------------------|----------------|----------------|--------|
| **Stage 1**           |           |                   |                |                |        |
| 5                     | 0.05<α<0.60 | F3                | 136.13         | 7.75×10$^{11}$ | 0.997  |
| 10                    | 0.05<α<0.50 | F4                | 146.35         | 1.39×10$^{13}$ | 1.000  |
| 15                    | 0.05<α<0.45 | F4                | 146.14         | 1.46×10$^{13}$ | 1.000  |
| 20                    | 0.05<α<0.45 | F5                | 167.25         | 2.10×10$^{15}$ | 1.000  |
| Average               | 0.05<α<0.50 | F3-F5             | 148.97         | 5.32×10$^{14}$ | 0.999  |
| **Stage 2**           |           |                   |                |                |        |
| 5                     | 0.65<α<0.95 | F7                | 807.32         | 6.99×10$^{73}$ | 0.986  |
| 10                    | 0.55<α<0.95 | F4                | 413.05         | 1.46×10$^{137}$| 0.992  |
| 15                    | 0.50<α<0.95 | F2                | 179.73         | 6.72×10$^{155}$| 0.990  |
| 20                    | 0.50<α<0.95 | F1                | 90.01          | 3.56×10$^{7}$  | 0.987  |
| Average               | 0.55<α<0.95 | F1-F7             | 227.60         | 4.85×10$^{36}$ | 0.990  |
| **Overall**           |           |                   |                |                |        |
| 5                     | 0.05<α<0.95 | F2                | 131.95         | 2.49×10$^{11}$ | 0.988  |
| 10                    | 0.05<α<0.95 | F2                | 125.19         | 8.26×10$^{10}$ | 0.985  |
| 15                    | 0.05<α<0.95 | F1                | 89.75          | 2.89×10$^{7}$  | 0.989  |
| 20                    | 0.05<α<0.95 | F1                | 90.24          | 3.85×10$^{7}$  | 0.988  |
| Average               | F1-F2     |                   | 109.28         | 8.28×10$^{10}$ | 0.987  |

Note: F – Reaction order mechanism

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Figure 4. Coats-Redfern plot for raw OPFP at different values of conversion
sition in hemicellulose and extractives, raw OPFP followed reaction-order mechanism with \( n \geq 3 \) whereby as heating rate was increased, reaction order also increased which indicates higher reaction rate, i.e. reactivity. As for stage 2, which is mainly associated to cellulose decomposition, raw OPFP similarly followed reaction order mechanism but \( n^{th} \) order decreased with increase of heating rate.

Expectedly, for overall torrefaction temperature range, reaction mechanism for raw OPFP followed reaction order mechanism, i.e. first reaction order \( (n = 1) \) indicated as F1 with good correlation coefficients \( (R^2) \) obtained in all stages, with \( R^2 \) values of 0.985 to 1. In addition, the average activation energy obtained during the overall temperature range was low at 109.28 kJ/kg that suggest easier start of decomposition stage. However, it should also be noted that for ROPFP at HR5 and HR10, values of activation energies are unexpectedly high and followed 2nd order reaction mechanisms (F2), which suggest the existence of more complex reaction involved at these heating rates, as suggested by few researchers [33,35,36]. In comparison to apparent activation energies obtained via C-R and KAS methods, it was observed that the overall \( E_a \) values for C-R method is much lower by more than This observation might be caused by an assumption of isoconversional method, such as: KAS method which disregards any reaction mechanism to calculate kinetic parameters, whereas model-fitting method determine kinetic parameters using a mass dependent function [36–38].

4. Conclusions

Physico-chemical analyses and thermal behavior of raw and torrefied oil palm frond pellets prepared at different torrefaction temperatures have been studied. Improvements on the properties were achieved via torrefaction treatment where high heating values, fixed carbon and elemental carbon contents significantly increased. The properties are desirable for treated biomass to be used as a potential bioenergy feedstock. Thermal degradation behavior via TGA showed hemicellulose was removed thus producing torrefied biomass nearing the decomposition behavior of coal. Via dynamic kinetic analysis, the variation of activation energies was obtained between the two methods applied (KAS and C-R methods) in which Coats-Redfern method is deemed to be a more reliable method to determine the kinetic parameters where from this study. It was found that thermal decomposition of raw OPFP follows reaction order mechanism.

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