

Quality Assessment of some Nigerian Branded Honey

ABSTRACT

Three branded honey samples purchased from different supermarkets in Ekiti State, Southwest Nigeria were analyzed for their pollen contents and for some physicochemical attributes. The pollen composition of the three branded honey investigated revealed important honeybee plants such as *Launea sp.*, *Hymenocardiaacida*, *Alchornea cordifolia*, *Phyllanthus sp.*, *Danelliaoliveri* and the family members of Fabaceae, *Myrtaceae* and *Combretaceae*. The result of the proximate compositions revealed low moisture content, crude protein, crude fat, crude fiber, high carbohydrate and energy content while the physicochemical parameters revealed that the specific gravity, pH, total acidity, Hydroxymethyl Furfural (HMF) and total sugar agreed with the international standard. Generally, the result showed that the three honey samples were rich in floral pollen types and equally removed doubt or suspicious on their acclaimed geographical origin. Also, the physicochemical results of the three branded honey samples prove that they are of good quality and are all suitable for human use.

Keywords: honey bee, pollen, physicochemical, floral, Ekiti State.

1. Introduction

Melisopalynology is an applied branch of palynology dealing with the study of pollen grains in honey samples and its application in apiculture. Honey is one of the major bee product obtained from nectar, sugary secretion and pollen as well as **nectaries**. Honey is highly valuable syrup and it is preferred to other sweetens because of its nutritional, medicinal and industrial purposes [1]. Because of various uses of honey from food to medicine, it is of great interest to carry out complete evaluation of honey in order to formulate values and ranges of various honey characteristics.

Branded and unbranded honeys are available in the market. Most of the honey sold in grocery stores is not raw honey. In some cases, such honey has been heated or adulterated by addition of syrup or other food ingredients. As an industry, it has become necessary to address what happens to honey after it has been removed from the bee hive, blended and filtered. Consumers are unaware of the quality of honey they consume and imminent danger in adulterated honey. Honey adulteration could result in serious health problems. In order to protect the consumers and also to give them quality for their money, several stringent measures have been set by various countries and organization for accessing the authenticity and quality of honey produced for commercial uses. The International Honey Commission (IHC) has propose certain constituent criteria, such as moisture content, electrical conductivity, reducing sugar, fructose and glucose content, sucrose content, minerals, free acidity and **HydroxyMethyl furfural** (HMF), as quality criteria for honey [2].

In addition to these criteria, there is need for **melisopalynological** analysis in order to determine botanical and geographical origin of the honey. Authentication of honey origin through its botanical elements [3] provide reliable information which help to curb the practice of wrong labeling of honey. Also, information obtained from honey characterization allows the packaging and storage of honey in appropriate conditions so as to preserve their qualities and **savour** (flavor)

of honey or savor) [4]. Lawal *et al.* [5] reported that honey is produced in bee hives in large quantity in Nigeria. However, Ayodele *et al.* [6] was of the opinion that despite the fact that Nigeria has great potential of bee keeping, only a small quantity of honey produced is commercially important due to adulteration and poor handling during processing. Honey farmers are now branding and rebranding their product in order to build consumer confidence and enhance patronage to this highly valued commodity.

Albeit, many authors have conducted several research work on quality assessment of Nigeria honey. Agbagwa *et al.* [7] carried out a comparative study on some honey sample in Nigeria and Manuka honey in order to establish the quality of Nigeria honey. Assessment of quality attribute of natural honey from Adamawa state, North eastern Nigeria was reported by Igwe *et al.* [8]. Quality assessment of honey sourced from natural and artificial apiaries in Ekiti State, Nigeria, was investigated by Oyeyemi [9]. Data on complete analysis of branded Nigeria honey are scanty. The need to restore consumer's confidence and attract more patronage of Nigeria honey both home and abroad prompted this study.

Therefore, the study was carried out to access the quality attribute of some branded honey samples in Nigeria.

2. MATERIALS AND METHODS

Three branded honey samples were purchased from the supermarkets in Ado Ekiti, Ekiti State, Nigeria. The branded honey samples were taken to the Department of Agricultural Extension Laboratory, Faculty of Agricultural Science, Ekiti State University for physicochemical analysis.

2.1 Physicochemical Analysis

Physicochemical parameters such as moisture content, pH, total acidity, specific gravity, HMF and total sugar were determined.

2.2 Determination of the pH

A pH digital meter was calibrated with buffers at pH 4 and 10. Sample solution was taken in the beaker and inserted. When the first reading was completed, the electrode was washed with distilled water and dried-up with tissue paper. Similarly, as a continue series, all other samples were determined accordingly [10].

2.3 Determination of moisture content

Two grams each of the honey sample was weighed and transferred into a pre-weighed crucible. The crucible was kept in an oven at 100- 105⁰ C overnight. After this, they were removed and cooled in a desiccator and re-weighed. The loss in weight was then calculated as the percentage moisture content ^[11] using the following formular:

$$\text{Moisture} = \frac{\text{Weight of fresh honey sample} - \text{Weight of dry honey sample}}{\text{Weight of fresh honey}}$$

2.4 Determination of specific gravity

The specific gravity (SG) of the honey samples was obtained as the ratio of the weight of sample to that of equal volume of water.

$$SG = \frac{W_{sp} - W_p}{W_{wp} - W_p}$$

Where;

W_p = Weight of the pycnometer

W_{sp} = Weight of sample + pycnometer

W_{wp} = Weight of water + pycnometer

2.5 Determination of HydroxyMethyl Furfural(HMF)

For HMF, 5 g each of the honey samples was accurately weighed into a beaker and dissolved in approximately 25 ml of water and quantitatively transferred into a 50 ml volumetric flask. 0.5 ml of carez solution I was added into the sample and mixed followed by further addition of 0.5 ml of carez solution II, mixing and finally topping it up to the mark with water. The prepared sample solution was filtered through Whatman filter paper no. 42. The first 10 ml of the filtrate were rejected while 5 ml of the filtrate was transferred into each of the two test tubes. 5 ml of distilled water was added into one of the test tubes and mixed well (the sample solution). 5 ml of 0.2% sodium bisulphate was added into the second test tube and mixed well (the reference solution). The absorbant of the sample solution was determined against the reference solution at 284 and 336 nm in 10mm quartz cells within 1hour. The determinations were done in triplicates. The HMF values were calculated using the formular shown below.

$$\text{HMF in mg/kg} = (A_{284} - A_{336}) \times 149.7 \times 5 \times D/W$$

Where A_{284} = Absorbance at 284nm

A_{336} = Absorbance at 336nm

149.7 = constant

D = dilution factor in case dilution was necessary

W = weight in g of the honey samples

The results were expressed in mg/kg to the nearest 1 decimal place.

2.6 Determination of total sugar

For this solution, 5 g of sample was taken into a beaker and 100 ml of warm water was added to it. The solution was stirred until all the soluble matters were dissolved and filtered through Whatman filter paper into a 250 ml volumetric flask. 100 ml of the solution was pipetted and prepared into a conical flask, after which 10 ml of diluted hydrogen chloride (HCl) was added and boiled for 5min. On cooling, the solution was neutralized to phenolphthalein with 10%NaOH and kept in a 250 ml volumetric flask [12]. This solution and the reading was(were) calculated as follows:

$$\text{Total sugar (\%)} = \frac{\text{Factor (4.95)} \times \text{dilution (250)} \times 2.5}{\text{Titre} \times \text{wt. of sample} \times 10}$$

2.7 Determination of the energy values

The energy values of the samples were determined as follows:

$$\text{Energy (Kcal/100g)} = (\% \text{ Crude Protein}) \times (\% \text{ Crude Fat}) \times (\% \text{ Carbohydrate})$$

2.3 Pollen analysis(2.8 Pollen analysis)

The honey samples were subjected to qualitative and quantitative analyses following the methodology recommended by the International Commission for Bee Botany^[13]. The examination and photomicrography of pollen grains were done using the Trinocular Olympus CH30 Microscope with X40 objective lens and a DE 1.3 MegaPixel Digital Camera attachment. The pollen types were identified with the help of reference slides and relevant literature from Nigerian plants in the Palynology Laboratory of the Department of Archeology, University of Ibadan, Oyo State, Nigeria. The pollen types were identified to generic, species and in some cases to family level. Pollen grain numbers were quantified using the techniques put forward by [14]. Results were expressed as frequency classes using the method suggested by [13].

3.0 Results(3. Results)

Results of the proximate composition of the three branded honey samples (A1, A2 and A3) are presented in Table 1. The moisture contents ranged between 12.09 to 15.50%, crude fat (0.30-0.41%), crude protein (0.68-0.84%), crude fiber (0.12-0.25%), ash (0.44-0.65%), Carbohydrate (80.19-85.31%) and energy content (351.12-347.95 kcal/100kg). The physiochemical parameters of the three honey samples showed pH range between 3.48 to 4.39, specific gravity (1.28-1.41), total acidity (28.20-56.62meq/kg), HMF (34.92-135.70mg/kg) and total sugar (68.23-70.15g). The pollen analysis results of the three honey samples were shown in Table 3. Seventeen different pollen types were identified among which the taxa were further identified to species level, 4 to generic level and 5 were identified to family level.

The quantitative pollen analysis of A2 honey sample revealed 20 different pollen types belonging to 10 families and 15 of the pollen types were identified to species level, 7 to generic level and 4 to family level. The result obtained for honey sample A3 revealed that 17 pollen families were identified in the sample. A further identification revealed 11 pollen types (species level), 5 were identified to generic level and 4 to family level. The pollen frequency of the pollen types found in the three honey samples showed that none of the pollen types occurred as “very frequent” and “frequent” classes. However, seventeen pollen types were “sporadic” and thirty-four were “rare”. Fig.1. depicted the photomicrographs of some pollen grains identified in the three honey samples as shown under the microscope.

4.0 Discussion(4. Discussion)

The physiochemical data of any honey sample is important for storage and marketing purposes. The results obtained for the three honey samples purchased from the groceries in Ado-Ekiti were compared with the EUC [15]. The results showed that the moisture contents of the three branded honey samples were low and were found to be within the limit of not more than 21% as prescribed by [16] and European Union Standard of honey sample [15]. Moisture content is one of the basic quality characteristics of honey. It contributes to honey viscosity, density, specific gravity, refractive index, fermentation and **savour**[2]. The 3 branded honey samples investigated

were found to be acidic in character. Their pH values are similar to the pH values of the Nigerian honey (3.55-4.40) reported by Oyeyemi ^[9] and Ethiopia honey (3.82-4.45) reported by Nigusie *et al.* [17]. The observed pH values for these honey samples were within the acidic range of pH and low enough to inhibit the growth of microbial growth.

Table 1: Physicochemical composition of branded Nigerian honey samples

Parameters	A1	A2	A3
Specific gravity	1.41	1.40	1.38
pH	4.39	3.84	4.26
Total acidity (meq/kg)	28.20	56.62	30.10
HMF(mg/kg)	35.70	34.92	78.90
Total sugar (g)	70.15	68.23	69.28

A1= Shalomisreal honey, Benue State, A2= Kingsway honey, Oyo State, A3= Afe Babalola University honey, Ekiti State.

The results of the specific gravity of the three branded honey examined fall within the prescribed standard limit. Codex Alimentarius Commission[16] prescribed the specific gravity range of honey sample to be between 1.38-1.45. The findings of this study conform to the study of Ndifeet *al.*[18] who reported a specific gravity ranging from 1.41-1.44 for Nigerian honey. HMF content of A1, A2 and A3 honey samples were found to fall within the limit of 40mg/kg prescribed in normal honey. HMF is a breakdown of fructose formed slowly during storage and very quickly when honey is heated. The amount of HMF found in honey sample could be used as a guide to storage period and the amount of heating which has taken place. High levels of HMF (> 100mg/kg) can also be an indicator of adulterated with inverted sugars. The low HMF in honey sample A1 and A2 indicated that they are fresh and of good quality.

Table 2: Proximate composition of branded Nigerian honey samples

Parameters	A1	A2	A3
Moisture content	12.09	12.63	15.50
Crude fat (%)	0.34	0.41	0.30
Crude protein (%)	0.84	0.76	0.68
Crude fiber (%)	0.12	0.25	0.20
Ash (%)	0.44	0.65	0.58
Carbohydrate (%)	80.19	85.31	82.74
Energy (kcal/100g)	351.12	347.95	336.38

Crude protein determination gave results that were similar to those obtained by Adeniyi *et al.*[19] who reported protein content of 0.69% and 0.74% for bitter and sweet honey in Nigeria. Meanwhile the obtained values were low compared to the report of Sohaimy and Shahata[20] for honey samples obtained in Egypt. The crude protein contents showed that honey is not an adequate source of dietary protein. It is well known that honey contains a trace amount of protein usually originated from pollens which is a natural and protein-rich food source[21]. The concentration of protein in honey varies depending on their botanical or geographical origin and storage period.

The range of ash content of the three honey samples was higher than 0.09%-0.021 % reported by Ouchemouki[22] for Algeria polyfloral honey. However, the results were within the limit of ash content proposed by the Codex Alimentarius Standards [23]. Honey ash content is a reflection of its richness in organic minerals and is determined by the botanical source [24].

The result of the carbohydrate contents (80.19-85.31%) obtained in this study was higher than the earlier reports of Buba *et al.* [25] and Adeniyi *et al.* [19]. The higher carbohydrate content in the three honey samples analyzed could be as a result of wide foraging activities of the bees on varieties of **necteriferous** plants. Glucose and fructose are the major component of carbohydrate in honey. The total sugar content in honey samples analyzed was slightly higher when compared with previous work of Abdulkhalia and Swales[26]who reported a range of 79.0-84.0% for some honey sample from the West Bank, Palestine.

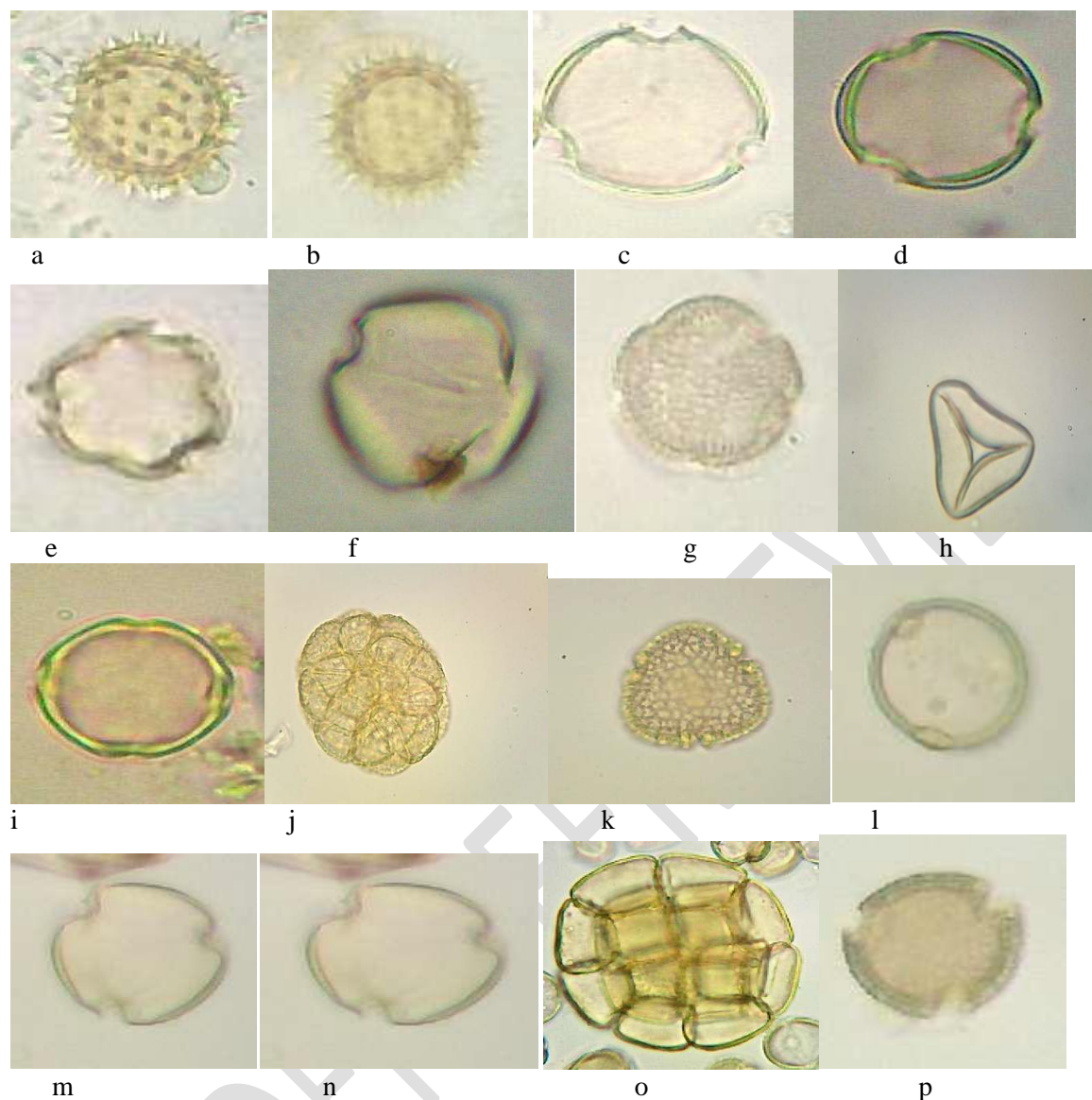
Table 3: Percentage pollen frequency class of branded Nigerian honey samples

Sample	Pollen type	%Frequency	Frequency class
A1	<i>Launea sp.</i>	7.62	Rare
	<i>Elaeisguineensis</i>	5.71	Rare
	<i>Tridax procumbens</i>	2.86	Sporadic
	<i>Danielia oliveri</i>	2.86	Sporadic
	<i>Combretum sp.</i>	11.43	Rare
	<i>Diospyros sp</i>	3.81	Rare
	<i>Alchornea cordifolia</i>	9.52	Rare
	<i>Hymenocardiaacida</i>	9.52	Rare
	<i>Phyllantus sp.</i>	8.57	Rare
	<i>Tetrapleura tetraptera</i>	3.81	Rare
	<i>Mitrocarpusscaber</i>	1.91	Sporadic
	<i>Vitellaria paradoxa</i>	3.81	Rare
	Asteraceae	4.76	Rare
	Fabaceae	5.71	Rare
	Combretaceae	7.62	Rare
	Myrtaceae	5.71	Rare
	Solanaceae	2.86	Rare
A2	<i>Cocos nucifera</i>	2.08	Sporadic
	<i>Bombax sp.</i>	1.62	Sporadic
	<i>Daniella oliveri</i>	3.01	Rare
	<i>Dolenix regia</i>	2.55	Sporadic
	<i>Combretum sp.</i>	3.94	Rare
	<i>Terminalia sp.</i>	1.16	Sporadic
	<i>Diospyros sp.</i>	2.08	Sporadic
	<i>Alchornea cordifolia</i>	10.64	Rare
	<i>Bridellia sp.</i>	5.32	Rare
	<i>Hymenocardiaacida</i>	6.48	Rare
	<i>Mallotussubutalus</i>	2.78	Sporadic
	<i>Phyllanthus sp.</i>	15.57	Rare

	<i>Acacia sp.</i>	2.08	Sporadic
	<i>Tetrapleura tetraptera</i>	1.85	Sporadic
	<i>Triplochiton scleroxylon</i>	1.16	Sporadic
	Fabaceae	2.08	Sporadic
	Combretaceae	8.80	Rare
	Myrtaceae	7.87	Rare
	Rubiaceae	5.82	Rare
A3	<i>Launea sp.</i>	5.53	Rare
	<i>Cocos nucifera</i>	1.38	Sporadic
	<i>Elaeisuineensis</i>	14.29	Rare
	<i>Hyphene sp.</i>	1.38	Sporadic
	<i>Aspillia Africana</i>	3.69	Rare
	<i>Tridax procumbens</i>	2.76	Sporadic
	<i>Emilia sp.</i>	1.38	Sporadic
	<i>Vernonia amygdalina</i>	6.45	Rare
	<i>Bombax sp.</i>	1.84	Sporadic
	<i>Alchornea cordifolia</i>	10.60	Rare
	<i>Phyllanthus sp.</i>	15.67	Rare
	<i>Parkia biglobosa</i>	5.99	Rare
	<i>Dombeya buetnerii</i>	5.53	Rare
	Asteraceae	11.06	Rare
	Combretaceae	5.53	Rare
	Fabaceae	1.83	Sporadic
	Solanaceae	5.07	Rare

The result of pollen analysis indicated that the honey samples were rich in different pollen types but were in low percentage. Also the result is a reflection that the honey samples are produced from different pollen and nectar plant sources. Pollen grains of *Lannea sp.*, *Elaeisuineensis*, *Phytallus sp.*, *Combretum sp.*, *Tridax procumbens*, *Danielliaoliveri*, *Diospyros sp.*, *Alchornea cordifolia*, *Hymenocardiaacida*, *Tetrapleura tetraptera*, *Microcarpousscabar*, Asteraceae, Fabaceae and Myrtaceae were identified in A1 honey sample.

The pollen analysis revealed that honey sample A2 composed of pollen grains of *Danielliaoliveri*, *Delonix regia*, *Alchornea cordifolia*, *Mallotussubulatus*, *Combretum sp.*, *Terminalia sp.*, *Bridellia sp.* and *Diospyros sp.* The pollen grains of *Lannea sp.*, *Cocos nucifera*, *Elaeisuineensis*, *Tridax procumbens*, *Emilia sp.*, *Vernonia amygdalina*, *Bombax sp.*, *Alchornea cordifolia*, *Phyllanthus sp.*, *Pakia biglobosa*, *Dombeya buetnerii*, Fabaceae, Solanaceae and Combretaceae were recorded in the honey sample A3.



a-b. Asteraceae, c-d. Caesalpinaceae, e. *Combretum sp.*, f. *Danielliaoliveri* g. *Microcarpuscabra*, h. *Elaeisqueensis*, i. *Vitallariaparadoxa*. j. *Parkia biglobosa*, k. *Bombax sp.*, l. *Hymenocardiaacida*, m. Solanaceae, n. *Mallotussabulatus*, o. *Acacia sp.* and p. *Bridellia sp.* Mag. X 400

Fig.1 Photomicrographs of pollen types obtained from A1, A2 and A3 honey samples. a- b Asteraceae, c-d. Caesalpinaceae, e. *Combretum sp.*, f. *Danielliaoliveri* g. *Microcarpuscabra*, h. *Elaeisqueensis*, i. *Vitallariaparadoxa*. j. *Parkia biglobosa*, k. *Bombax sp.*, l. *Hymenocardiaacida*, m. Solanaceae, n. *Mallotussabulatus*, o. *Acacia sp.* and p. *Bridellia sp.* Mag. X 400.

The pollen identified in the honey samples revealed the botanical and geographical origin of the branded honey samples. The results portrayed the true vegetation types of the three honey samples and give credence to the acclaimed labeling of the honey samples by the respective producer. For many consumers, good quality honey is expected to be visually clean and clear.

Honey that contains good pollen appears cloudy hence makes it look unappealing for consumers. Identification and quantifying of pollen in honey is one of the best ways to determine the range of nectar types used by the bees to produce honey and therefore label it correctly based on actual foraging resources. The botanical source may be labeled if the honey is obtained mainly from a particular source. Such honey must also have the organoleptic, physicochemical and microscopic characteristics of the acclaimed origin.

5. CONCLUSION

A complete analysis of honey that involved pollen and physicochemical investigation is needed before any honey sample could be certified as suitable for human use. There is a need for the consumers to request for certificates or analysis results to prove the origin and composition of the honey. Adulterated honey does more harm to their consumers, hence, the need for honey authentication no matter its brand and acclaimed source.

The results of the pollen analysis indicated that these branded honey samples are rich in pollens which confirmed that they are produced from different types of pollen and/nectar plant sources (floral origin). The results also affirmed their geographical origin as claimed by their respective producer. The data obtained from the proximate and physicochemical compositions of the branded honey samples fall within the limit of international standards which prove that these branded honey are of good quality and are suitable for human use.

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