ESTIMATION OF THERMOPHYSICAL PROPERTIES OF VEGETABLE OILS FOR CREATION DATABASE TO ENGINEERING CALCULATIONS

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I. INTRODUCTION

The decline in available oil reserves and stricter environmental regulations has motivated global interest in studies for the discovery and improvement of renewable energy sources, especially biofuels.

Vegetable oils and fats animal’s types used in the manufacture of biofuel, can cause changes in their properties. In other words, biodiesel can present different properties by the chemical composition of raw materials, for example, a low performance in relation to the properties of cold flow [1].

The literature reveals that there is a large volume of researches conducted in the design of processes and manufacture of biofuels from vegetable oils, but the experimental data available for thermodynamic properties of vegetable oils and/or predict methods are limited [2-6].

The scarcity of data on the thermophysical properties of biodiesel components and vegetable oils can be related to the difficulty in obtaining them in pure form and in the performance of experiments, as these substances can degrade under severe conditions [7], [8]. However, vegetable oils aren’t composed only by a triacylglycerol. Oils and fats can contain dozens of different triacylglycerols (TAG), as tripalmitin, tristearin and triolein.
Diacylglycerols (DAG), monoacylglycerols (MAG) and free fatty acids (FA) are also components of vegetable oils and have few properties reported in the databases, as well as methyl and ethyl esters [4], [9-12].

Group Contribution (CG) methods have been used in recent years to estimate some physicochemical properties of organic substances, such as normal boiling point, critical properties and acentric factors [3], [11], [13]. The GC considers that the properties of molecules depend on the nature of atoms and the types of chemical bonds within molecules [14]. In addition to GC, methods with the principle of contributor fragments have been used for calculations that consider that the properties of molecules are established from the contributions of their functional groups [8]. These methodologies need only small amounts of information about the substances in order to obtain a reliable estimate. Although there are many methods in the literature, each of them has built-in premises and practical limits that must be applied. Therefore, selecting the most appropriate models for predicting thermophysical properties is extremely important for achieving realistic results in a process simulation.

II. METHODOLOGY

II.1 MODELING OF PREDICTING THERMOPHYSICAL PROPERTIES

For the development of property estimates, electronic spreadsheets were created in MS-Excel® with equation sequences for the calculations of the methods.

For the application of the methods of estimating properties, the molecules of the compounds of interest (TAGs, DAGs, MAGs and FA) were drawn and, then, the functional groups identified and quantified.

Example: A molecule of triacylglycerol PLO, formed by the junction of palmitic fatty acids (P), linoleic (L) and oleic (O) in a glycerol molecule. Its chemical formula is represented by \( \text{C}_{33}\text{H}_{100}\text{O}_{6} \). Figure 1 represents the triacylglycerol PLO molecule.

![Figure 1: Chemical structure of triacylglycerol PLO. Source: Authors, (2020).](Image)

The groups present in the molecular structure represent in Figure 1 were counted for the methods of contribution of groups showed in Table 1, JOBACK [15] and GANI [16].

| JOBACK groups | Score | GANI groups | Score |
|---------------|-------|-------------|-------|
| CH\(_3\)      | 3     | CH\(_3\)    | 3     |
| >CH\(_2\)     | 42    | >CH\(_2\)   | 40    |
| >CH-          | 4     | >CH-        | 1     |
| -COO-         | 3     | -CH\(_2\)-COO-(C) | 2 |
| =CH-          | 3     | -CH-        | 1     |

Source: Authors, (2020).

Each of these groups has a contribution value depending on the property to be calculated and method used. Thus, it’s possible to use the equations of the methods safely to predict the properties.

Equations necessary for the calculations of GC methods and fragments approach are described below.

**Normal boiling temperature** (\( T_b \)): \( T_b \) is the temperature at which the vapor pressure equals the external or atmospheric pressure.

Developed an advanced method of group contribution based on the groups of the thermodynamic model *Universal Functional Activity Coefficient* (UNIFAC) [16-17]. In addition, the authors added the so-called “second order” contributions that allow differentiating molecules from isomers, molecules that have the same groups, located together or not, resonance structures, among others [8].

The equations of the JOBACK and GANI models for \( T_b \) estimation are represented by equations 1 and 2.

**JOBACK:**

\[
T_b = 198 + \sum_k N_k (T_{bk})
\]

(1)

**GANI:**

\[
T_b = 204.359 \times \ln \left( \sum_k N_k T_{bk} \right)
\]

(2)

where \( N_k \) represents the number of groups of type \( k \) and \( T_{bk} \) the contribution to the normal boiling temperature of groups \( k \).

The semi-empirical approach of [18], [19] was also used to calculate of boiling temperature of TAGs, DAGs and MAGs is implemented the basis of fragments of the constituents. A relationship between the boiling temperature of each of the fragments of the TAGs (fatty acids and glycerol) and pressure was applied. Therefore, the experimental data of [20], [21] on the temperature of each of the fragments were adjusted to Equation 3.

**ZONG:**

\[
T_b^a = a \times \ln(P) + b
\]

(3)

where \( a \) and \( b \) are the adjustment parameters of each fragment. \( T_b^a \) is the boiling temperature of fragment \( a \). \( P \) is the vapor pressure (mmHg). Knowing the value of \( T_b^a \) for each of the fragments, it was possible to calculate the value of \( T_b^\text{TAG} \) using Equation 4:

**\( T_b^\text{TAG} \):**

\[
T_b^\text{TAG} = \frac{\sum N_{\text{Frag.}a} \times T_b^a}{3}
\]

(4)
T_b^{TAG} represents normal boiling temperature of triacylglycerol (K) and N_{frag} number of fragments of a.

**Critical properties** (P_c, V_c, T_c and Z_c): Pressure (P_c), volume (V_c), temperature (T_c) and compressibility factor (Z_c). Critical are parameters used in many volumetric, thermodynamic and transport correlations based on the theory of corresponding states to estimate gas and liquid properties.

According to [8], the experimental determination of these properties is difficult, as chemical degradation can occur at high temperatures, especially for long chain components.

The critical point of glycerol was defined by [22] through an unconventional methodology called pulse-heating, being the only experimental measure available in the literature. The critical temperature obtained was 850 K and the critical pressure was 75 bar. As glycerol begins to undergo thermal degradation from 510 temperature obtained was 850 K and the critical pressure was 75 bar. According to [8], the experimental determination of these properties is difficult, as chemical degradation can occur at high temperatures, especially for long chain components.

The estimation methods evaluated in this study were: [15], [16], [23], [24]. The first method requires only information on the structure of the compound, while the other methods require values of molar mass and boiling temperature. The methods are represented by Equation 5 to Equation 16.

**JOBACK:**

\[
T_c = T_b[0.584 + 0.965 \sum_k N_k(T_{ck}) - \left( \sum_k N_k(T_{ck}) \right)^2]^{-1}
\]  

\[
P_c = [0.113 + 0.0032 N_{at} - \sum_k N_k(P_{ck})]^{-2}
\]  

\[
V_c = 17.5 + \sum_k N_k(V_{ck})
\]

**GANI:**

\[
T_c = 181.28 \times \ln(\sum_k N_k T_{ck})
\]  

\[
P_c = [\sum_k N_k P_{ck} + 0.10022]^{-2} + 1.3705
\]  

\[
V_c = \left( \sum_k N_k(V_{ck}) - 0.00435 \right) \times 1000.0
\]

**AMBROSE:**

\[
T_c = T_b \left[ 1 + \left( 1.242 + \sum_k N_k(T_{ck}) \right)^{-1} \right]
\]  

\[
P_c = M_M \left[ 0.339 + \sum_k N_k(T_{ck}) \right]^{-2}
\]

**LYDERSEN:**

\[
T_c = \frac{0.567 + \sum_k N_k(T_{ck}) - \left( \sum_k N_k(T_{ck}) \right)^2}{\sum_k N_k(T_{ck})}
\]

\[
P_c = \frac{M_M}{(0.34 + \sum_k N_k(P_{ck})^2}
\]

\[
V_c = 40 + \sum_k N_k(V_{ck})
\]

\[
\omega = -\log \left( \frac{P_{vp}}{T_c^{0.7}} \right) - 1.0
\]

where P_{vp} is vapor pressure, T_c critical temperature and P_c critical pressure.

The definition of the equation (Equation 17) arose because monatomic gases (Air, K-r, Xe) have \( \omega \approx 0 \), except for noble gases (He, Ne) and some others (e.g. Rn). All other species have positive values up to 1.5. To obtain \( \omega \) values is necessary to know the constants T_c, P_c and the property P_{vp} at reduced temperature, T/T_c = 0.7.

Pitzer rule (PITZER) and Kesler-Lee (LEE) were selected in this work to determined \( \omega \). Equation 18 was presented by [25] for \( \omega \) calculation:

\[
Z_c = 0.291 - 0.080 \omega
\]

According to [8], Equation 18 can be used to predict the acentric factor of strongly polar substances and associative substances. However, the authors indicate that if the critical parameters were not adequately estimated for the substance there isn’t guarantee of the accuracy of the desired property.

A set of equations was developed by [26] to evaluate properties of chemical compounds such as molecular mass, normal boiling temperature, critical pressure, critical temperature and acentric factor. For the latter, the following correlations were developed (Equation 19 and Equation 20). Such models depend on the reduced boiling temperature (T_{br} = T_b/T_c) and the characterization factor of Watson (K_w).

\[
\omega = -7.904 + 0.1352K_w - 0.007465K_w^2 + 8.359T_{br}
\]

\[
+ \frac{1.408 - 0.01063K_w}{T_{br}}
\]

\[
(19)
\]
For $T_{br} < 0.8$:

$$\omega = \frac{-\ln P_c - 5.92714 + 5.0964 + 12.8862 \ln T_{br} - 0.169347 \frac{\rho_{br}}{T_{br}}}{15.2518 - 15.2518 \ln T_{br} + 0.435777 \frac{\rho_{br}}{T_{br}}}$$ (20)

The Watson factor can be defined by Equation 21:

$$K_w = \frac{MT^1}{d_{15\degree C}}$$ (21)

where $MT_b$ is the average of the normal boiling temperature given in °R and $d$ is the relative density measured at 15°C.

**Enthalpy formation and Gibbs free energy of formation in the standard state:** The knowledge of standard enthalpy of formation ($\Delta H^0_f$) is important to provide the enthalpy reaction for the formation of a mol of a substance from the chemical elements that constitute it, in its reference states.

The Gibbs free energy formation ($G^0_f$) is defined as Gibbs’ energy reaction for the formation of a mol of a substance from the chemical elements that constitute it in its reference states. According to [27], Gibbs free energy can be obtained from calorimetric data (enthalpy and entropy, from calorific capacities), equilibrium constants in chemical reactions, electrochemical measurements and spectroscopy data, when in the gas phase.

Rarely values for the calculation of the reaction Gibbs energy are found. Methods of contributions from groups such as [15], [28], [16] and others, have been of great help in predicting reliable estimates of values of formation enthalpy, absolute entropy and Gibbs free energies.

Groups contribution to $G_f$ (298.15 K) and $H_f$ (298.15 K) were obtained by [16] through equations (Equation 22 and Equation 23).

**GANI:**

$$H^0_f = 10.835 + \sum N_k(H_{fk})$$ (22)

$$G^0_f = -14.828 + \sum N_k(G_{fk})$$ (23)

$N_k$ represents the number of groups and $H_a$ and $G_a$ the group’s contribution to enthalpy and Gibbs energy, respectively.

The method of [28] can also be used to estimate formation enthalpy and absolute entropy of chemical compounds in the gas phase, considering the standard state. In this method, contributions are given only to atoms with valence greater than one. Groups are linked to key atoms with specific nomenclature (Equation 24, Equation 25, and Equation 26). For example, group C-(-C)(H)$_3$ is a reference to the carbon atom (key atom) bound to another carbon and three hydrogens.

**BENSON:**

$$\Delta H^0_f(298,15K) = \sum N_k(\Delta H_{fk}^0)$$ (24)

$$\Delta G^0_f(298,15K) = \Delta H^0_f(298,15K) - 298,15\Delta S^0_f$$ (25)

$$\Delta S^0_f(298,15K) = \sum N_k(\Delta S_{fk}^0) - RT\ln \sigma + l\eta$$ (26)

where $\sigma$ is the symmetry of the molecule and $\eta$ the number of isomers. R is the universal constant of gases.

The Joback and Reid method [15] is an extension of the Lydersen method [24]. It's a method that uses additive contributions and no contribution to interactions between groups. According to the authors, it is not a high-precision method, but works well for many chemical species. Among these properties, those of interest for this work are Gibbs free energy estimates of standard formation and enthalpy of standard formation. Equation 27 and Equation 28 are used by this method to estimate these two properties.

**JOBACK:**

$$H^0_f = 68.29 + \sum N_k(H_{fk})$$ (27)

$$G^0_f = 53.88 + \sum N_k(G_{fk})$$ (28)

$N_k$ represents the number of groups, $H_a$ contribution to enthalpy formation and $G_a$ contribution to Gibbs free energy of formation of groups k.

The highlight the similarity of Equation 22 and Equation 27, for the calculation of the enthalpy of formation in the GANI and JOBACK methods, respectively. The main difference in these equations lies in the identification of group k. It’s known that the GANI method considers second-order interactions, consequently, the groups identified by this method may be different from that obtained by JOBACK. Similarly, it happens in Equation 23 and Equation 28 for Gibbs energy calculation.

**Vapor pressure (P$_{vap}$):** The fragment-based method [19] and the Clausius-Clapeyron equation were applied to estimate vapor pressures of TAGs using Equation 29. As it wasn’t possible to find experimental data for vapor pressures of unsaturated TAGs, then was considered that the chains of saturated and unsaturated fatty acids have identical vapor pressures when they have the same number of carbon atoms for the fragment approach.

**ZONG:**

$$\log P(T) = \frac{-\Delta G^0_{vap}}{R\theta ln10} + \frac{\Delta H^0_{vap}}{R\theta ln10} \left( \frac{1}{\theta} - \frac{1}{T} \right)$$ (29)

where $P$ is the vapor pressure (Pa), $T$ is the absolute temperature (K), R is the constant of the gases, $\theta$ is the reference temperature (θ=298.15 K), $\Delta H^0_{vap}$ is the vaporization enthalpy at the reference temperature $\theta$, and $\Delta G^0_{vap}$ is the Gibbs energy of vaporization at the reference temperature $\theta$.

A group contribution method was proposed to estimate vapor pressure and heats of vaporization as a function of the temperature of organic liquids found in the oil and biofuel industries [29]. The regression of the parameters was based on an extensive database, composed of fatty acids, esters of methyl, ethyl, propyl and butyl, tri-, di- and monoacylglycerols and hydrocarbons. This methodology improved the predictions obtained by [18] method, due to the inclusion of new experimental data of esters and acylglycerols (in addition to hydrocarbons) and critical points. Like this, the equation 30 was used to calculated of vapor pressure (P$_{vap}$). The pressure is in Pa and temperature in K.
A linear equation to estimate heat of vaporization was used by [25]. $\Delta H_{\text{vap}}$, as a function of reduced temperature ($T_r$) and acentric factor $\omega$. This expression was derived in an analytical equation [31], through an approximation of the correlation of 0.6 < $T_r$ < 1.0 (Equation 35).

$$\frac{\Delta H_{\text{vap}}}{R T_c} = 7.08 (1 - T_r)^{0.354} + 10.95 \omega (1 - T_r)^{0.456} \quad (35)$$

[32] applied another group contribution method to estimate vaporization heat for several substances as a function of reduced temperature (Equation 36).

$$\Delta H_{\text{vap}} = A (1 - T_r)^{\alpha} \exp(-\alpha T_r) \quad (36)$$

$\Delta H_{\text{vap}}$ represents vaporization heat, $T_r$ at reduced temperature, and $A$ and $\alpha$ are contributing parameters of the group. [19] used Equation 29 to predict vaporization enthalpy. The vaporization heat appears from the correlation of vapor pressure by Antoine's equation. Temperature dependence with vaporization heat was not evaluated.

### Density ($\rho$): Rackett's modified equation was used to estimate the density of liquid vegetable oils [33]. First, the density of the liquid mixture of free fatty acids was estimated and then a correction factor was added to describe the shape of the TAGs (Equation 37). No correction factors were presented for DAGs and MAGs.

$$\rho = \frac{\sum x_i MM_i}{R \left( \frac{\sum x_i T_i}{P_{ci}} \right) \left( \sum x_i Z_{RAi} \right)^{1 + (1 - T_r)^2}} + F_c \quad (37)$$

where $x_i$, $MM_i$, $T_{ci}$, $P_{ci}$, and $Z_{RAi}$ are molar fractions, molecular mass, critical temperature, critical pressure and Rackett parameter, respectively, of each fatty acid $i$. $T_r$ is the reduced temperature. The density is given in kg/m$^3$.

The fragment-based approach was proposed by [19] to estimate the molar volume of each fragment with dependence on temperature, and then estimated the molar volume of the liquid based on the composition and contribution of each fragment (Equation 38).

$$v^L = \sum_A N_{\text{frag}A} v_A^L(T) \quad (38)$$

$v_A^L$ is the molar volume of the liquid of each fragment and $N_{\text{frag}A}$ is the number of fragments A in a component.

[34] extended the method of group contribution developed by [35] to predict the densities of pure liquid compounds (Equation 39).

$$\rho = \frac{MM}{v} = \frac{MM}{\sum n_i \Delta v_i} \quad (39)$$

where MM molecular mass and $v$ molar volume.

### Heat capacity ($c_p$): The fragment-based method to estimate the heat capacity of TAGs, DAGs and MAGs was described by [19]. Fragments of FAs and MAGs were expressed as linear temperature-dependent equations (Equation 40). Unsaturated fatty acid fragments were assumed to be identical.

$$c_p = c_{p,\text{fr}} + c_{p,\text{r}} \quad (40)$$
\[ C_p^L = \sum_A N_{frag,A} C_{p,A}(T) \]  

where \( N_{frag,A} \) the number of fragments A in the component and \( C_{p,A}^{A} \) is the contribution of the calorific capacity of the fragment in component A (J/kmol K).

Ceriani et al. [30] extended their method of contribution of groups previously used for the prediction of vapor pressure of organic liquids to develop a heat capacity model with the same set of functional groups in addition to a new linear relationship as a function of the contribution of the group (Equation 41).

\[ C_p^{li} = \sum_k N_k (A_k + B_k T) \]  

\( N_k \) represents the number of K groups in the molecule, \( A_k \) and \( B_k \) are obtained from regression parameters.

Heat capacities for TAGs and vegetable oils was estimated by [36], primarily by Rowlinson-Bondi equation [37][38]. Equation 42 estimated the heat capacities of pure fatty acids. They then added a correction factor, based on study of [33], to predict the density of triacylglycerols. Resulting in an expression with dependence on the molecular mass of the oil (Equation 43).

\[ \frac{c_p^{AG} - c_p^{AG,0}}{R} = 1.45 + 0.45(1 - T_r)^{-1} + 0.25\omega[17.11 + 25.2(1 - T_r)^{1/3} T_r^{-1} + 1.742(1 - T_r)^{-1}] \]  

\( c_p^{AG} \) is the liquid specific heat of fatty acids, \( c_p^{AG,0} \) the specific heat of the ideal gas, \( R \) the universal constant of gases, \( T_r \) the reduced temperature and \( \omega \) the acentric factor.

\[ C_p(estimated) = C_p(AG) + F_c \]  

\( F_c \) correction factor dependent on the molecular mass of the oil.

II.2 CHARACTERIZATION OF VEGETABLE OILS

To evaluate the methods of prediction of properties correctly, one should make the choice of consistent data, that is, the composition of the studied oil sample should be as real as possible, in order to ensure that the conclusions are effective.

For soybean, sunflower and palm oil, which were used as raw material in this work, the composition used in triacylglycerols was that obtained by [39], [40], [41], respectively. These authors determined the oil profile in TAGs by high performance liquid chromatography (HPLC).

For macauba pulp oil, as the composition in TAGs, DAGs and MAGs was not found in the literature, two characterization methodologies were proposed. The first is an approach in which oil is represented by a mixture of Mixed TAG, either by (A) high resolution gas chromatography or (B) computational methodology proposed by [42]. The second approach characterizes vegetable oil as a mixture of fatty acids.

Approach 1:

(A) High Performance Gas Chromatography Analysis (GC)

The sample of macauba pulp oil was diluted in toluene at the final concentration of 0.7%. GC analysis was performed without derivatization on a thermo stabilized fused silica capillary column of TG-5 HT from Thermo brand of 15 meters x 0.25 mm x 0.10 micrometers. The analysis was performed with hydrogen flow from 1.5 mL to 50°C under constant pressure. The initial temperature of the column was 50°C, with a temperature increase schedule of 15°C/minute to 180°C, with ramps of 7°C/minute up to 230°C and up to 350°C with programming of 10°C/min, remaining at this temperature for another 25 minutes. The injector was maintained at 320°C, in the flow division mode of 1:50 and 1 microliter of solution was injected. The detector was maintained at 380°C.

Internal normalization was performed to quantify triacylglycerols. Standards of palmitic acid, linoleic, monoolein, monopalmitin, diolein, dipalmitin, tripalmitin and triolein from Sigma and Nu Chek brands were used for identification. They were dissolved in toluene PA.

(B) Computational Analysis

The computer program was assembled in Microsoft Excel based on equations to calculate the molar percentage of triacylglycerols according to the distribution and molar composition of fatty acids present in vegetable oil.

For the characterization of the pulp oil of the macauba to be performed, the fatty acid profile was necessary. Thus, the mass composition obtained by [43].

For the construction of the oil profile studied in TAGs, 33 triacylglycerol molecules with 99 fatty acid radicals were considered, and the number of each radical in these 33 molecules was proportional to the percentage shown in composition [43]. These FA radicals were randomly distributed among the 33 molecules of TAGs.

Thus, the thermophysical properties were estimated for each mixed TAG by applying the specific prediction method and then the mixing rule was used to estimate the oil property according to the composition of mixed triacylglycerols.

Approach 2: The second approach is based on the characterization of vegetable oil as being a mixture of fatty acids. That is, the fatty acid composition of the oil is used to determine the properties of interest. This methodology has been the most used by researchers, because there is a greater amount of experimental data on the properties of pure fatty acids.

III. RESULTS AND DISCUSSIONS

Three statistical parameters have been used to determine the accuracy of the performed predictions: absolute deviation (AD), relative deviation (%RD) and absolute average relative deviation (%AARD):

\[ \text{Absolute deviation (AD)} = |x_{\text{exp,i}} - x_{\text{calc,i}}| \]  

\[ \%RD = \frac{|x_{\text{exp,i}} - x_{\text{calc,i}}|}{X_{\text{exp,i}}} \times 100 \]  

\[ \%AARD = \frac{\sum |x_{\text{exp,i}} - x_{\text{calc,i}}|}{n} \times 100 \]  

X is the property to be evaluated, n is the number of experimental data, and the subscripts “exp” and “calc” indicate the experimental and calculated property, respectively.
III.1 ESTIMATION OF THERMODYNAMIC PROPERTIES

The estimated properties for TAGs, DAGs, MAGs and FAs, which will be discussed below, are available in the supplementary materials.

Normal boiling temperature (Tb): First, normal boiling temperatures for triacylglycerols were estimated, whose experimental data were found in the literature. The contribution methods of JOBACK and GANI groups, and the method based on ZONG fragments were evaluated and the estimated data are represented in Figure 2.

![Figure 2: Estimated normal boiling temperatures (K) for PPP (tripalmitin), SSS (tristearin) and OOO (triolein). Source: Authors, (2020).](image)

Figure 2 shows the T_b estimated for three simple triacylglycerols, PPP, SSS and OOO, compared to experimental data obtained in [44], [45] [38]. The results show that the T_b values calculated by the JOBACK method have high deviations (AARD>100%). The lowest absolute average relative deviation was reached when the ZONG method was used (AARD=1.9%), followed by 21% for GANI. Thus, the ZONG method was used in this work to calculate T_b for vegetable oils, in the TAG approach.

Since no experimental data from The MAGs and DAGs were found in the literature, the method adopted for the determination of T_b was the ZONG method, which observed lower deviations for the TAGs analyzed, as shown above.

The relative deviations of the estimate of T_b for fatty acids according to JOBACK and GANI are presented in Figure 3. The experimental data used are found in [21] and [46]. It’s noteworthy that it wasn’t possible to apply the fragment-based methodology proposed by [19] for these compounds.

![Figure 3: Relative deviations (%) for normal boiling temperature for fatty acids in the JOBACK and GANI methods. Source: Authors, (2020).](image)

By the analysis of Figure 3, it is observed that the greatest deviation of the GANI method occurred in unsaturated fatty acid C16:1. The absolute average relative deviation (AARD) found for saturated fatty acids was 15.4% for the JOBACK method and 2.9% for GANI. On the other hand, unsaturated FAs gave AARD equal to 17.6% and 2.5% for the same methods, respectively. This analysis shows the agreement of the estimation of T_b by the GANI method for both saturated and unsaturated AGs.

A T_b curve as a function of the molar mass of saturated and unsaturated FAs was constructed to improve previous analysis (Figure 4). In this figure, the boiling points of saturated fatty acids (continuous line) show an almost linear characteristic. In addition, it can be noted the proximity of the experimental data with the values estimated by the GANI method.

![Figure 4: Normal boiling temperatures (K) estimated as a function of molar mass for fatty acids. X represents unsaturation. Source: Authors, (2020).](image)

It’s important to highlight that, in a simulation study of biodiesel production, an error in the value of T_b will imply in the erroneous prediction for other properties, such as critical properties. This fact leads to erroneous results when properties, such as viscosity and density, are calculated using equations and correlations incorporated into commercial simulators.

Critical properties (P_c, V_c, T_c and Z_c): Critical properties were estimated for TAGs, DAGs and MAGs for AMBROSE [23], JOBACK [15], GANI [16] and LYDERSEN [24] methods.

It’s important to mention that only critical temperature data from some AGs were found [46]. No values were available for TAGs, DAGs and MAGs. However, they were considered enough to point out the most precise method.

The AARD obtained for the predicted critical temperatures for the AGs compared to the experimental data are 3.31%, 0.52%, 3.57% and 2.40%, respectively, for the JOBACK, GANI, LYDERSEN and AMBROSE methods.

It’s noted that the GANI method presents an average of the smaller relative deviations for fatty acids (AARD=0.52%). In addition, this method is the only one among those evaluated that requires, as input, only the molecular structure (ASPEN TECH, 2014). Thus, this methodology was chosen to predict the critical properties of the components of vegetable oils.

Critical properties were estimated for several compounds involved in biodiesel production by [47]. Fatty acids, triacylglycerols and methyl esters were analyzed. The authors compared the data obtained with experimental data and pointed to GANI as the most accurate method, compared to the other two studied, [15] and [48].

The critical properties were estimated using the methods of [23] and [15] by [49]. According to the authors, in these two methods the critical temperature estimation is dependent on the normal boiling temperature. Therefore, the critical temperature was also evaluated by the authors by [50], in which the critical temperature is independent of the normal boiling temperature. According to [49], the Joback method presented the lowest...
deviations for critical volume and Ambrose’s method presented the smallest deviations from the data available in [46] for critical temperature and pressure.

[51] evaluated the critical temperature by the methods of [50], [15] and [24]. Comparing the results with predicted values in [46], the authors selected the Lydersen and Joback groups and highlighted the success of these methods because they considered the experimental value of normal boiling temperature in the prediction of critical temperature.

Acentric factor (\(\omega\)): The data estimated by [25], [26] and [29] showed similar results, as can be seen in Table 2.

| Components | Pitzer | Kesler-Lee |
|------------|--------|------------|
| LLL        | -0.6524 | -0.6343    |
| LL         | -0.3183 | -0.3183    |
| L          | 1.0146  | 1.0692     |

Source: Authors, (2020).

In this work, the Pitzer rule was adopted for the determination of the acentric factors of all compounds, as it’s the most used method according to [37].

Enthalpy of formation and Gibbs free energy formation in the standard state \((H^0_e G^0_e)\): Initially, the formation enthalpy data were evaluated for TAG (triolein-OOO), DAG (diolen-OO) and MAG (monolein-O) compared to data available in the literature [52], [53].

Table 3 shows the absolute deviations obtained for each method used in estimating this property. It is noted that the evaluated methods presented relative mean deviations below 3%. The lowest AARD was observed when GANI (0.75%) was used, followed by JOBACK (0.95%). However, the GANI method requires normal boiling temperature as input. Thus, to avoid the error associated with a previously estimated measurement, the method chosen in this work was JOBACK. This requires as input only the molecular structure of the compounds. The Gibb’s free energies also were calculated by JOBACK for TAGs, DAGs and MAGs.

Table 3: AD and AARD of the standard energy of formation of TAG, DAG and MAG.

| DIPPR 801(kJ/mol) | Absolute deviation (kJ/mol) | JOBACK | GANI | BENSON |
|-------------------|-----------------------------|--------|------|--------|
| \(OOO\)           | -2161                       | 14.415 | 1.078| 35.053 |
| \(OO\)            | -1670                       | 14.844 | 16.090 | 44.916 |
| \(O\)             | -1175                       | 15.345 | 14.681 | 45.660 |
| AARD(%)           | 0.954                       | 0.754  | 2.733 |

Source: Authors, (2020).

For the FAs, only data from stearic acid formation enthalpy (-884.7 kJ/mol) were obtained in the literature. The absolute deviations (AD) found for the methods of JOBACK, GANI and BENSON, respectively, were 119.58, 121.14 and 389.22 kJ/mol. Therefore, the first method was chosen to predict the formation energy for fatty acids.

For Gibbs free energy, the method chosen for fatty acids was BENSON [37]. According to [54], this is more accurate in estimating this property compared to GANI and JOBACK methods, considering the effects of neighboring atoms on calculations.

Vapor pressure (\(P_{vap}\)): The vapor pressure was initially evaluated for simple TAGs by the group contribution method [29] and fragment-based method [19], [55]. Figure 5 compares the estimated \(P_{vap}\) data compared to experimental data for tristearin (SSS) and tripalmitin (PPP), obtained in [56].

The methods of [29], [19], [55] showed similar predictions for TAGs as shown in Figure 5. The means of the deviations of the estimated steam pressures, in the temperature range of 230 to 313°C, for the simple triacylglycerols tripalmitin (PPP) and tristearin (SSS) were respectively 15.01 and 20.04 for ZONG and 6.63 and 19.58 for CERIANI. This resulted in AARDs for the ZONG method of 17.52% and for CERIANI 13.11%. By the analysis of the AARDs, the method of [29] showed higher accuracy, so it was pointed to the estimation of steam pressure for TAGs.

Moreover, according to [29], the proposed method can represent any number of fatty and hydrocarbon compounds with the lowest number of parameters and can extrapolate the temperature (critical point), by correctly describing the vapor pressures and vaporization heats.

Enthalpy of vaporization (\(H_{vap}\)): Table 4 shows the estimated values for enthalpy of vaporization compared to data obtained from [46].

According to the analysis in Table 4, the lowest AARD was observed for the Basarova method (AARD=2.68%), followed by Ceriani (AARD=10.05%) and Pitzer (AARD=25.55%). It is noteworthy that, for the methods of [25] and [32], it is necessary to inform the critical temperatures and pressures, and the acentric factors of the TAGs.

Table 4: Relative deviations (RD) and absolute average relative deviation (AARD) of enthalpy of vaporization.

| Temp. \(\text{°C}\) | Exp. \(\text{kJ/mol}\) | RD (%) |
|------------------|-------------------|--------|
| SSS              | 247-314           | 164.63 | 13.84 | 27.74 | 2.89 |
| PPP              | 232-300           | 158.53 | 6.27  | 23.36 | 2.47 |
| AARD(%)          | 10.05             | 25.55  | 2.68  |

Source: Authors, (2020).

For [31] indicated Basarova and Svboda method to predict enthalpy of vaporization values when the Ceriani and Meirelles method (2004) was used [31]. In turn, [29] corrected the temperature limitations of the previously proposed method [18]. So, the method adopted in this study to predict \(\Delta h_{vap}\) of TAGs, DAGs and MAGs was Ceriani [29].

Density (\(\rho\)): Table 5 compares density predictions by [19], [33], [34], [55] to experimental data of simple TAGs [57-59].
All four methods evaluated ensure accuracy by predicting density for TAGs in the temperature range from 80°C to 100°C. The means of the deviations found were low, not exceeding 4.0%.

The method described in [33] doesn’t have adequate correction factors for the DAGs and MAGs of the Rackett equation to estimate the density of vegetable oils. In addition, the method of contribution of groups showed in [34] isn’t particularly designed for predictions of lipid compound density. In other words, there isn’t specific group to describe the portion of glycerol in TAGs, DAGs, MAGs and oils. Therefore, groups CH2, CH, CH2OH, and CHOHe should be used as substitutes when describing the portion of glycerol. To improve the accuracy of the method, data should be developed for the glycerol group.

While the methods described by [19], [55] were based on fragments, generally more accurate than group contribution methods. Also, can be to observe an increasing in order of AARDtotal of densities estimated like: ZONG (1.6%) < IHMELS (1.7%) < HALVROSEN (2.0%). Then, the ZONG method was chosen to predict the densities of TAGs, DAGs and MAGs

Figure 6 illustrates the good agreement of the [34] method for the calculation of the density of fatty acids. For this comparison, experimental data from [60] were used and [61]. In this, continuous lines are estimated data and points are experimental values, equal colors mean the same fatty acid.

The AARD (%) obtained for fatty acids for the [34] method was low (0.30%). Showing that the method is accurate when estimating the density for these compounds.

Heat capacity (cp): The deviations of estimated heat capacities compared to experimental data from [58] and [36] can be seen in Table 6. The data were estimated in the temperature range of 60-180°C.

The results obtained, as shown in Table 6, showed that the method [19] have greater accuracy than [30] and [36]. In this specific case, the AARD for the ZONG method was 0.88%, while for [30] and [36] were 3.26% and 1.51%, respectively. It is worth mentioning that the method of [36] is applicable only to TAGs.

Therefore, the methodology proposed by [19] was adopted for the calculations of this property for TAGs, DAGs and MAGs.

III.2 CHARACTERIZATION OF OILS

Approach 1:

(A) High Performance Gas Chromatography Analysis (GC)

The results of chromatographic analysis can be seen in Table 7.

As in this technique a non-polarized capillary column was used, the analysis is qualitative and simplified, because the TAGs are separated into groups having the same number of carbon atoms, not being possible the determination of unsaturation.

Table 7: Composition of macauba pulp oil obtained by GC.

| Sample | Macauba pulp oil |
|--------|------------------|
| TAGs   | Rep. 1 | Rep. 2 | Examples of possible TAGs |
| C48    | 0.77924 | 0.80121 | PPP |
| C50    | 9.96296 | 9.79972 | PPS, PPO, PPL, PPLn |
| C52    | 42.1804 | 41.928  | SSP |
| C54    | 47.077  | 47.471  | SSS, OOO, LLL |

Note: There isn’t differentiation between C18:0, C18:1 e C18:2.

A more refined separation of TAGs occurs when a polarized column is used in chromatographic analysis. Thus, in a polarized column, the separation of TAGs, such as POO and PLS, which are TAGs with the same number of carbons and degree of unsaturation, but with different polarity, can be achieved. In both non-polar and polarized columns, there is no discrimination between isomeric triacylglycerols such as POO and OPO [42].

(B) Computational Analysis

Table 8 shows the proportions of triacylglycerols through the composition of fatty acids obtained from the computational program created in this work according to the methodology cited by [42].

Table 8: Characterization of macauba pulp oil in TAGs.

| Groups | TAGs | % m/m | Groups | TAGs | % m/m |
|--------|------|-------|--------|------|-------|
| C50:1  | POP  | 8.0   | C52:4  | PLnO | 4.0   |
| C50:1  | PPS  | 2.0   | C54:2  | SOO  | 4.0   |
| C52:2  | PPO  | 28.0  | C54:3  | OOO  | 26.0  |
| C52:3  | OPnO | 4.0   | C54:4  | OOL  | 14.0  |
| C52:3  | PLO  | 2.0   | C54:5  | OLL  | 2.0   |
| C52:4  | LOPn | 6.0   |        |      |       |

Source: Authors, (2020).
The identification of triacylglycerols by groups with different numbers of carbon atoms and by different combinations of saturated and unsaturated fatty acids were made by comparing the percentage data obtained from the GC with those provided by the program created on the computer. The data in Table 9 show that the proportions of each component calculated by the computer (PC) are close to the average obtained by chromatographic analysis (GC).

Table 9: Composition of macauba pulp oil in triacylglycerols by GC and using a program elaborated on the computer (PC).

| Number of Carbons | Composition (%) |
|-------------------|-----------------|
|                   | PC              | GC              |
| C48               | 0.00            | 0.79            |
| C50               | 10.00           | 9.88            |
| C52               | 44.00           | 42.05           |
| C54               | 46.00           | 47.27           |

Source: Authors, (2020).

Thus, the list of triacylglycerols presented in Table 8, obtained by the computer program, will be considered as the composition of the macauba pulp oil for this present study. Since the mixed triacylglycerol approach is closer to reality, and the results of chromatographic analysis show that the program used for the distribution of mixed TAGs meets the data obtained experimentally.

III.3 SELECTION OF THE MACAUBA OIL CHARACTERIZATION METHOD

The choice of the appropriate characterization method is important in the prediction of thermophysical properties. Some criteria to be considered in the adoption of the method are ease of use, rigorous characterization, and satisfactory predictions. Table 8 and [43] showed the composition of macauba oil in terms of TAGs and fatty acids, respectively.

Predictions of some properties were made to compare the two approaches to oil characterization. Table 10 shows predictions of density and calorific capacity of macauba pulp oil for the approach in mixed TAGs and fatty acids.

Table 10: Densities and heat capacities by the two approaches of characterization of macauba pulp oil.

| Method | TAG-Mixed | Fatty acids |
|--------|-----------|-------------|
| Density | [33] | 0.17 | 0.16 |
|         | [19]  | 0.97 | ND   |
|         | [34]  | 1.79 | 1.83 |
| Heat capacity | [36] | 2.17 | 2.50 |
|         | [19]  | 2.20 | ND   |
|         | [30]  | 0.56 | 0.47 |

Source: Authors, (2020).

According to the analysis of Table 10, it can be observed that the two approaches had similar estimates, with small average of relative deviations. However, it’s recommended to use the characterization of oils in mixed TAGs, because it is closer to the real.

III.4 PROPERTIES OF VEGETABLE OILS

It’s known that vegetable oil is a mixture of various TAGs, DAGs, MAGs and free FAs. In this work was adopted that vegetable oils were composed only of TAGs and DAGs. To determine the properties of vegetable oils, their compositions presented above, and the predicted properties of the compounds present in the mixture were used, using the ideal mixing rule (equation 47).

\[ P_{\text{mixture}} = \sum_{i=1}^{n} x_i P_{\text{pure},i} \] (47)

The methods adopted to predict the thermophysical properties of TAGs, DAGs and MAGs, consequently of vegetable oils, are summarized in Table 11.

Table 11: Methods adopted to predict the thermophysical properties of vegetable oils.

| Properties | Methods |
|------------|---------|
| T_b        | Zong et al. (2010a and 2010b) |
| T_c, P_c e V_c | Constantinou and Gani (1994) |
| Acentric factor(\(\omega\)) | Pitzer (1995) |
| Enthalpy formation(H_0) | Joback and Reid (1987) |
| Gibbs Energy(G_0) | Joback and Reid (1987) |
| P_vap | Ceriani et al (2013) |
| H_vap | Ceriani et al (2013) |
| \(\rho\) | Zong et al. (2010a e 2010b) |
| \(c_p\) | Zong et al. (2010a e 2010b) |

Source: Authors, (2020).

Therefore, the estimated properties for vegetable oils were compared with experimental data available in the literature. The representation of temperature-dependent thermophysical properties, such as vaporization enthalpy (H_vap), heat capacity (\(c_p\)), density (\(\rho\)) and vapor pressure (P_vap) is shown in Figure7(a), Figure 7(b), Figure 7(c) and Figure 7(d), respectively.

By the analysis of Figure 7 (a), it is observed that the enthalpy of vaporization of vegetable oils decreases with increasing temperature. Heat of vaporization for soybean oil was determined by [62], H_vap=184.9 kJ/mol. The predicted value for soybean oil in this study was in the range of 216 to 174 kJ/mol, showing that there is agreement of the predicted given with the experimental. By the analysis of Figure 7 (b), it can be observed that the calorific capacities of the analyzed vegetable oils increased linearly with the increase in temperature from 20 to 80°C. Similar behavior was observed by [63].

Density and viscosity for babassu, buriti, Brazil nut, macadamia and grape seed oils were obtained by [38]. The densities found were from 912 to 876 kg/m³, in the temperature range of 20-70°C. [64] determined the density of palm oil, \(\rho\)=888 kg/m³ at 50°C, close to the value found in this study (880 kg/m³). For sunflower oil, these authors found density in the range of 880 to 945 kg/m³ in the temperature range of -20 to 80°C. Similar values were found, as shown in Figure 7 (c) [65] determined for the pulp oil of macauba densities in the range of 918.5 to 925.3 kg/m³,showing again that the estimates calculated in this study reached good precision.

Vapor pressure of various vegetable oils was determined by [66]. The authors found vapor pressures for soybean oil in the range of 2.6kPa to 73.5kPa, ranging from 250-330°C. Similar values were observed, as can be seen in Figure 7 (d).

Table 12 shows the rest of the thermophysical properties that were estimated for the different vegetable oils under study, which can be able to use as input data for engineering calculations like simulations and optimize process.
Figure 7: Estimated properties dependent on temperature (a) Vaporization enthalpy, (b) Heat capacity, (c) Density and (d) vapor pressure for sunflower (▲), palm (■), soybean (♦) and macauba pulp (●) oil.
Source: Authors, (2020).

Table 12: Estimated properties for vegetable oils by the methods adopted in this work.

| Properties  | Sunflower | Soy | Palm | Macauba |
|-------------|-----------|-----|------|---------|
| T_b (K)     | 583.34    | 662.40 | 615.54 | 739.08 |
| T_c (K)     | 836.34    | 950.30 | 879.71 | 1051.57 |
| P_c (N/m²)  | 2.95 x 10⁵ | 3.36 x 10⁵ | 3.17 x 10⁵ | 3.86 x 10⁵ |
| V_c (m³/kmol)| 2.80      | 3.18  | 2.91  | 3.48    |
| Z_e         | 0.04      | 0.05  | 0.03  | 0.03    |
| H_f (J/kmol)| -1.40 x 10⁹ | -1.76 x 10⁹ | -1.84 x 10⁹ | -2.09 x 10⁹ |
| G_f (J/kmol)| -6.95 x 10⁷ | -2.16 x 10⁸ | -3.99 x 10⁸ | -3.73 x 10⁸ |
| ω           | -0.59     | -0.66 | -0.62 | -0.75   |

Source: Authors, (2020).

Finally, the values predicted in this study are in acceptable ranges when compared with experimental data of similar vegetable oils. The difference in the composition of oils, especially in the number of unsaturation, causes changes in thermodynamic properties, as seen in the previous analyses.

IV. CONCLUSIONS

This work showed that the modeling of properties of TAGs, DAGs, MAGs and fatty acids is closely linked to the estimation of base parameters, such as normal boiling temperature, critical temperature, pressure and volume and acentric factor. In addition, it was shown that the use of experimental data to validate the method should be carried out with caution, considering that the nature of many available data is predicted.

Prediction models of properties such as group contribution and fragment approach were evaluated to determine the normal boiling point, critical properties, vapor pressure, liquid density, calorific capacity and vaporization enthalpy of fatty acids, triglycerides, diacylglycerols and monoacylglycerols. It was possible to arrive at values similar to the experimental values of the properties to vegetable oils with the application of selected methods for each property.

The estimation of the thermophysical properties of vegetable oils is of paramount importance when one wishes to evaluate biofuel production processes by simulation technique. This is because experimental data of these raw materials are scarce in the literature. Therefore, a database with values close to the actual is required for the use of simulators to have correct predictions. Thus, in this work a database was created that can be used as input parameters in simulation projects.

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