Rubber seeds from a Nigerian hybrid rubber tree were dried in an oven at 110°C for 2 h to 7 wt% moisture content and cold-screw pressed at 30°C to obtain an oil yield of 28 wt%. Chemical compositional, nuclear magnetic resonance (NMR) and Fourier transform infra-red spectroscopy (FTIR) analyses were conducted on the oil sample extracts. Chemical analysis of the oil indicated iodine value of 136.07 g I₂/100 g oil, peroxide value of 9.45 O₂/kg oil, saponification value of 189.5 mg KOH/g oil, and confirms its suitability for applications, such as biodiesel production, oleochemicals synthesis, polyurethane composites and water-reducible alkyd resins, pharmaceutical products, plasticizers, adhesives, and surfactants. Rubber seed oil is a promising substitute to linseed oil as semi-drying oil for oily paint formulation. ¹H NMR analysis revealed that the fatty acid compositions consist of linoleic acid (34.22 wt%), oleic acid (28.6 wt%), linolenic acid (18.6 wt%), and saturated fatty acids (18.57 wt%). FTIR analysis indicated fingerprint regions of 1461 to 585 cm⁻¹ which can be used to check adulteration of the oil. The NMR spectra (¹H and ¹³C) of the oil are similar to those of other vegetable oils with well-identified peaks and regions that can be used to authenticate the quality of the oil.

Key words: Rubber seed oil, characterization, nuclear magnetic resonance (NMR), Fourier transform infra-red spectroscopy (FTIR), quality assessment, technical applications.

INTRODUCTION

Edible vegetable oils have several food applications and they are increasingly being used for other purposes such as resinous pigments and drying oils in paints and coatings production (Aigbodion and Bakare, 2005). To reduce costs associated with the use of edible vegetable oils in the food industry, non-edible oils of good industrial applications are highly sought to replace edible oils (Roschat et al., 2017). In recent times, with the development of new technologies and research future areas expansion on non-edible oils applications to replace edible oils, detailed characteristic property assessment of oils quality are imperative to the entrepreneurs (Reshad et al., 2015). Therefore, novel oils characterization to determine their possible applications...
and to forestall adulteration because of price differentials with low quality oils is paramount for high quality derivable products (Barison et al., 2010).

Recently, several researchers studied rubber seed oil (RSO) usefulness to authenticate its industrial application as a renewable resource material to complement the fast depleting non-renewable mineral oils of fossil origins (Onoji et al., 2019). Rubber tree accounts for 99% of the world’s natural rubber (NR) latex used for the production of several rubber products (Atabani et al., 2013). Currently, the seeds are underutilized and allowed to rot away in the plantations, apart from a minimal use in subsequent tree plant breeding process in Nigeria (Onoji et al., 2017). The rubber seeds used in this study were collected from the 40-hectare (ha) NIG800 series plantations of Rubber Research Institute of Nigeria (RRIN), Iyanomo, Benin-City (Aigbodion and Bakare, 2005). The foreign (Malaysia, Sri Lanka, etc.) rubber seeds (RRIM 600, RRIM 501, PB 28/59, PB 5/63, RRIM 628, RRIC 45, RRIM 614, AVROS 1581, RRIM 605, PB 5/51, GT1, RRIM 62, RRIM707, PR107 and PB217) used as parentage stock in these plantations were pre-germinated in pre-nursery beds (Umar et al., 2010). Ground and polythene bags nursery techniques were used for seedlings production. Certified rubber seedlings obtained through budding techniques were used for planting in these plantations. Specie of Nigerian hybrid is a high-yield latex (3000 to 3500 dry NR/hectare/year) tree, resistant to wind and with a capacity to produce about 1200 seeds/tree/year (Onoji et al., 2020). The species has an average weight of 4 g/seed (Onoji et al., 2019), and a non-edible oil content yield of about 43 wt% (Onoji et al., 2016). Non-edible seed oil from rubber tree has been identified as a potentially promising material for several industrial applications such as, the production of biodiesel (Yang et al., 2011), oleochemicals (Hosamani and Katagi, 2008), polyurethane composites and water-reducible alkyd resins (Bakare et al., 2010), pharmaceutical products, plasticizers, adhesives, and surfactants (Onoji et al., 2016). De-oiled cake, after pretreatment to remove toxic material such as hydrogen cyanide, is suitable as a valuable source of proteins for farm animals and poultry feeds (Eka et al., 2010). RSO has been characterized as a semi-drying oil (Aravind et al., 2015), and of a high quality that makes it a promising substitute for linseed oil in paint formulation (Ebewele et al., 2010). In Nigeria, linseed oil is imported for use in the paint industry and other applications (Okienmen et al., 2005). However, available report shows that RSO was yet to have any commercial value in Nigeria, which has the capacity to produce 13,000 tons RSO/year (Okienmen et al., 2005). The oil can be processed into a semi-drying oil for the paint industry to reduce importation of linseed oil which depletes the Nigerian foreign exchange reserves. There is therefore a need to characterize the Nigerian hybrid RSO in order to authenticate its quality against any adulteration with respect to the aforementioned usage.

Vegetable oils authentication for product quality via spectroscopic methods use is well reported in literature (Sadowska et al., 2008). Gas chromatography coupled with mass spectrometry (GC-MS) has been in use for several decades (Skogge et al., 2007). However GC-MS, a destructive method, involves oil sample chemical modification (oxidation) with the tendency to produce unreliable results. In addition, GC-MS is cumbersome, cost in-effective, time consuming and may pose problems in result interpretations, especially the fatty acid compositions (Barison et al., 2010). Fourier transform infra-red spectroscopy (FTIR) is a non-destructive technique reported for use in the analysis of free fatty acids and to monitor oil quality to check adulteration (Valente et al., 2016) through the identification of the functional groups present in the sample (Bohre, 2013). Proton \(^{1}\text{H}\) and carbon-13 \(^{13}\text{C}\) nuclear magnetic resonance (NMR) spectroscopy is a non-destructive technique used to determine the proportion of different acyl groups present in oils and fats, and other liquids in a very short time for sample preparation and experimental spectrum acquisition compared to GC-MS (Scano et al., 2008; Guillén and Ruiz, 2003a). \(^{1}\text{H}\) NMR spectra area signals are proportional to the number of hydrogen atoms that produce the signals (Guillén and Ruiz, 2003a). This method has many advantages because it does not involve chemical modification of sample like the GC-MS (Yeung et al., 2008).

In this study, seed oil was extracted by mechanical means notable for high-quality oil (Ebewele et al., 2010). This method is cost effective and can be adapted to small- to medium-scale entrepreneurship due to its low operating and maintenance costs. For instance, a demonstration-scale rubber seed oil mechanical extractor designed and fabricated by RRIN located in Benin City at the cost of US$1,050 has a capacity to extract 500 L of oil/day. The extracted oil physico-chemical properties were determined by using standard methods described elsewhere (Onoji et al., 2016), while FTIR and NMR spectroscopy were deployed to determine oil functional groups and fatty acids, respectively.

**EXPERIMENTAL**  
**Materials and reagents**  
Fresh and glossy rubber seeds used for this study were handpicked

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from the hybrid rubber estate plantation of RRIN. All reagents from BDH Chemicals Ltd., Poole England, and GFS Chemicals, Inc., 867 McKinley Ave., Columbus, OH 43223 and used in the analyses were of analytical grades.

Seed oil extraction

Seed kernels were separated from seed shells by gentle cracking using a laboratory mortar and pestle. The kernels were weighed, and dried in an oven at 110°C for 2 h to attain constant weight for moisture content determination. About 2 kg of fresh seed kernel was dried at 110°C to 7 wt% moisture content for a maximum oil yield (Igeleke and Omorusi, 2007). Seed oil was extracted at 30°C from the dried kernel using a mechanical oilseed screw press. Extracted oil was stored in screwed airtight plastic containers for physico-chemical characterization, and spectroscopic analysis.

Physico-chemical analysis of rubber seed oil

The standard methods adopted for the determination of physico-chemical properties of the rubber seed oil extracted using a 99.9% n-hexane solvent in a soxhlet extractor were obtained elsewhere (Onoji et al., 2016). The experiments were carried out in duplicate, and the average values recorded for accuracy of data.

Spectroscopic analysis of rubber seed oil

Oil infrared spectra were recorded using an FTIR spectrometer (Model: TENSOR 27, Bruker Optics Inc., USA) equipped with a detector, and interfaced to a personal computer operating under OPUS spectroscopy software supplied together with the instrument. A film of 2 μL of the RSO sample was deposited between two disks of KBr, in the absence of air (Guillén and Cabo, 2002). Sinclair et al. (1952) reported that the degree of unsaturation in vegetable oils correlates with some absorbance bands in the FTIR spectrum, hence the frequencies of such bands are closely related to the proportions of saturated, monounsaturated, and polyunsaturated acyl groups (Guillén and Ruiz, 2003a). Consequently all spectra were recorded between 4000 and 500 cm⁻¹.

Extracted oil ¹H NMR spectrum was recorded on Bruker ultra-shield TM 500-MHz NMR spectrophotometer. Approximately 0.2 g of oil sample was dissolved in 500 μL of deuterated chloroform (CDCl₃) as solvent (δ = 7.26 ppm) containing a small amount of tetramethylsilane (TMS) as an internal standard (δ = 0 ppm). This was then placed in 5 mm diameter NMR test tubes to commence the analysis (Guillén and Ruiz, 2003a). The acquisition data were: spectral width 8.1 kHz, relaxation delay 3s, 15 scans, and pulse width 30°.

¹³C NMR experiments were also carried out with Bruker ultra-shield TM 500-MHz NMR equipment. About 25 mg of RSO was dissolved in 1 mL of CDCl₃. Samples were placed in 5 mm NMR tubes at 20°C and analyzed within 2 days as described by Scano et al. (2008). The experiments were conducted at room temperature (28°C). ¹³C spectra were recorded at 235.2 ppm spectral width, relaxation delays 3s, and a total of 350 scans. The carbon atom in CDCl₃ was observed at 77.42 ppm. All other peaks were assigned with respect to it.

Fatty acid compositions of rubber seed oil using NMR spectroscopy

Unsaturation degree in oils and fats can also be determined from the proportion of olefinic hydrogen atoms obtained from ¹H NMR data, besides different acyl groups present in oil sample (Guillén and Ruiz, 2003b). The proportions of Linoleic (Ln), Linoleic (L), Oleic (O), and saturated (S) fatty acids present in the RSO were calculated from the areas of the identified peaks (1 – 10) in ¹H NMR spectra (Figure 1) using parameters shown in Equations 1 to 4 for oils containing similar acyl groups as that mentioned previously (Reshad et al., 2015; Guillén and Ruiz, 2003b).

\[
L_n (%) = 100 \left( \frac{n}{A + B} \right)
\]

\[
L(\%) = 100 \left[ \frac{1}{2} - \frac{1}{2}\left( \frac{A}{B} \right) \right]
\]

\[
O(\%) = 100 \left[ \frac{C}{10} - \frac{1}{2}\left( \frac{A}{B} \right) \right]
\]

\[
S(\%) = 100 \left[ 1 - \frac{C}{2A} \right]
\]

where A, is the area signal 1, corresponding to methyl hydrogen atoms of saturated, oleic (omega-9), and linoleic acyl groups; B is area signal 2, corresponding to methyl hydrogen atoms of linoleic acyl groups; C is area signal 5, corresponding to methylene hydrogen atoms in position α, in relation to one double bond (allylic protons); D is area signal 6, corresponding to methylene hydrogen atoms in position α, in relation to the carboxyl group; and E is area signal 7, corresponding to methylene hydrogen atoms in α position with two double bonds (bis-allylic protons).

RESULTS AND DISCUSSION

Physico-chemical properties of rubber seed oil

Mechanically extracted hybrid oil properties are presented in Table 1, with oil yield of 28 wt%. This yield closely agrees with the yield of 23 wt% reported by Aigbodion and Bakare (2005) on similar experiment with RSO. The oil parameters such as iodine value of 136.07 g I₂/100 g oil, peroxide value of 9.45 O₂/kg oil, saponification value of 189.5 mg KOH/g oil, and refractive index of 1.472 are within the range reported by other researchers on RSO (Reshad et al., 2015; Aravind et al., 2015).

FTIR and NMR spectroscopic analysis of rubber seed oil

Results obtained from oil FTIR and NMR analysis are geared towards providing an insight into the fatty acid structural configuration and, compositions that enables easy evaluation of its potential applications by entrepreneurs.

FTIR analysis of rubber seed oil

Figure 2 shows the FTIR spectrum of the extracted oil. The main peaks and their assignment to functional groups are presented in Table 2. The O–H stretching vibrations bands (for alcohols, carboxylic acids, and hydroperoxides) usually in the absorption range of 3200 to 3600 cm⁻¹ for vegetable oils (Ogbru and Ajiwe, 2016).
Onoji et al.

Figure 1. $^1$H NMR spectra of rubber seed oil.

Table 1. Physico-chemical properties of rubber seed oil.

| Parameter                             | Reshad et al. (2015)          | This study                   |
|---------------------------------------|-------------------------------|------------------------------|
| Colour                                | Golden yellow                 | Dark brown                   |
| Density, g/cm$^3$ at 25°C             | 0.910                         | 0.892 ± 0.01                 |
| Specific gravity at 15°C              | -                             | 0.91 ± 0.02                  |
| Oil content (wt%)                     | 40–50                         | 28 ± 0.2                     |
| Oil water content (wt%)               | -                             | 0.075 ± 0.01                 |
| Initial kernel water content (wt%)    | -                             | 9.85 ± 0.03                  |
| Iodine value, g I$_2$/100 g oil       | 113                           | 136.07 ± 0.04                |
| Peroxide value, meq. O$_2$/kg oil     | -                             | 9.45 ± 0.06                  |
| Saponification value, mg KOH/g oil    | 235.28                        | 189.5 ± 0.13                 |
| Acid value, mg KOH/g oil              | 24                            | 16.5 ± 0.1                   |
| Free fatty acid (%FFA as oleic acid)  | 12.12                         | 8.25 ± 0.05                  |
| Kinematic viscosity, mm$^2$/s at 40°C  | 30                            | 38.5 ± 0.04                  |
| Refractive index at 20°C              | 1.47                          | 1.472 ± 0.001                |
| Pour point, °C                        | −1.5                          | −5                           |
| Cloud point, °C                       | 3                             | 4.8                          |
| Cold filter plugging point, °C        | −                             | −0.62                        |
| Flash point, °C                       | −                             | 243                          |
| Fire point, °C                        | −                             | 254                          |
| Aniline point, °C (°F)                | −                             | 21 (69.8)                    |
| Boiling point, °C                     | −                             | 122                          |
| Freezing point, °C                    | −                             | −18                          |
| Higher heating value (HHV), MJ/kg     | 39.34                         | 39.81                        |
| Mean mol. weight of fatty acids (g/mol)| -                            | 295.5                        |
| Average mol. weight of RSO (g/mol)    | -                             | 924.56                       |

Values are mean ± standard deviation of duplicate data.
Figure 2. FTIR spectrum of rubber seed oil.

Table 2. Description of FTIR analysis of rubber seed oil.

| Wavenumber (cm⁻¹) | Functional group | Description of vibration                     |
|-------------------|------------------|---------------------------------------------|
| 3009              | Olefinic         | =C–H stretching                             |
| 2921-2853         | Methylene        | Strong peaks stretching vibration of C–H    |
| 1742              | Carboxyl         | −C=O stretching vibration of esters         |
| 1710              | Carboxylic acids | Stretching vibration of C=O                  |
| 1461              | Methyl, methylene| Medium signal due to bending vibration of alkyl C–H bonds |
| 1377              | Methyl           | Alkyl C–H bending vibrations                 |
| 1238              | Carboxylic acids, Esters | Stretching vibration of C–O             |
| 1160              | Esters           | C–O stretching vibration of ester groups    |
| 1118-1033         | Esters           | =C–O–C stretching vibration of ester groups |
| 721               | Methylene        | Bending (rocking) vibration of C–H          |
| 585               | Alkanes          | Weak signal due to C–H vibration            |

are absent in the RSO used for this study. The observed spectrum implies that triglyceride is the main component of the oil as a result of the strong presence of ester carbonyl group (C=O) at 1742 cm⁻¹. Ester group presence could be attributed to the stretching vibrations of C–O observed at 1160 cm⁻¹. Results on fingerprint obtained are within the range (1461 to 585 cm⁻¹) which provides useful information to detect oil adulteration. The results obtained are within the range reported by other researchers on RSO (Reshad et al., 2015).

NMR spectra analysis of rubber seed oil

Figure 1 depicts the ¹H NMR spectrum of RSO. The assignment of signals is listed in Table 3. Signal 1 is the overlap of doublet signals of methyl group protons that appears between 0.8 and 0.81 ppm. Signal 2 is a singlet methyl group proton, and appears at 0.89 ppm. Signal 3 appears between 1.19 and 1.23 ppm corresponding to linolenyl chains protons, and saturated methylene group of acyl chains, respectively. Signal 4 appears at 1.53 ppm,
Table 3. $^1$H NMR and $^{13}$C NMR analysis of rubber seed oil.

| $^1$H (ppm) | $^{13}$C (ppm) | Functional groups | Assignments |
|-------------|----------------|------------------|-------------|
| 0.80 - 0.89 | 13.91          | CH$_3$ (saturated, oleic, and linoleic acids) | All acyl chain except linolenic |
| 1.19        | 33.97          | CH=CHCH$_2$CH$_3$ | Linolenic chains |
| 1.23 - 1.4  | 29 - 31        | (CH$_2$)$_n$ saturated aliphatic | All acyl chains |
| 1.53        | 21.9           | CH$_2$=CH$_2$COOH | All acyl chains |
| 1.94 - 1.98 | 26.1           | CH$_2$CH=CH      | All unsaturated acyl chains |
| 2.22        | 33.97          | CH$_2$-COOH      | All acyl chains |
| 2.68; 2.72  | 24.9           | CH=CHCH$_2$CH=CH | Linolenic and linoleic chains |
| 4.05, 4.06; 4.22, 4.23 | 61.93 - 77.16  | CH$_2$-OCOR     | Triglycerides (glyceryl group) |
| 5.25        | -              | CH-OCOR          | Triglycerides (glyceryl group) |
| 5.26        | 126.99 - 131.99| CH=CH (olefinic group) | All unsaturated fatty acids |
| -           | 172.41 - 172.82| C=O (carbonyl group) | - |
| -           | 178.64         | COOH (carboxylic acids) | - |

Figure 3. $^{13}$C NMR spectra of rubber seed oil.

represents the methylene protons in the $\beta$ position. Signal 5 appears between 1.94 and 1.98 ppm, due to $\alpha$-methylene proton related to a proton in $\alpha$ position in relation to a single double bond (allylic protons). Signal 6 occurred due to the methylene protons in $\alpha$ position, and it appears at 2.22 ppm. Signal 7 is a signal overlap between 2.68 and 2.72 ppm, due to responses from $\alpha$-methylene protons related to the double bonds (bi-allylic protons). Signal 8 appears at 4.05, 4.06 and 4.22, 4.23 ppm, due to the protons on carbon atoms 1 and 3 of the glyceryl group. Signal 9 appears at 5.25 ppm, corresponding to the proton on carbon 2 atom of the glyceryl group, and it overlaps with Signal 10 at 5.26 ppm that represents olefinic protons of different acyl groups.

$^{13}$C NMR spectra analysis of rubber seed oil

Figure 3 shows the $^{13}$C NMR spectrum of the oil under analysis with well-defined four distinct regions (A, B, C and D): A (13.91 - 33.97 ppm) due to methyl or alkyl functional groups for saturated acids, oleic, and linoleic acids; B (61.93 - 77.16 ppm) due to glyceryl C-atoms; C (126.99 - 131.99) due to the presence of unsaturated alkenes; and D (172.41 - 178.64) due to the presence of carbonyl and carboxylic acid groups. The assignment of
Table 4. Fatty acid compositions of rubber seed oil via $^1$H NMR spectra.

| Fatty acid/systemic name                        | Chemical formulae  | Composition (%) |
|------------------------------------------------|-------------------|-----------------|
| Saturated (C$^ullet_{xx:0}$)                  |                   | Reshad et al. (2015) | This study |
| Oleic (C$^{18:1}$/ cis-9-Octadecenoic           | $C_{18}H_{34}O_2$ | 19.91           | 18.57      |
| Linoleic(C$^{18:2}$/ cis-9-cis-12-Octadecadienoic | $C_{18}H_{32}O_2$ | 27.06           | 28.6       |
| α-Linolenic (C$^{18:3}$/ cis-9-cis-15-Octadecatrienoic | $C_{18}H_{30}O_2$ | 39.86           | 34.22      |
| Total saturated                                 |                   | 13.17           | 18.6       |
| Total monounsaturated                           |                   | 19.91           | 18.57      |
| Total polyunsaturated (linoleic, linolenic)     |                   | 27.06           | 28.6       |
| Total unsaturated (oleic, linoleic, linolenic)  |                   | 53.03           | 53.82      |
|                                                   |                   | 80.09           | 81.42      |

XX means 14, 16 or 18.

The fatty acid compositions of vegetable oils depend to a large extent on the seed storage and processing time, extraction technique employed, and method of oil analysis etc. Based on Equations 1 to 4 and the identified peak areas of the $^1$H NMR spectrum (Figure 1), the fatty acids composition was quantified and tabulated in Table 4. The spectra analysis showed that the fatty acid compositions of the extracted RSO consist of linoleic acid (34.22 wt%), oleic acid (28.6 wt%), linolenic acid (18.6 wt%), and saturated fatty acids (18.57 wt%), and it compared favorably with results on RSO from other researchers (Reshad et al., 2015). The results show that linoleic acid is the predominant fatty acid present in this hybrid RSO and over 80% of the fatty acids are unsaturated.

Conversely, the oil may be categorized among the less nutritional quality due to the low profile of oleic acid (28.6 wt%) compared to olive oil (64.6 - 84.4 wt%) and canola (60 - 75 wt%) (Reshad et al., 2015). In addition, due to the high content of unsaturated fatty acids coupled with the low temperature properties, the oil is a promising material for biodiesel production, and as a substitute for linseed oil suitable for making drying oil for paint and varnish formulations.

**Conclusions**

In this study, rubber seed oil was mechanically extracted from the seeds of Nigerian hybrid rubber trees with oil yield of 28 wt%. An Adulteration of high-value RSO with low-grade oils often constitutes economic and commercial loss. Consequently chemical analysis to check for adulteration revealed that the RSO is non-edible and possesses significant potentials as substitute to the edible oils on different industrial applications. For instance, the oil can be used as a replacement for linseed oil as a semi-drying oil in the paint industry. The spectroscopic analytical methods (FTIR and NMR spectroscopy) employed in the study are non-destructive, simple, fast, reliable, cost-effective, and with no sample pre-treatment required compared to the GC-MS analytical tool which involves chemical modification of oil samples. The $^1$H NMR spectra analysis of the oil shows high degree of unsaturation which implies susceptibility to low temperature, hence suitable for low climatic regions. The observed FTIR fingerprint regions of the oil are adequate for adulteration detection of low quality RSO that may impair derived products. The –C=O, =C–O–C, and C–O vibrations in the FTIR bands of 1742, 1118-1033, and 1160 cm$^{-1}$ indicate the presence of strong ester groups supportive of biodiesel production. The findings of this study are useful parametric data for rubber seed oil identification, quantification, and authentication purposes.

**CONFLICT OF INTERESTS**

The authors have not declared any conflict of interests.

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