Characterization, kinetics and thermodynamics of epoxidation-esterification of *Irvingia gabonensis* kernel oil methyl ester

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**ABSTRACT**

Epoxidation-esterification of fatty acid methyl ester obtained from *Irvingia gabonensis* kernel oil (IGKO), as well as its characterization, kinetics and thermodynamics were the main focus of this study. The methyl ester obtained via base catalyzed transesterification was used for epoxidation-esterification modification process. Epoxidation kinetics and thermodynamics parameters were also investigated. The properties of the IGKO and epoxidized-esterified *Irvingia gabonensis* kernel oil (IGKO) methyl ester (MIGKO e) were determined using standard methods. Rate constant K and activation energy Ea for the epoxidation process was found to be of the order $10^{-5}$ Lmol$^{-1}$s$^{-1}$ and 46.02 kJ/mol, respectively. ΔG, ΔH, and ΔS values for the epoxidation process were (94.74–101.42 kJ mol$^{-1}$), 43.30 kJ mol$^{-1}$, and $–167.20$ J mol$^{-1}$ K$^{-1}$, respectively, indicating the non-spontaneous, endothermic, and endergonic nature of the process. The physicochemical characteristics of MIGKO e were: $9.1^\circ$C, 298°C, 840 kg/dm$^3$, 13.84 mm$^2$s$^{-1}$, 1.351 mg KOH/g oil, 1.01 mg/kg and 39.78 kV, for pour point, flash point, density, viscosity, acid value, moisture content and dielectric strength, respectively. The MIGKO e properties indicated its potential for use as a bio-transformer fluid, upon further treatment with pour point depressant.

1. Introduction

Increase in global energy demand, instability in world petroleum price, as well as the non-biodegradability of fossil fuels, have increased the demand for bio-oils used in bio-fuels production, as alternative to conventional fuels (Adepoju, 2014). Bio-oils used in the production of products like biodiesels, bio-transformer fluids and bio-lubricants, etc., which are substitute to conventional petroleum base fuels are gaining ground due to their high biodegradability nature, environmental friendliness, availability, energy conservation and management potentials (Adepoju and Olamide, 2014). Most bio-fuels and bio-products like the aforementioned ones are produced by transesterification or alcoholsysis process (Knothe et al., 2007). Other processes include esterification, epoxidation, hydrotreatment and oleaginous microorganisms' treatment (Shote et al., 2009; Agu et al., 2019). These bio-fuels and bio-products are often produced using vegetable oils extracted from seeds/nuts.

However, the primary challenges associated with the production of these bio-fuels and bio-products are limited raw materials and cost of production. Hence, these shortcomings, limit final products (biodiesels, bio-transformer fluids and bio-lubricants etc.) availability (Berhanu and Amare, 2012). In other words, to ensure production competitiveness and availability for commercial purposes, the readily availability and affordability of these vegetable oils raw materials becomes very essential. As a result, several countries like Brazil that depends so much on bio-oils for their energy needs have focused more on the use of vegetable oils for this purpose (Chhetri et al., 2008). Thus, the sustainable supply of less expensive vegetable oils, like *Irvingia gabonensis* kernel oil (IGKO) becomes very essential (Dorado et al., 2006).

Over the years, many researchers have produced the biodiesel, bio-transformer fluid and bio-lubricants from vegetable oil sources. For instance Ong et al. (2013), Sadia et al. (2013) and Rashid et al. (2014) produced biodiesel from *Jatropha curcas* oil, wild safflower (*Carthamus oxyacantha* Bieb) seed oil and milo (*Thespesia populnea*) seed oil,
respectively. Similarly, in transformer fluid production, successes have been reported on bio-transformer oil production from Terminalia catappa L. kernel oil (Agu et al., 2019), Jatropha curcas oil (Beltran et al., 2017) and high oleic sunflower oil (Viertel et al., 2014). Also, in bio-lubricant production, Menkiti et al. (2017a, 2017b) and Aji et al. (2015) produced bio-lubricant from Jatropha curcas oil and Neem seed oil, respectively. As such, there is need to evaluate the feasibility of modifying Irvingia gabonensis kernel oil (IGKO) and its methyl ester by the epoxidation-esterification process for use as transformer fluid.

Irrespective of the modification method used, the need for modification of vegetable oil structure is critical, since it helps to ensure its applicability for use as different products. Therefore, some of the methods of vegetable oil modifications include, but not limited to esterification (Sharma et al., 2006), esterification (Mohammed et al., 2007), epoxidation (Campanella et al., 2010; Giubota-Rosie et al., 2014; Goud et al., 2006) transesterification (Agu et al., 2019; Abdelmalik, 2014; Sanchez et al., 2014), and epoxidation-esterification (Abdelmalik et al., 2011; Agu et al., 2019). Irrespective of method(s) adopted, the unsaturated groups of the fatty acid chains in triglycerides in vegetable oils are intentionally changed to obtain more complex structures. This is to ensure improvements in the low-temperature properties of the oil samples, as well as their oxidative stability (Campanella et al., 2010; Agu et al., 2019).

Of all these methods of vegetable oil modification, classical epoxidation of vegetable oil procedure is of great interest to researchers because of its advantages. These benefits include; high product yield of the process, low cost of the peracid (often acetic acid) synthesis, recyclability of vegetable oil procedure is of great interest to researchers hence, the justification of modifying Irvingia gabonensis kernel oil (IGKO) and its methyl ester by the epoxidation-esterification process for use as transformer fluid.

For instance, with respect to bio-transformer oil production using epoxidation-esterification method, Agu et al. (2019) have assessed the feasibility of using modified Terminalia catappa L. oil as transformer oil. This method was also used by Abdelmalik et al. (2011) for bio-transformer oil production using Palm kernel oil. For bio-lubricant production, Campanella et al. (2010) explored the possibility of producing lubricants from soybean and sunflower oils, by the chemical modification of the oils using epoxidation method. Also, Hwang et al. (2003) worked on the biolubricant production by acid-catalyzed oxirane ring-opening reactions using epoxidized soybean oil, using alcohols and subsequent esterification of the obtained hydroxyl group with acid anhydride. Similarly, Giubota-Rosie et al. (2014) successfully evaluated the role of epoxidation reaction in the modification of Camelina sativa oil for possible use as biodiesel. Also, the epoxidation of waste used oil was successfully used in the synthesis of biodiesel by Kongyi et al. (2013).

Even though vegetable oils epoxidation has been comprehensively studied and reported in literature, it has been discovered that the epoxidation of alkyl ester of base oil, instead of the vegetable oil itself, gives better properties of the intended final product. This is attributed to better product stability (thermo-oxidative) and pour point (Agu et al., 2019; Abdelmalik et al., 2011). Also, the alkyl group with different numbers of double bonds in the fatty acid methyl ester (FAME) significantly affects the epoxidation efficiency, as well as the product selectivity (Huang et al., 2015). As such, several works have been carried out on the epoxidation of alkyl ester of various vegetable oils. For example, Campanella et al. (2008) successfully carried out epoxidation of soybean fatty acid methyl ester (FAME) using performic acid. Similarly, Borugadda and Goud (2014) carried out castor oil methyl esters epoxidation using heterogeneous ion-exchange resin (IR-120) as a catalyst. Epoxidation of methyl esters was also successfully carried out by Shuangfei and Lisheng (2011) using SO3H-functional Boehmite acidic ionic liquid catalyst. Furthermore, the epoxidation of methyl esters obtained from olive and linseed oils was successfully carried out by Huang et al. (2015). During epoxidation reaction, it is essential to understand the kinetics and thermodynamics of the process.

A number of researchers have also successfully carried out research on the kinetics and thermodynamics of vegetable oil and methyl ester epoxidations. For instance, Iikudoria et al. (2007) successfully carried out kinetics and thermodynamics of epoxidation of the methyl esters of Parkia Biglobosa seed oil and obtained the rate constant of the order, activation energy, enthalpy ΔH, entropy ΔS and free energy ΔG values of 10^{-2} L mol^{-1} S^{-1}, 51.963 K J/mol, 13.8 K J/mol, ~ 3.55 K J/mol and 1.51 K J/mol. Similarly, Cai et al. (2008) successfully epoxidized sunflower oil and obtained rate constant of the order 10^{-6} L mol^{-1} S^{-1} and activation energy of 85.21 K J/mol. They also obtained thermodynamic parameter values of 82.73 K J/mol, ~ 83.73 K J/mol and 110.01 K J/mol, for the enthalpy ΔH, entropy ΔS and free energy ΔG, respectively. In a similar manner, Okieimen et al. (2002) successfully carried out the epoxidation of rubber seed oil, with rate constant of the order, activation energy, enthalpy ΔH, entropy ΔS and free energy ΔG values of 10^{-6} L mol^{-1} S^{-1}, 15.7 K J/mol, 15.2 K J/mol, ~ 3.94 K J/mol and 25.44 K J/mol. Also, Naidir et al. (2012) studied the kinetics and thermodynamics of the epoxidation of trimethylolpropane ester, with the rate constant of the order and activation energy values of 10^{-2} L mol^{-1} S^{-1} and 69.4 K J/mol. In the work of Cai and Wang (2011), they successfully studied the epoxidation of unsaturated fatty acid methyl ester (FAMEs) in the presence of SO3H-functional, and obtained the rate constant of the order, activation energy, enthalpy ΔH, entropy ΔS and free energy ΔG values of 10^{-4} L mol^{-1} S^{-1}, 45.5 K J/mol, 42.7 K J/mol, ~ 201.3 K J/mol and 112.1 K J/mol. Goud et al. (2006), successfully studied the epoxidation of mahua oil (Madhunica indica), and obtained the rate constant of the order, activation energy, enthalpy ΔH, entropy ΔS and free energy ΔG values of 10^{-6} L mol^{-1} S^{-1}, 14.5 K J/mol, 13.8 K J/mol, ~ 51.1 K J/mol and 30.6 K J/mol.

However, to the best of our knowledge, there is no report available on the extensive study of the epoxidation of Irvingia gabonensis kernel oil (IGKO) methyl ester using peracetic acid (PAA). Also, there is no literature information on the kinetics and thermodynamics of Irvingia gabonensis kernel oil methyl ester epoxidation, hence, the justification and novelty of the present study. Thus, it has becomes paramount to close these existing research gaps. Therefore, this work focuses on the extensive study of the epoxidation kinetics and thermodynamics of Irvingia gabonensis kernel oil methyl ester, using peracetic acid (PAA). Also, the physicochemical properties of the obtained product were analyses as well.

2. Materials and methods

2.1. Materials

The primary raw material, Irvingia gabonensis kernels (IGK) were bought in Nkwo-Agu Market, Umuaa, Nigeria. Conrars laboratory chemical vendor located in Enugu, Nigeria supplied all analytical grade chemicals/reagents used in research and they were used without further purification. These reagents include n-hexane, KOH, acetic acid, H2SO4. Hydrogen peroxide, NaHCO3, NaCl, Magnesium sulphate, ethyl acetate, acid anhydride, boron trifluoro diethyletherate, and methanol. The entire reagents were of analytical grade with 99 % purity levels.

2.2. Extraction of Irvingia gabonensis kernel oil (IGKO)

Extraction of oil from IGK followed the procedure described in the work of Menkiti et al. (2015). The entire extraction process was repeated times, after which the average value determined and noted. N-hexane was used for the solvent extraction process using soxhlet apparatus to extract IGKO. Percentage oil yield determination was carried out in line
2.3. Transesterification experiment

2.3.1. Acid catalyzed esterification

The acid catalyzed esterification of the IGKO followed the procedure as described in our earlier work (Agu et al., 2019). In the earlier described procedure, a reaction glass vessel was used in heating IGKO sample at 60 °C. Thereafter, H₂SO₄ (1.0 % w/w) catalyst and 99 % pure methanol (30 % v/v) solution was heated at 30 °C for a period of 4 h. This solution was then poured into the glass vessel that contains IGKO to ensure esterification reaction. With the aid of a stirrer operating at 600 revolutions per minute (RPM), the mixture was stirred for an hour. Using a separating funnel, the content was decanted into it and allowed for 2 h, to ensure proper/complete separation. The top layer which consists of methanol-water fractions was discarded, while the oil phase (methyl ester) was used as starting material for base-catalyzed transesterification reaction.

2.3.2. Base catalyzed transesterification

The transesterification of the methyl ester followed the procedure described in the work Agu et al. (2019). 50 ml of methyl ester sample was decanted into a 150 ml conical flask. It was heated to 60 °C using a water bath. Afterwards, potassium methoxide solution was obtained by dissolving KOH pellets in methanol, contained in a 250 ml beaker. The potassium methoxide solution and methyl ester were mixed at a ratio of 6:1. The mixture was eventually stirred vigorously at 500 revolutions per minute (RPM) for 2 h using magnetic stirrer. Thereafter, it was kept in separating funnel for 24 h without further stirring, to ensure proper sedimentation. The obtained methyl ester was washed with distilled water. Demeoisturization of the obtained methyl ester was carried out at 100 °C.

Eq. (2) was used to estimate % methyl ester yield.

\[
\text{% Methyl ester yield} = \frac{\text{Mass of methyl ester produced (g)}}{\text{Mass of oil sample used (g)}} \times 100
\]  

(2)

The obtained methyl ester sample is modified *Irvingia gabonensis* kernel oil (MIGKO⁴). The scheme is presented in Figure 1. A similar reaction scheme had earlier been presented in our previous publication (Agu et al., 2019).

2.4. Epoxidation experiments

2.4.1. Epoxidation reaction

The epoxidation reaction was carried out according to the method described by Agu et al. (2019). 100 g of the MIGKO⁴ sample was measured into a flask and heated to 70 °C, prior to the addition of peracetic acid (PAA). The PAA was obtained by mixing 14 wt % of acetic acid and 2.5 % of H₂SO₄. Thereafter, hydrogen peroxide (16 wt %) was added. After the introduction, the reaction proceeded with constant stirring (1200 RPM) for 7 h. Afterwards, 5 % NaHCO₃ was used to purify the sample, followed by saturated NaCl addition, so as to get the epoxide. Finally, anhydrous magnesium sulphate was used to dry the product, while a rotary evaporator was used for solvent removal.

Figure 2 shows the MIGKO⁴ epoxidation reaction for epoxide [epoxy methyl ester of *Irvingia gabonensis* kernel oil (EMIGKO⁰)] production. A similar reaction scheme had earlier been presented in our previous publication, Agu et al. (2019).

2.4.2. Esterification ring opening reaction

The ring-opening reaction was carried out according to the method described in Agu et al. (2019). EMIGKO⁰ (20 g) was measured and then added to 10 ml of ethyl acetate with constant stirring. 4 g acid anhydride was then introduced, followed by boron trifluoro diethyletherate (1 ml). The reaction proceeded with constant stirring (1200 RPM) for 7 h. Thereafter, 5 % NaHCO₃ was used to purify the obtained branched methyl ester sample, followed by the addition of 10 % NaCl solution. Subsequently, the branched methyl ester sample was obtained, after NaCl addition. Finally, anhydrous magnesium sulphate was used to dry the product, while rotary evaporator was applied for solvent removal. Distillation method was used to remove the unreacted anhydrides at reduced pressure and 80 °C.

Figure 3 show the esterification ring-opening of the EMIGKO⁰ (epoxide) reaction scheme for the production of modified/epoxidized *Irvingia gabonensis* kernel oil methyl ester (MIGKO²) sample. Similar reaction scheme had earlier been presented in our previous publication by Agu et al. (2019).

2.5. Physicochemical characterization of IGKO and MIGKO² samples

Using AOAC approved techniques (1990), acidity/acid value (AOAC 969.17), iodine value (AOAC 993.20), density (AOAC 985.19) and moisture content (AOAC 926.12) of the samples were estimated. Similarly, ASTM D445 (2011) was used for viscosity determination, while the breakdown voltage (dielectric strength) was analyzed following the IEC 60156 (2003) standard. Furthermore, flash and pour points were determined using ASTM D93 (2012) and ASTM D97 (2012) standard procedures respectively. These properties were measured in triplicates while obtained average values were recorded.

2.6. Epoxidation kinetics

The in situ epoxidation reaction generally takes place in two steps: (i) formation of peroxy acid and (ii) reaction of peroxy acid with the unsaturated fatty acids of the methyl ester (MIGKO⁴). Eqs. (3) and (4) are...
the equations for the formation of peroxy acid and reac
tion of peroxy acid with the unsaturated fatty acids of the methyl esters respectively.

\[
\text{RCOOH} + \text{H}_2\text{O} \rightarrow \text{RCOOH} + \text{H}_2\text{O}
\]  
(3)

\[
\text{unsaturated fatty acid} \rightarrow \text{peroxy acid} \rightarrow \text{epoxide} \rightarrow \text{acid}
\]  
(4)

Assuming that the rate determining step for the epoxidation reaction is peroxy acid formation, while peracid concentration is assumed to be unchanged during the entire reaction process, the epoxidation rate is then expressed as shown in Eq. (5) (Arumugam et al., 2014; Gan et al., 1995).

\[
\frac{d[\text{EP}]}{dt} = k([\text{H}_2\text{O}_2]_0 - [\text{EP}]) \cdot \text{[RCOOH]}_0
\]  
(5)

Where subscript 0 denotes initial concentrations, \( k \) denotes the rate constant of the epoxidation reaction and EP denotes epoxides.

Hence,

\[
\ln \left( \frac{[\text{H}_2\text{O}_2]_0 - [\text{EP}]}{[\text{RCOOH}]}_0 \right) = \frac{k}{t} + \ln [\text{H}_2\text{O}_2]_0
\]  
(6)

According to Eq. (6), plot of \( \ln \left( \frac{[\text{H}_2\text{O}_2]_0 - [\text{EP}]}{[\text{RCOOH}]}_0 \right) \) vs. time should yield straight lines for those reactions with insignificant oxirane degradation. However, reactions with substantial ring opening were seen to deviate from linearity. As such, their respective rate constants were obtained using the initial slopes.

### 2.7. Activation energy calculation

The energy level of the molecules to initiate a chemical reaction is generally represented as the activation energy.

For a second order model, rate constants increase with temperature and may be described by the Arrhenius law in Eq. (7).

\[
K = A \exp \left( \frac{-E_a}{RT} \right)
\]  
(7)

Where: \( K \) = the extraction rate constant (L g\(^{-1}\) min\(^{-1}\))  
\( A \) = the temperature independent factor (L g\(^{-1}\) min\(^{-1}\))  
\( E_a \) = the activation energy (J mol\(^{-1}\))  
\( R \) = the gas constant (8.314 J mol\(^{-1}\) K\(^{-1}\))  
\( T \) = the absolute suspension temperature (K)

Eq. (8) is the Linearized form of Eq. (7). From the plot of ln (K) against \( 1/T \) plot derived from Eq. (8), the values of the Activation energy (\( E_a \)) and the temperature independent factor (\( A \)) would be obtained from the slope and the intercept, respectively.

\[
\ln(k) = \ln (A) + \left( \frac{-E_a}{R} \right) \frac{1}{T}
\]  
(8)

### 2.8. Thermodynamics studies of epoxidation reaction

In the epoxidation of *Irvingia gabonensis* kernel oil methyl esters (MIGKO\(^3\)), Eq. (9) was used to estimate the average enthalpy of activation \( \Delta H \) of the reaction (Okieimen et al., 2002).

\[
\Delta H = E_a - RT
\]  
(9)

Where \( E_a \) is the activation energy calculated from the Arrhenius plot, \( T \) is temperature, and \( R \) is the universal gas constant.

Similarly, the average entropy of activation, \( \Delta S \), as well as the Gibbs free energy of activation, \( AG \), were obtained using equations, 10 and 11, respectively (Okieimen et al., 2002).

\[
k = \frac{RT}{Nh} \cdot e^{\Delta S/k} \cdot e^{-E_a/RT}
\]  
(10)

\[
\Delta G = -RT \ln K
\]  
(11)

Where \( k \), \( N \) and \( h \) are the rate constant, Avogadro (6.02 \times 10\(^{23}\) mol\(^{-1}\)) constant and Planck (6.63 \times 10\(^{-34}\) J) constant, respectively.

### 2.9. Thermogravimetric Analyses (TGA) of Irvingia gabonensis kernel

The thermal stability/behavior of IGK was investigated using a Mettler-Toledo TGA/SDT A851e thermogravimetric analyzer under nitrogen atmosphere. A total of 8.96 mg of the *Irvingia gabonensis* IGK sample was placed in an aluminium vessel and covered with a pierceable cap. Prior to the measurement of the IGK sample, the cap was perforated and the sample was placed in a nitrogen N\(_2\) atmosphere. A total of 8.96 mg of the IGK sample was placed in an aluminium vessel and covered with a pierceable cap. Prior to the measurement of the IGK sample, the cap was perforated and the sample was heated from 24.52 °C to 800 °C at a heating rate of 3–4 °C/min. Nitrogen N\(_2\) was used as an inert atmospheric gas at 100 ml min\(^{-1}\) flow rate. The different weight loss phases and loss-rate temperatures for moisture and other combustible materials loss steps, alongside the overall percentage weight loss were determined using STARe Thermogravimetric Analysis Software.

### 3. Results and discussions

#### 3.1. Oil yield of Irvingia gabonensis kernel

As already reported in our earlier publications, the oil yield of *Irvingia gabonensis* kernel (IGK) was 68.80 % (by dry mass basis) (Agú et al., 2020). This oil yield obtained, differ from other IGK oil yields reported elsewhere, 67.33 % (Ekpe et al., 2018), 73.83 % (Matos et al., 2009) and 69.76 % (Zou et al., 2015) elsewhere. Variations in oil yields were associated with the geographical location, solvents of extraction, varieties of Irvingia gabonensis kernel, as well as seeds/kernels harvest period. However, when compared to other oil seeds, the oil yield obtained for IGK was significantly higher than those of *Terminalia catappa* L. kernel (60.45 %), Soybean (20.4 %) Colocynthis vulgaris Shred Seeds and Cottonseed (24.82 %), as reported by Agú et al. (2019), Nikolić et al. (2009), Agú et al. (2018) and Sharif et al. (2019). Hence, the economic viability of IGK oil has been established, as well as its high oil yield; unlike seeds like *Chrysophyllum albidum* with lower oil yield.
3.2. Physicochemical properties of *Irvingia gabonensis* kernel oil (IGKO) and the epoxidized methyl ester (MIGKO*$_e$)

The properties of *Irvingia gabonensis* kernel oil (IGKO) and the epoxidized methyl ester of *Irvingia gabonensis* kernel I oil (MIGKO*$_e$) samples are presented in Table 1.

3.2.1. Dielectric strength

Dielectric strength is the maximum voltage needed to cause dielectric breakdown of the material. It is also expressed as the maximum electrical strength that a material, like oil can constantly withstand without failure of its properties. Its importance cannot be overemphasized because it is vital in maintaining reliable operation of power transformers, in order to avert transformer failure (Agu et al., 2019). The dielectric strength (DS) (breakdown voltage) values for IGKO and MIGKO*$_e$ were 25.83 ± 0.001 kV and 39.78 ± 0.001 kV, respectively (see Table 1). With reference to the values of the DS of IGKO and MIGKO*$_e$, it was noticed that the breakdown voltage changed (increased) after the modification of the IGKO; with MIGKO*$_e$ having higher DS value. Hence, the reason for the 39.78 ± 0.001 kV value, obtained for MIGKO*$_e$ sample. This increase in the DS, resulting from the modification of the IGKO, was associated to esterification and transesterification processes used for methyl ester production; as well as the choice of the methyl ester over the oil for the epoxidation-esterification modification process (Menkiti et al., 2017b). The breakdown voltage of IGKO, obtained in this work is comparable, though slightly higher than those of crude palm oil (17–23 kV) (Azis et al., 2014) and palm kernel oil (25 kV) (Usman et al., 2012). Although the breakdown voltage of IGKO was slightly less, compared to 30.61 kV for *Terminalia catappa* L. kernel (Agu, 2019); it was significantly lower, compared to 60 kV reported for coconut oil by Abyseundara et al. (2001). These variations in DS of IGKO and other vegetable oils in literature were linked to differences in triglycerides of various vegetable oils; as a result of their diverse polar natures (Agu et al., 2019; Shah and Tahir, 2011).

Furthermore, for the epoxidized methyl ester of *Irvingia gabonensis* kernel oil (MIGKO*$_e$), its dielectric strength (DS) was found to be lower, compared to epoxy methyl esters of palm kernel oil (42.58 kV) (Abdelmalik et al., 2011) and *Terminalia catappa* L. kernel oil (50.05 kV) (Agu et al., 2019). However, it was higher, compared to that of purified calabash seed oil transformer fluid (24 kV) (Oyelaran et al., 2020). Just like in the base vegetable oils, the difference in the DS of MIGKO*$_e$ and other epoxy methyl esters of various vegetable oil in literature, was attributed to their respective diverse triglycerides polar natures (Agu et al., 2019). Also, this difference in the DS of MIGKO*$_e$ and other works in literature, could be associated to diverse modification process routes, and reagents used (Agu et al., 2019).

3.2.2. Moisture content

The primary cause of poor vegetable oil quality, when compared to mineral oils is high moisture. This is attributed to its potential to increase tendency for oil to conduct electricity, as against its primary role of insulation especially for oils like transformer fluids (Menkiti et al., 2017b). It is therefore important to state that high moisture content in oils, results in reduction in DS of oil, as well as its insulation/cooling properties. As noted in Table 1, moisture contents for IGKO and MIGKO*$_e$ were 3.75 ± 0.001 mg/kg and 1.01 ± 0.001 mg/kg, respectively. Lower moisture value was noticed for MIGKO*$_e$, when compared to IGKO. This decrease in the moisture content of MIGKO*$_e$ sample was associated with catalytic reactions of esterification, transesterification, and the epoxidation-esterification modification processes. This is because oil modification, as seen in this work, lowered the moisture content of the final product (MIGKO*$_e$) (Menkiti et al., 2017b). Furthermore, the moisture content of IGKO was found to be higher, compared to those of *Terminalia catappa* L. kernel oil (2.1 mg/kg) (Agu et al., 2019) and soya bean oil (2.0 mg/kg) (Usman et al., 2012). However, moisture content variations were attributed to the method used to extract oil from the seeds (Ikye et al., 2013). This is because it has been established that the moisture content of solvent (method) extracted oils, is less than the traditional cold maturation extracted oils (Agu et al., 2018). Nevertheless, for the modified oil sample obtained by epoxidation-esterification of *Irvingia gabonensis* kernel oil methyl ester (MIGKO*$_e$), its moisture content was found to be lower than the 1.9 ± 0.001 mg/kg, reported for bio-transformer fluids obtained from palm oil; but the same as the 1.0 ± 0.001 mg/kg obtained from coconut (Aimi and Hussin, 2014). However, the moisture content of MIGKO*$_e$ in this work was higher, compared to 0.18 ± 0.03 mg/kg for epoxidized waste cooking oil methyl ester (Paul et al., 2021) and 0.65 ± 0.001 mg/kg for epoxidized *Terminalia catappa* L. kernel oil methyl ester (Agu et al., 2019). As earlier reported for vegetable oils, the differences in moisture contents of MIGKO*$_e$ sample, and those of bio-transformer fluids produced from coconut, *Terminalia catappa* L. kernel and palm oils, were linked to extraction and modification methods adopted (Agu et al., 2019).

3.2.3. Flash and pour points

As it relates to transformer fluid, flash point is the temperature during which constituents (especially hydrocarbons) present in the oil begins to evaporate, leading to flash when in contact with source of light. The implication is that the flash point must be kept above temperature of 140 °C. Conversely, the least temperature at which oil begins to flow at specific standard condition is referred to as the pour point. In temperate regions with icy/snow climate particularly during winter, pour point of transformer oil is a vital property of valuable consideration. This is due to the fact that if oil temperature drops below the pour point, transformer oil functionality of convection flow stops, resulting in obstruction of the transformer cooling (Agu, 2019). As presented in Table 1, with the epoxidation-esterification modification process, flash point of *Irvingia gabonensis* kernel oil (IGKO) sample increased, while pour point decreased. Hence, there was increase in flash point, and corresponding decrease in pour point of modified oil (MIGKO*$_e$), after the modification process. It is vital to state that the decline in pour point of IGKO resulting from the oil modification was linked to decline in the unsaturation nature

| Property                  | Unit | TO* | IGKO* | MIGKO* | Standard method |
|---------------------------|------|-----|-------|--------|----------------|
| Electric strength         | kV   | 40-60| 25.83 ± 0.001 | 39.78 ± 0.001 | IEC 60156 |
| Moisture content          | mg/kg| <20 | 3.75 ± 0.001 | 1.01 ± 0.001 | AOAC 926.12 |
| Pour point                | °C   | -48 | 17 ± 0.5 | 9 ± 0.5 | ASTM D97 |
| Flash point               | °C   | 152 | 285 ± 0.5 | 298 ± 0.5 | ASTM D93 |
| Density, 20 °C            | g/cm³| 870 | 900 ± 0.5 | 840 ± 0.5 | AOAC 985.19 |
| Viscosity, 40 °C          | mm²/s| 10  | 19.37 ± 0.001 | 13.84 ± 0.001 | ASTM D445 |
| Acidity/Acid value        | mg KOH/g oil | <0.01 | 5.18 ± 0.001 | 1.351 ± 0.001 | AOAC 969.17 |

* conventional mineral transformer oil.

* *Irvingia gabonensis* kernels oil.

* Modified* *Irvingia gabonensis* kernels oil obtained by epoxidation-esterification of the methyl ester.

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Table 1. Physicochemical properties of Modified *Irvingia gabonensis* kernel oil samples...
of the oil sample, emanating from the modification processes (Adekunle et al., 2016). From the Table 1, the flash points of IGKO and MIGKOe samples were 285.0 ± 0.5 °C and 298.0 ± 0.5 °C, respectively. Furthermore, the flash point of IGKO was found to be higher than those of coconut oil (225 °C), Terminalia catappa L. kernel oil (260 °C) and Jatropha oil (146 °C), as reported by Aimi and Hussin (2014), Agu et al. (2019) and Aliyu and Tijjani et al. (2017), respectively. For Irvingia gabonensis kernel oil (IGKO) samples reported by other researchers, Bello et al. (2011) reported that a flash point value of 300 °C. Similarly, the flash point (298.0 ± 0.5 °C) of MIGKOe, was found to be higher, compared to those of palm kernel alkyl ester with 148 °C (Abdelmalik, 2014) and epoxidized Terminalia catappa L. kernel oil methyl ester 280 °C (Agu et al., 2019). Also, the flash point of MIGKOe was higher than mineral transformer fluid with 152 °C (Menkiti et al., 2017b). On the other hand, Table 1 shows that pour point values of IGKO and MIGKOe samples were 17 ± 0.5 °C and 9 ± 0.5 °C, respectively. These values were higher, compared to the -48 °C value, for conventional transformer fluid. Higher pour points of IGKO and MIGKOe were due to unsaturated fatty acid ester presence in oil, as well as its leftover in the MIGKOe sample; despite the ester enrichment processes (Agu et al., 2019). IGKO pour value was found to be less than 28 °C, reported for Irvingia gabonensis kernel oil by Bello et al. (2011). This value was also found to be greater than the 2 ± 0.15 °C, 3 ± 0.5 °C and 1.15 ± 0.10 °C, reported for Moringa seed, Terminalia catappa L. kernel and castor seed oils, by Aliyu and Tijjani (2017), Agu et al. (2019) and Aliyu and Tijjani (2017), respectively. Pour point of MIGKOe was lower than -8 ± 0.5 °C, 2.40 ± 0.03 °C, 4 °C and 8 °C values, reported by Agu et al. (2019), Paul et al. (2021), Oyelaran et al. (2020) and Borugadda and Goud (2014), for epoxidized Terminalia catappa L. kernel oil methyl ester, epoxidized waste cooking oil methyl ester, purified calabash seed oil transformer fluid and epoxidized castor oil methyl ester, respectively. The difference in pour point of MIGKOe and other epoxidized methyl ester samples and the purified calabash seed oil sample, was linked to the adopted extraction and modification methods (Demirbas, 2009). However, using pour point depressants, there is possibility of improvement in Pour point of MIGKOe (Agu, 2019).

3.2.4. Density

Density is defined as the substance mass per unit volume. The density of oil is very important as it helps/guides in pump designs, since most pump systems are designed to pump a fluid of a specific density. With respect to transformer fluid, it helps to ensure that proper cooling takes place in the system during operation (Agu, 2019). From Table 1, the densities of IGKO and MIGKOe samples were 900 ± 0.5 g/cm³ and 840 ± 0.5 g/cm³, respectively. The density of MIGKOe was seen to be lower, compared to that of IGKO (Table 1). This reduction in IGKO density value after modification, which is evident in the obtained value for MIGKOe, was caused by the difference in density of IGKO, compared to other vegetable oil seeds/nuts (Menkiti et al., 2017b). With reference to other results in the literature, it was found that density of IGKO was lower, compared to 930 g/cm³ for Irvingia gabonensis kernel oil (Bello et al., 2011). The difference in density of IGKO in this study, compared to other IGKO reported in the literature, could be associated with diverse morphologies of the seeds/kernels, and methods of extracting the oils (Ilyaka et al., 2013). However, the density of IGKO (900 ± 0.5 g/cm³) in this study was observed to be higher, compared to those of Jatropha (720 ± 0.50 g/cm³), Moringa (520 ± 0.40 g/cm³), Castor (740 ± 0.30 g/cm³) and cotton (850 ± 0.40 g/cm³) oils (Aliyu and Tijjani, 2017). Also, the difference in density of IGKO, compared to other vegetable oil seeds/nuts could be associated with diverse morphologies of the seeds, as well as the method of extracting the oils (Agu et al., 2019). This is associated with the fact that better purity levels, hence lower densities, are obtained when solvent extraction using soxhlet extractor is used, unlike the adoption of cold maturation method (Agu et al., 2018). Furthermore, the density (840 ± 0.5 g/cm³) of the modified oil sample obtained by epoxidation-esterification of Irvingia gabonensis kernel oil methyl ester (MIGKOe) was found to be lower than the 850 ± 0.5 g/cm³ and 956 g/cm³ values, which were reported for purified calabash seed oil used as bio-transformer fluids and epoxidized castor oil fatty acid methyl ester, by Oyelaran et al. (2020) and Borugadda and Goud (2014), respectively. On the other hand, the density (840 ± 0.5 g/cm³) of MIGKOe sample was higher than 773.8 ± 3.71 g/cm³, reported for epoxidized waste cooking oil methyl ester, by Paul et al. (2021). Also, as earlier stated for vegetable oils, this difference in the density of the MIGKOe sample and those of other bio-transformer fluids, was linked to extraction methods adopted; and the modification routes used for the processes (Agu et al., 2019). From Table 1, it is worthy of mention that the density of the final product obtained by epoxidation-esterification of Irvingia gabonensis kernel oil methyl ester (MIGKOe), was close to the standard stipulated for the density (870 g/cm³) of conventional mineral transformer oil.

3.2.5. Viscosity

It is evident from Table 1 that the viscosity of IGKO, significantly decreased after the modification process, giving rise to the final product (MIGKOe sample), with much lower viscosity value. Viscosity of IGKO and MIGKOe samples were 19.379 ± 0.001 mm²/s and 13.84 ± 0.001 mm²/s, respectively (See Table 1). This was due to the modification reactions processes routes adopted, which improve product (MIGKOe sample) purity, viscosity, as well as its thermo-oxidative stability state (Agu et al., 2019; Abdelmalik et al., 2011). As observed in Table 1, the viscosity of the obtained IGKO (19.37 ± 0.001 mm²/s) was found to be lower than 45 mm²/s reported for Irvingia gabonensis kernel oil in literature, by Bello et al. (2011). As earlier inferred, the differences in viscosities of IGKO in this study, and other IGKO reported in literature was likened to the extraction method used, in addition to the differences in their respective species (Agu et al., 2019). IGKO viscosity (19.37 ± 0.001 mm²/s) in this work was found to be higher than the viscosity of other seeds/nuts oils like, Jatropha (16.50 ± 0.12 mm²/s), Moringa (11.80 ± 0.02 mm²/s), and Castor (18.10 ± 0.12 mm²/s) (Aliyu and Tijjani, 2017). However, the viscosity of IGKO in this work was found to be lower, compared to those of Sunflower oil (34.5 mm²/s) (Rouabeh et al., 2019), Coconut oil (29 mm²/s) (Aimi and Hussin, 2014), Cotton oil (29.40 ± 0.12 mm²/s) (Aliyu and Tijjani, 2017), Palm oil (29.2 mm²/s) (Aimi and Hussin, 2014), Terminalia catappa L. kernel oil (20.29 ± 0.001 mm²/s) (Agu et al., 2019) and Olive oil (110.3 mm²/s) (Rouabeh et al., 2019). The differences in viscosity of these other seeds/nuts reported in the literature and that of IGKO in this study, was attributed to the difference in the morphologies of the seeds/nuts, in addition to the extraction method chosen for extraction purposes (Agu et al., 2019). Furthermore, the viscosity (13.84 ± 0.001 mm²/s) of the modified oil sample (MIGKOe) was found to be higher than 8.8 mm²/s (Oyelaran et al., 2020), and 12.15 ± 0.07 mm²/s (Paul et al., 2021), reported for purified calabash seed oil used as bio-transformer fluids and epoxidized waste cooking oil methyl ester respectively. However, its viscosity was lower, compared to the 170.85 mm²/s, 35.81 mm²/s and 55.13 mm²/s, reported for epoxidized soybean oil, epoxidized castor oil fatty acid methyl esteran epoxidized trimethylolpropane (TMP) ester, by Adhavarye and Erhan (2002), Borugadda and Goud (2014) and Naidir et al. (2012), respectively. However, the 13.84 ± 0.001 mm²/s value obtained for MIGKOe sample, was higher than 3.2 mm²/s, reported for IGKO biodiesel, by Bello et al. (2011). These observable differences, was likened to the variations in the structural formations of the molecules of the triglyceride in the respective base oils, together with the differences in the modification methods/process routes (Agu et al., 2019).

3.2.6. Acid value

As observed in Table 1, the acid values of IGKO and MIGKOe samples were 5.18 ± 0.001 mg KOH/g oil and 1.351 ± 0.001 mg KOH/g oil,
respectively. From Table 1, it was observed that MIGKO\(t\) sample had lower acid value, when compared to IGKO. This decrease in acid value as evident in the MIGKO\(t\) sample could be attributed to the fact that the prevalent free fatty acids in IGKO were significantly removed after the modification process (Agu et al., 2019). Acid value of IGKO was higher, compared to other IGKO reported in literature by 4.67 mg KOH/g oil (Zou et al., 2013) and 1.2 mg KOH/g oil (Bello et al., 2011). However, IGKO acid value was less, compared to 9.40 mg KOH/g oil, as reported for Irvingia gabonensis kernel oil, by Etong et al. (2014). As earlier inferred in other earlier discussed properties, the differences in acid values of IGKO in this study, and other IGKO reported in literature was due to the extraction method used, in addition to the differences in their respective species (Agu et al., 2019). Nevertheless, acid value of IGKO in this work was higher, compared to those of Jatropha (3.45 ± 0.12 mg KOH/g oil), Moringa (4.47 ± 0.12 mg KOH/g oil), Castor (2.12 ± 0.08 mg KOH/g oil), and Cotton (3.72 ± 0.51 mg KOH/g oil) oils, reported by Aliyu and Tijjani (2017). These differences in the acid values of these other seeds/nuts reported in literature and that of IGKO in this study, was attributed to the difference in the morphologies of the seeds/nuts, in addition to extraction method used, as oils obtained by solvent extraction method, have higher acid value, compared to those obtained by cold maturation. Also, in solvent extraction method, solvent type significantly influences acid value of obtained oil (Agu et al., 2019). Furthermore, for epoxidized-esterified Irvingia gabonensis kernel oil methyl ester (MIGKO\(t\)), its acid value (1.351 ± 0.001 mg KOH/g oil) was found to be higher than 0.142 mg KOH/g oil, 1.08 mg KOH/g oil and 0.59 mg KOH/g oil, reported for purified calabash seed oil used as bio-transformer fluids (Oyelaran et al., 2020), epoxidized castor oil fatty acid methyl ester (Borugadda and Goud, 2014) and epoxidized methyl ester of Parkia biglobosa seed oil (Ikhuoria et al., 2007), respectively. Also, the acid value of MIGKO\(t\) (1.351 ± 0.001 mg KOH/g oil) was still higher, compared to 0.20 ± 0.04 mg KOH/g oil for epoxidized waste cooking oil methyl ester (Paul et al., 2021) and 0.09 mg KOH/g oil for epoxidized soybean oil (Adhvarye and Erhan, 2002). Furthermore, acid value of MIGKO\(t\) was also greater, compared to <0.01 mg KOH/g oil recommended for mineral transformer oil by IEC standard. As earlier reported for vegetable oils, the difference in acid value of MIGKO\(t\) and that of purified calabash seed oil used as bio-transformer fluids was associated with adopted modification procedure (Agu et al., 2019). However, it is important to state that high acid value in transformer fluid, results in its absorption by the paper insulating materials in the transformer; hence resulting in increased degradation of the entire transformer insulation system (Kouassi et al., 2018).

### 3.3. Kinetics of the epoxidation of Irvingia gabonensis kernel oil methyl ester (MIGKO\(t\))

Figure 4 shows the plots for the in situ epoxidation of Irvingia gabonensis kernel oil methyl ester (MIGKO\(t\)) at different temperatures. According to Equ (6), the plot of In \([\text{H}_2\text{O}_2(\text{EP})]\) against time results in straight lines, specifically for reactions that had insignificant degradation of oxirane (see Figure 4). In this study, deviations from linearity were observed for the reactions (at 55, 65 and 75 °C) with substantial ring opening (Figure 4). This was attributed to the epoxide degradation (Ikhuoria et al., 2007). In cases like this, epoxidation reaction rate constants were gotten using initial slopes.

The values of rate constants, gotten from linear portion of the plots for the studied sample (MIGKO\(t\)) at different temperatures are presented in Table 2. Rate constants gotten for MIGKO\(t\) sample are of the order of 10\(^{-5}\) Lmol\(^{-1}\)s\(^{-1}\) (Table 2). The rate constants obtained for the sample, were less than the 10\(^{-4}\) Lmol\(^{-1}\)s\(^{-1}\), as reported by Cai and Wang (2011) for the epoxidation of unsaturated fatty acid methyl ester. Although, this obtained rate constant for the epoxidation of the MIGKO\(t\) sample was of the same order of 10\(^{-5}\) Lmol\(^{-1}\)s\(^{-1}\), reported for oleic acid derived from palm kernel oil (PKO), by Jalil et al. (2018). However, the obtained rate constants of the studied sample was greater, compared to 10\(^{-6}\) Lmol\(^{-1}\)s\(^{-1}\) for epoxidations of rubber seed oil (Okieimen et al., 2002), palm olein methyl ester (Gan et al., 1992) and mahuwa oil (Goud et al., 2006). Similarly, the obtained rate constants in this work were also greater than the 10\(^{-7}\) Lmol\(^{-1}\)s\(^{-1}\), as reported by Ikhuoria et al. (2007) for the kinetics of epoxidation of methyl esters of Parkia Biglobosa seed oil. This difference in rate constants for epoxidation of MIGKO\(t\) sample, compared to other methyl esters and vegetable oils, was due to differences in hydrogen bonding present in molecular chains of these samples, as well as interaction between these molecules (Saalah et al., 2017; Bayo et al., 2001).

Furthermore, it is important to state that the values of rate constants for the studied sample were temperature dependent. Hence, elevation of reaction temperature increases the rate constant. The reason behind this is that elevated temperature favors formation of peracetic acid, resulting in both accelerated reaction and increased oxirane conversion, in addition to enhanced rate of oxirane ring opening (Cai and Wang, 2011). For instance, there was over sevenfold increase in the rate constant for the epoxidation of MIGKO\(t\), when temperature was increased from 35 to 75 °C. The results obtained in this study were slightly different from that obtained at temperature of 45–75 °C by Cai et al. (2008) with a fourfold increase in the rate constant value for the epoxidation of soybean oil. Also, Ikhuoria et al. (2007) reported a threefold increase in the rate constant value for the epoxidation of Parkia Biglobosa seed oil when the reaction temperature was increased from 50 to 70 °C.

### 3.4. Thermodynamics of the epoxidation of Irvingia gabonensis kernel oil methyl ester

The values of activation energy, calculated from the slope of the linear dependence of InK verses 1/T (K\(^{-1}\)) (Eq. 9) [see Figure 5] for the epoxidation of the Irvingia gabonensis kernel oil methyl esters (MIGKO\(t\)) are presented in Table 3. From the table, the activation energy for MIGKO\(t\) sample is 46.02 kJ/mol. The value of the activation energy obtained for the studied sample was significantly lower, compared to those reported by Gan et al. (1992) (63.20 kJ/mol), Goud et al. (2006) (60.60 kJ/mol),

![Figure 4. Plot of In \([\text{H}_2\text{O}_2(\text{EP})]\) vs. time for the epoxidation of MIGKO\(t\) by peracetic acid.](image-url)
Furthermore, the activation energy (46.02 kJ/mol), obtained for the epoxidations of unsaturated fatty acid methyl esters and soybean oil, respectively. However, the thermodynamic parameters, calculated from Eqs. (9), (10), and (11), respectively, are also presented in Table 3. From Table 3, it was observed that the value of \( \Delta G \) for the epoxidation of MIGKO \( -1 \) (Table 3), was significantly lower than those reported by Jalil et al. (2018) (47.82 kJ mol \(-1\)), Gan et al. (1992) (146.0 kJ mol \(-1\)) and Goud et al. (2006) (146.0 kJ mol \(-1\) \( \times 1 \)) for the epoxidations of soybean oil, palm olein methyl ester and mahua oil, respectively. However, in the works of Ikhuoria et al. (2007) and Okieimen et al. (2002), they reported that the \( \Delta S \) values for epoxidation of Parkia Biglobosa seed oil and rubber seed oil were 3550 J mol \(-1\) \( \times 1 \) and 213.6 J mol \(-1\) \( \times 1 \), respectively. These values were lower than the \( \Delta S \) values for MIGKO \( -1 \) epoxidation.

The value of \( \Delta G \) for the epoxidation of MIGKO \( -1 \) sample was found to be in the range of 94.74–101.42 kJ mol \(-1\) (Table 3). These positive values of \( \Delta G \) indicate the non-spontaneous nature of the reaction (Cai and Wang, 2011). From Table 3, it was observed that the \( \Delta G \) values for the MIGKO \( -1 \) sample studied, increased with temperature. Thus, indicating that the Gibb’s free energy of a substance is temperature dependent (Lin et al., 2008; Cai and Wang, 2011). These values were similar in range with those reported by Cai et al. (2008) (110.01 kJ mol \(-1\)), Lin et al. (2008) (111.39 kJ mol \(-1\)), Cai and Wang (2011) (112.1 kJ mol \(-1\)), Goud et al. (2006) (106.3 kJ mol \(-1\)) and Okieimen et al. (2002) (127.9 kJ mol \(-1\)) for the epoxidation of sunflower oil, soybean oil, unsaturated fatty acid methyl ester, mahua and rubber seed oil, respectively. However, the \( \Delta G \) range of values (94.74–101.42 kJ mol \(-1\)) for the epoxidation of MIGKO \( -1 \) sample was higher, compared to those for the epoxidation of soybean oil (25.98 kJ mol \(-1\)), sunflower oil (24.98 kJ mol \(-1\)) and rapeseed oil (22.78 kJ mol \(-1\)) (Jalil et al., 2018).

3.5. Thermogravimetric Analyses (TGA) of Irvingia gabonensis kernel (IGK)

TGA was used to investigate the thermal decomposition and thermal stability behaviors of Irvingia gabonensis kernel (IGK) at heating rate of 3–4 °C/min. The decomposition of the IGK sample is shown in Figure 6. The thermograms show three weight loss phases for the IGK sample. Similar results have been reported by a number of researchers. Sugumaran et al. (2012) reported that during the thermal decomposition of banana empty fruit bunch and Delonix regia fruit pods, four and three weight losses were respectively observed. Similarly, Vichaphund et al. (2014) and Lotus et al. (2011) both reported three weight loss phases for the thermal decomposition of physic nut and as-spun fibers, respectively.

From Figure 6, the first phase for the IGK sample with range of 24.52–71.05 °C corresponded to the loss of moisture. This moisture loss represents 3.49 % initial weight losses in IGK. Similar results of moisture losses can be found in the works on the thermal decompositions of Chrysophyllum albidum (Sokoto et al., 2016), Jatropha curcas L. (Vichaphund et al., 2014), and banana empty fruit bunch (Sugumaran et al., 2012).

The second phase weight loss for the IGK (71.05–244.14 °C), could be attributed to (correspond to) the loss of residual moisture, oils and other volatile materials. This decomposed material at this phase represents 25.43 %, losses in the initial mass of the IGK sample. Furthermore, the third phase weight loss for the IGK (244.14–284.33 °C), corresponds to 60 % losses in the initial mass of the sample. At this phase, there was major decomposition of organic compounds like starches, fibers, cellulose biomass for IGK, as evident in its huge weight losses. Synonymous results have been reported in the literature for the thermal decompositions of Terminalia catappa L., Chrysophyllum albidum and natural fiber by Dos Santos et al. (2016), Sokoto et al. (2016) and Yao et al. (2008), respectively.
The thermal decomposition of IGK sample occurred within the temperature range of 244.14–384.33 °C, with maximum decomposition of 60%. As could be seen in the curve in Figure 6, the complete pyrolysis of IGK sample occurred at 384.33 °C. This third phase thermal decomposition temperatures of the studied IGK sample is comparable to the degradation temperature range of 200–500 °C for Chrysophyllum albidum cake (Sokoto et al., 2016), safflower seed (Onay, 2007) and Cherry seed (Duman et al., 2011). However, this third phase decomposition temperature of the IGK sample was lower than 400–700 °C for rapeseeds cake (Onay and Koçkar, 2004).

Finally, beyond the decomposition temperature 384.33 °C, for the IGK sample, there was no significant change in the mass of the sample with further temperature increase. Hence, indicating the high thermal stability of the samples. Therefore, the residue left after the volatilization of combustible substances was 11.18 % of the initial mass of the IGK sample, used for the analyses. Practically, the high decomposition temperature of the sample is an indication of its high thermal stability.

4. Conclusion

Based on experimental conditions, after the modification of IGKO, MIGKOe had acceptable properties that were comparable to mineral transformer oil properties. Hence, the properties MIGKOe significantly conformed to the stipulated IEC standard for transformer fluid. At extraction conditions of 55 °C, 150 min and 0.5 mm, 68.80 % was the maximum oil yield, extracted from Irvingia gabonensis kernel (IGK). Rate constant K and activation energy Ea for the epoxidation process was found to be of the order 10^-5 Lmol^-1s^-1 and 46.02 kJ/mol, respectively. The ΔG, ΔH, and ΔS values obtained for the epoxidation reaction indicates non-spontaneous, endothermic, and endergonic nature of the process. Finally, high decomposition temperature of the Irvingia gabonensis kernel (IGK) samples, as obtained from the Thermogravimetric Analyses (TGA) results, is an indication of the high thermal stability of the sample. In other words, IGKO and MIGKOe samples could be used in transformers as dielectric fluids.

Declarations

Author contribution statement

C.M. Agu: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.
A.C. Agulanna, C.H. Kadurumba: Performed the experiments.
P.C. Nnaji, E.L Udokporo: Contributed reagents, materials, analysis tools or data.
M.C. Menkiti: Analyzed and interpreted the data.

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Additional information

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