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Abstract:

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Oil Recovery from Palm Kernel Meal Using Subcritical Water Extraction in a Stirred Tank Reactor

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Abstract: Palm kernel meal (PKM) is one of the main byproducts of the oil palm industry. PKM can be obtained as the result of solvent or mechanical extraction of palm kernel oil; in both cases, meal has a remaining oil content that could be recovered. In this work, PKM coming from a mechanical pressing extraction system with an initial oil content between 7 to 8% (wt.) was treated with subcritical water in a batch stirred reactor. To find the proper operational conditions, a three-step experimental process was performed. Extraction temperature, reaction time, particle size and alkaline catalyst usage were selected as process factors. After subcritical extraction, the system was cooled down and depressurized; then oil phase was separated by centrifugation. After extraction, meal was oven-dried at 80 °C. A maximum recovery of 0.034 kg-oil/kg-meal was obtained at 423 K, 720 s and particles smaller than 0.001 m. The experimental procedure showed consistent extraction yields of 40% without modifying the quality of the obtained oil.

Keywords: palm kernel meal; subcritical water extraction; oil recovery; clean separation

1. Introduction

In recent years, interest in improving sustainability of agricultural material processing has been increasing. In addition, environmental phenomena have become significantly unstable, directly impacting agricultural business competitiveness. Palm kernel oil processing hasn’t been sheltered from that reality, so trying to improve oil recovery has become a necessity for this sector. Palm kernel meal (PKM) is a by-product in the palm kernel oil (PKO) production process. PKM is used as a basis for the manufacture of animal feed [1]. Palm kernel oil represents between 46% and 53% of the weight of the kernel [2]. Its chemical composition is different from crude palm oil and it is semi-solid at room temperature [3].

Worldwide, Colombia it is ranked as the fifth-largest palm kernel meal producer; its 148,000 MT in 2016 placed Colombia as the first American maker of this feedstock. The commercial value of this material it is rated at approximately 46.7USD/ton, which is lower than other competitors, such as rice bran or soybean meal. However, pressed palm kernel meal still has an oil content between 7 to 8% wt, this vegetable oil has a market value around 617.15USD/ton.

One option that could become a valid sustainable alternative is subcritical water processing [4]. Subcritical water refers to a region where fluid temperature varies between 373 K and 547 K in a liquid state. At those conditions some properties, like polar moment shift, make water more similar
to conventional organic solvents, making it suitable for lipid extraction and hydrolysis [5], creating a sustainable method to decrease hazardous organic solvent consumption. The thermodynamic properties of water are based on the strength and structure of their hydrogen bonds [6]; which are governed by the relations between water atoms [7]. Therefore, a minimum change in one of the bonds affects the entire volume of water. Particularly at the subcritical region, changes in the resistance of these bonds are reflected in the dielectric constant and heat of vaporization change [6]. At lower temperature, the hydrogen bonds are stronger implying a high dielectric constant. The opposite occurs at high temperatures, since the dielectric constant is decreased [8,9], leading to an increase in the solubility of a normally hydrophobic compound [6].

Oil recovery from previously pressed fibers using subcritical water, using different catalysts, has been investigated in previous studies [10,11]. Also, the complementary use with different solvents was evaluated in [12]. Studies on palm empty fruit bunch liquefaction using alkaline catalysts have also been conducted [13,14]. Subcritical water has also been used for phenolic compound extraction from several oil palm byproducts [15], carotenoids [16], and carbohydrates, such as sugars [17] and cellulose [18] from palm fibers and leaves. This study presents the findings from tests of an experimental batch extraction process of oil from palm kernel meal using subcritical water under different experimental conditions.

2. Materials and Methods

2.1. Raw Material

Mechanically pressed palm kernel meal was collected from PALMACEITE S.A. (Aracataca, Magdalena, Colombia) in 5 kg plastic bags that were stored at approximately 23 °C to prevent lipidic photo-oxidation. Potassium hydroxide (85% of purity) was provided by Riedel de Haen (Wunsstorfer Str. 40, 30926 Seelze, Germany). High performance liquid chromatograph (HPLC) grade hexane was supplied by EMD Millipore corporation (Billerica, MA, USA). Ultra high purity argon gas was used as inert gas for reactor headspace during the extractions and Ultra high purity helium, hydrogen, air and nitrogen, all supplied by The Linde Group (Klosterhofstrasse 1, Munich, Germany) were used as carrier and detection gases during the chromatographic analysis. Distilled water from the experimental laboratory was used during the experiments.

2.2. Meal Preparation

Palm kernel meal was separated in 5 fractions using number 8, 20, 40 and 60 standard American Society for Testing and Materials (ASTM) sieves and a shaker all provided from Pinzuar Ltd.a. (Carrera 108 No. 18-31, Bogota, Colombia). The sieving process produced $2.38 \times 10^{-3} \text{ m}$, $0.84 \times 10^{-3} \text{ m}$, $0.42 \times 10^{-3} \text{ m}$, $0.25 \times 10^{-3} \text{ m}$ and a fine size particle fractions that were labeled and stored separately for further processing.

2.3. Subcritical Water Extraction

Subcritical water extraction was performed in a $2.5 \times 10^{-4} \text{ m}^3$ stirred tank batch reactor Model 3230 (Figure 1) supplied by AMAR Equipments Pvt. Ltd. (6, Parmar Industrial Estate, Kale Marg, Bail Bazaar, Kurla (West) Mumbai–400 070, India). Palm kernel meal, catalyst, and water mixture were added. Argon gas was introduced to replace the air on the reactor headspace and establish a $6 \times 10^5 \text{ Pa}$ base pressure. Reaction temperature and agitation were established using a NEURO100 PPI Controller (Process Precision Instruments, 101 Diamond Industrial Estate, Navghar, Vasai Road (E), Dist. Palghar—101210, Maharashtra, India) and a frequency inverter (Schneider Electric Co., 43-45, bd Franklin Roosevelt 92500 Rueil Malmaison, France) respectively. Pressure was elevated from base level to operational condition by temperature action, and was monitored also using a NEURO100 PPI Controller. After the determined reaction time passed, temperature was decreased using the water cooling system of the vessel. After the system was cooled, vessel was depressurized, and the mixture
was poured in $15 \times 10^{-6}$ m$^3$ centrifuge tubes. Later, the tubes with the samples were centrifuged at $1262 \times g$ for 1200 s in a Thermo Scientific CL-22 bench centrifuge (168 Third Avenue, Waltham, MA, USA 02451). After centrifugation, oil layer was extracted by pipetting and separated for further analysis.

**Figure 1.** Stirred tank batch reactor.

### 2.4. Soxhlet Extraction

Between 0.005 and 0.01 kg of palm kernel meal were mixed with $150 \times 10^{-6}$ m$^3$ of hexane for standard Soxhlet extraction at 343 K for 6 h in order to determine total oil content of the material.

### 2.5. Gas Chromatography

Gas Chromatography with Thermal Conductivity Detector (GC-TCD) was performed with a Hewlett-Packard 5890 series II, a TCD detector (135 mL/min He), using a DB-23 Agilent Tech capillary column with a 0.25 mm internal diameter, 60 m length and 0.25 µm film thickness. The Fatty acids composition was performed by oil conversion to fatty acids methyl esters, according to Cocks and Van Rede [19]. About 50 mg of oil and mixed with 950 µm of n-hexane, then was added 50 µm of CH$_3$NaO, and the mixture was stirred in a vortex mixer and let it rest for 60 min after 1 µL collected supernatant was injected in the GC for 40 min and 30 °C. The Standards used (fatty acids oleic, myristic, lauric, linoleic, linolenic, palmitic, palmitoleic, arachidic, caprylic, caproic, capric and stearic) were an analytic grade (Sigma-Aldrich Inc., St. Louis, MO, USA). Nitrogen was used as the analyte carrier gas.

### 2.6. Extraction Yield Determination

Extracted lipid fractions were separated from the aqueous and solid phases by centrifugation at 5000 min$^{-1}$ for 1500 s. Then, the separated oil was extracted by pipetting. The extracted oil was weighed and the yield was determined by Equation (1).

$$Yield = \frac{m_e}{m_S}$$  \hspace{1cm} (1)
where $m_e$ is the obtained oil mass by subcritical water extraction in g., and $m_s$ is the obtained oil mass by Soxhlet extraction in g.

### 2.7. Experimental Execution

The experiment was designed in stages with independent objectives. Initially, an exploration process was performed to determine the probable experimentation region. For this purpose, $16 \times 10^{-3}$ kg of $600 \mu$m meal were loaded to the reactor with $100 \times 10^{-6}$ m$^3$ of water. The particle size, meal mass and water load amounts were maintained on every run. In the exploration stage, temperature was varied in 10 K steps from 393 to 493 K; reaction time was varied between 300, 600, 1800, 2700 and 5100 s and rotation speed was varied between 400, 600 and 800 min$^{-1}$. Also, at this stage, alkaline catalyst action was evaluated by adding $9 \times 10^{-3}$ kg of KOH (9%wt) for the catalyzed runs. For the second stage, a $2^4$ screening design was applied to determine the significance level of the experimental region factors; in this case, reaction temperature was varied between 418 and 428 K, reaction time from 480 to 720 s, meal mass load from $12 \times 10^{-3}$ to $20 \times 10^{-3}$ kg, and particle size varied between $600 \mu$m (Mesh 18) and $212 \mu$m (Mesh 70). Finally, on the third stage a central composite design was applied, and with the results a regression model was built and a statistical optimization process using Statgraphics Centurion XVI (Statgraphics Technologies, Inc., Virginia, USA) was executed. The regression model obtained was built according to the structure shown in Equation (2).

$$Y = \beta_0 + \sum_{i=1}^{3} \beta_ix_i + \sum_{i=2}^{3} \sum_{j=1}^{2} \beta_{ij}x_ix_j + \sum_{i=1}^{3} \beta_{ii}x_i^2 + \epsilon$$  \hspace{1cm} (2)$$

where $Y$ is the regression model response, $x_i$, $x_j$ correspond to the process factors, $\beta_0$ is the intercept term, $\beta_i$ correspond to the linear terms coefficients, $\beta_{ij}$ are the interactions terms coefficients, $\beta_{ii}$ are the quadratics terms coefficients, and $\epsilon$ is the modeling error term. Model adequacy was tested using lack of fit and Durbin-Watson tests. Also, residual structure was analyzed and standard ($R^2$) and adjusted ($R^2_{adj}$) determination coefficients were evaluated to determine the model degree.

### 3. Results and Discussion

#### 3.1. Experimental Region Exploration

Phase I exploration resulted in a qualitative evaluation of lipid phases obtained with different treatments, with and without catalyst, looking to determine on which temperature region extraction seemed to appear. It was found that at a temperature below 413.15 K a lipid phase was not appreciable, and at a temperature above 433.15 K a burnt lipid phase was appreciable.

An increase in the rotation speed improved the mass transfer solvent-solute, due to the film thickness decrease around the particles. However, this parameter was not significant through the range studied. Therefore, this parameter was set in 600 min$^{-1}$ during the experiments to avoid agglomeration and sedimentation of particles, which would cause mass transfer decrease.

Reaction time was set at 600 s and experiments were done with and without catalyst; initial yields obtained were 35% without catalyst and 15% with KOH as catalyst, also KOH-treated samples had a significant change in smell which could imply quality failures on reclaimed products. Figure 2 show obtained lipid phases.
Obtained lipid profiles for the initial extracts by gas chromatography analysis are shown in Figure 3.

In Figure 3 it can be seen that the lipid phase obtained without basic catalyst was richer in lauric acid, which is desirable for applications in the cosmetic industry.

**Figure 2.** Extracted lipid phases.

**Figure 3.** Fatty acid profiles for samples without catalyst, with catalyst and standard (Error bars represent 1.96%).
3.2. Screening Experimentation

Due to the fatty acid profiles obtained, screening was done without catalyst, a factorial design \((2^k)\) was performed for this purpose with process temperature, reaction time, meal mass and particle size as experimental factors, Table 1 shows experimentation order and results.

| Run | Temperature (K) | Time (s) | Meal Mass \((\times 10^{-3} \text{ kg})\) | Particle Size | Yield (%wt) |
|-----|-----------------|----------|---------------------------------|--------------|-------------|
| 1   | 428.15          | 720      | 12                              | A            | 0           |
| 2   | 418.15          | 720      | 20                              | A            | 0           |
| 3   | 428.15          | 480      | 12                              | B            | 8.22        |
| 4   | 428.15          | 480      | 20                              | A            | 1.85        |
| 5   | 418.15          | 480      | 20                              | B            | 16.19       |
| 6   | 418.15          | 720      | 12                              | B            | 10.87       |
| 7   | 428.15          | 720      | 20                              | B            | 24.55       |
| 8   | 418.15          | 480      | 12                              | A            | 5.44        |
| 9   | 428.15          | 480      | 20                              | B            | 7.4         |
| 10  | 418.15          | 720      | 12                              | A            | 3.82        |
| 11  | 418.15          | 720      | 20                              | B            | 24.56       |
| 12  | 418.15          | 480      | 20                              | A            | 2.47        |
| 13  | 428.15          | 480      | 12                              | A            | 2.80        |
| 14  | 428.15          | 720      | 20                              | A            | 8.51        |
| 15  | 418.15          | 480      | 12                              | B            | 13.03       |
| 16  | 428.15          | 720      | 12                              | B            | 13.08       |

A: 2.36mm > particle diameter > 1mm. B: particle diameter < 1mm.

Particle size was managed as a qualitative factor where A and B mean above and through mesh 18. System pressure varied between \(7.2 \times 10^5 \text{ Pa}\) to \(14.6 \times 10^5 \text{ Pa}\) depending on the temperature operational point of every run.

3.3. Statistical Analysis

Analysis of variance (ANOVA) was used to determine factor and interaction effect significance on the extraction yield; Table 2 shows analysis results for individual factors.

| Source       | Sum of Squares | Degrees of Freedom | Mean Square | F-Ratio | p-Value |
|--------------|----------------|--------------------|-------------|---------|---------|
| Temperature  | 6.21           | 1                  | 6.21        | 0.26    | 0.6205  |
| Time         | 48.96          | 1                  | 48.96       | 2.05    | 0.1804  |
| Meal Mass    | 49.95          | 1                  | 49.95       | 2.09    | 0.1764  |
| Size         | 540.68         | 1                  | 540.68      | 22.60   | 0.0006  |
| Total Error  | 263.22         | 11                 | 23.93       | 0.26    | 0.6205  |
| Total        | 909.02         | 15                 |             |         |         |

From Table 2 it can be inferred that particle size has a higher significant impact on the extraction yield compared to the other process factors. In addition, process temperature showed the least impact over the response variable mainly due to the small range of variation applied. However, the range size was selected according to the exploring experimentation, where it was shown that above 443.15 K, the lipid phase obtained had physical changes that affected the final product quality. The experimental data was also evaluated looking for autocorrelation signs, the Durbin-Watson statistic calculated was 1.96 with a \(p\)-value of 0.60, which confirmed the absence of correlation between the data. Likewise, homoscedasticity was checked, comparing residual values dispersion ranges obtained at the different factor levels. Due, to its poor significance level, process temperature was discarded from complete
analysis; Table 3 illustrates new ANOVA without temperature and including the interaction between the remaining experimental factors.

Table 3. ANOVA results for the screening experiment excluding reaction temperature from the sources of variation.

| Source          | Sum of Squares | Degrees of Freedom | Mean Square | F-Ratio | p-Value |
|-----------------|----------------|--------------------|-------------|---------|---------|
| B = Reaction Time| 48.96          | 1                  | 48.96       | 3.12    | 0.0978  |
| C = Meal Mass   | 49.95          | 1                  | 49.95       | 3.18    | 0.0948  |
| D = Particle Size| 540.68         | 1                  | 540.68      | 34.42   | 0.0000  |
| BC              | 61.74          | 1                  | 61.74       | 3.93    | 0.0661  |
| CD              | 50.59          | 1                  | 50.59       | 3.22    | 0.0929  |
| Total Error     | 157.1          | 10                 | 15.71       |         |         |
| Total           | 909.02         | 15                 |             |         |         |

Table 3 confirms significant influence of particle size over the extraction yield, but also shows that BC and BC interactions have a higher impact on the yield compared even with the single factor effect. Therefore, they were included on detailed experimentation.

3.4. Response Surface Experimentation

A $2^3$ rotatable central composite design with reaction time, meal mass and particle size as codified experimental factors with 4 center points was done to determine extraction yield optimum levels. In this case particle size wasn’t taken as a qualitative factor, instead several milling diameters were used. Table 4 shows the experiment runs and their results.

Table 4. Results for the screening experiment excluding reaction temperature from the sources of variation.

| Run | Particle Size (mm) | Time (min) | Meal Mass (g) | Obtained Yield (%wt) |
|-----|--------------------|------------|---------------|----------------------|
| 1   | −1                 | −1         | 1             | 21.99                |
| 2   | −1                 | −1         | −1            | 18.60                |
| 3   | 1                  | −1         | −1            | 8.08                 |
| 4   | 1                  | 1          | 1             | 1.41                 |
| 5   | 0                  | 1.68179    | 0             | 7.16                 |
| 6   | −1                 | 1          | −1            | 18.20                |
| 7   | 0                  | 0          | 0             | 13.52                |
| 8   | 0                  | 0          | −1.68179      | 7.95                 |
| 9   | 0                  | 0          | 0             | 13.18                |
| 10  | 1                  | −1         | 1             | 2.83                 |
| 11  | 1                  | 1          | −1            | 2.62                 |
| 12  | 0                  | 0          | 0             | 3.63                 |
| 13  | 0                  | 0          | 0             | 8.51                 |
| 14  | −1.68179           | 0          | 0             | 37.97                |
| 15  | −1                 | 1          | 1             | 30.59                |
| 16  | 0                  | −1.68179   | 0             | 7.17                 |
| 17  | 1.68179            | 0          | 0             | 0.43                 |
| 18  | 0                  | 0          | 1.68179       | 12.74                |

Table 5 shows that particle size has a marked inverse relation with extraction yield. Meanwhile reaction time and meal mass behavior as individual factors has a more random result.

3.5. Response Surface Statistical Analysis

After performing an initial analysis of variance, reaction time showed no significant effect on the extraction yield in this experiment; Therefore, it was excluded from the final analysis. Tables 5 and 6 shows the initial ANOVA and ANOVA with only the remaining factors and interactions, respectively.
Table 5. Initial ANOVA for the response surface experiment.

| Source      | Sum of Squares | Degrees of Freedom | Mean Square | F-Ratio | p-Value |
|-------------|----------------|--------------------|-------------|---------|---------|
| A = Particle Size | 1386.68       | 1                  | 1386.68     | 127.11  | 0.0000  |
| B = Time    | 0.12           | 1                  | 0.12        | 0.01    | 0.9172  |
| C = Meal Mass | 22.00          | 1                  | 22.00       | 2.17    | 0.1789  |
| AA          | 154.69         | 1                  | 154.69      | 15.25   | 0.0045  |
| AB          | 28.27          | 1                  | 28.27       | 2.79    | 0.1335  |
| AC          | 61.61          | 1                  | 61.61       | 6.07    | 0.0390  |
| BB          | 7.27           | 1                  | 7.27        | 0.72    | 0.4219  |
| BC          | 21.39          | 1                  | 21.39       | 2.11    | 0.1845  |
| CC          | 1.70           | 1                  | 1.70        | 0.17    | 0.6932  |
| Total Error | 81.13          | 8                  | 10.14       |         |         |
| Total       | 1764.86        | 17                 |             |         |         |

Table 5 shows that p-value for factor C was higher than 0.05. Nevertheless, this source was not excluded from ANOVA because the interaction AC was significant (p-value = 0.0335). Because of this, C factor cannot be excluded from the statistical analysis. During the statistical analysis, non-significant factors were removed, obtaining the results as shown in Table 6.

Table 6. ANOVA for the response surface experiment.

| Source      | Sum of Squares | Degrees of Freedom | Mean Square | F-Ratio | p-Value |
|-------------|----------------|--------------------|-------------|---------|---------|
| A = Particle Size | 1386.68       | 1                  | 1386.68     | 98.59   | 0.0000  |
| C = Meal Mass | 22.00          | 1                  | 22.00       | 1.56    | 0.2425  |
| AA          | 173.32         | 1                  | 173.32      | 12.32   | 0.0066  |
| AC          | 61.61          | 1                  | 61.61       | 4.38    | 0.0659  |
| Lack-of-fit | 15.23          | 4                  | 3.81        | 0.27    | 0.8896  |
| Total Error | 126.59         | 9                  | 14.06       |         |         |
| Total       | 1785.43        | 17                 |             |         |         |

The AC interaction effect is very close to the significance level, therefore it was kept in the model. Also, the AC interaction is significant in the initial ANOVA. The diagnostics were all positive showing that all assumptions were accomplished with the following results. The observed lack of fit was non-significant. To check experiment validity, residual behavior was analyzed, Figure 4 shows a normal probability plot of the residuals obtained for the response surface experiment.

![Normal Probability Plot](image-url)

Figure 4. Normal probability plot of obtained residuals.
Figure 4 does not present evidence of any abnormal behavior from the residuals obtained, this could validate the normality assumption in the response surface experiment. The homoscedasticity check was validated comparing residual value dispersion ranges obtained at the different factor levels.

As an autocorrelation check, the Durbin-Watson statistic was calculated; obtained d-statistic was 1.51035 with a p-value of 0.1822 which confirmed the absence of correlation between residuals.

3.6. Regression Model

With the data from the response surface experiment, a regression model was obtained, due to the significant interaction, a quadratic structure was used. The R² adj value of the model is 89.613%. Equation (3) shows obtained expression.

\[
Yield = 9.32972 - 10.0766S + 1.26938M + 3.56488S^2 - 2.775SM
\]  

(3)

where \( S \) is the particle size in mm, and \( M \) is the meal mass in g.

With the obtained expression, a response surface was built where the optimal yield region was identified. Figure 5 shows the top view of the surface.

In Figure 5 it can be seen that the maximum yield region is located at 0.212 mm particle size and 22.7 g palm kernel meal mass, at a temperature of 423.15 K and reaction time 640.2 s. A computational optimization process using STATGRAPHICS Centurion XVI was applied to obtain the recommended operation point. According to the optimization exercise, the optimum obtained extraction yield will be 40.69%.

4. Conclusions

This work shows that it is feasible to perform effective oil extraction from palm kernel meal at moderate temperature and pressure conditions. The optimal region was found to be near 150 °C and 7 bars, which is significantly lower than results in other studies, where the process took place above 200 °C and 20 bars. That has a positive impact both in energy usage and preventing fatty acid degradation.

Subcritical water extraction can become a viable option to efficiently recover oil from palm kernel meal without compromising meal quality, a maximum recovery of 0.034 kg oil/kg meal was obtained.
at 423 K in a 720-s batch, using particles with a size of $0.212 \times 10^{-3}$ m. In addition, using KOH as a catalyst led to no improvement in extraction yield, but had an effect on fatty acid profile.

A lower fatty acid content reduced the risk of palm kernel meal rancidity, which allows increased storage time. Also, the lipid phase without catalyst was 6% richer in lauric acid than present palm kernel oil, and the residual palm kernel meal still had a remaining average oil content of 3.5%, which is desirable for the cosmetic and animal feed industries.

Sub-critical water extraction presents no problems with a residual content of toxic solvents in the finished product, and the solvent removal after extraction is no longer necessary, leading to energy savings.

In this work, the cost of the process is not estimated, but operations such as grinding can be expensive, especially with lignocellulosic materials, which can turn this into a negative factor for the process.

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