

Solubility and Nucleation of Methyl Stearate as a Function of Crystallization Environment

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Supporting Information

ABSTRACT: Crystallization studies of methyl stearate from supersaturated dodecane, kerosene, and toluene solutions reveal strong evidence that solvent choice influences solubility and nucleation behavior. Solute solubility is less than ideal with toluene, kerosene, and dodecane, respectively, exhibiting the closest behavior to ideality, the latter consistent with the highest solvation. Polythermal crystallization studies using the Kashchiev–Borissova–Hammond–Roberts (KBHR) model [Kashchiev et al. *J. Phys. Chem. B* 2010, 114, 5441; Kashchiev et al. *J. Cryst. Growth* 2010, 312, 698; Camacho et al. *CrystEngComm* 2014, 16, 974] reveal a progressive nucleation (PN) mechanism with crystallite interfacial tension (γ_{eff}) values between 0.94 and 1.55 mJ/m², between 1.21 and 1.91 mJ/m², and between 1.18 and 1.88 mJ/m² for dodecane, kerosene, and toluene, respectively. Nucleation rates at the critical undercooling lie between 4.56×10^{16} and 1.79×10^{17} nuclei/mL·s, with the highest rates associated with crystallization from kerosene solutions. Iso-supersaturation nucleation rates are the highest for dodecane ranging from 2.39×10^{17} to 3.63×10^{18} nuclei/mL·s. Nucleation in toluene appears to be hindered by its relatively higher interfacial tension, which is associated with nucleation rates about an order of magnitude less than those obtained for dodecane.

1. INTRODUCTION

The study of diesel and biodiesel fuel crystallization is of importance to the fuel industry as poor cold-flow properties of these mixtures can cause operability problems within vehicle engines, pipelines, and vehicle tanks under cold weather conditions due to the formation of crystals at low temperatures. To date most research has focused on the crystallization within diesel fuel,^{4–15} which mostly comprises alkanes, with much less emphasis being placed on the study of biodiesel fuel. First-generation biodiesel fuels are generally a mixture of both saturated and unsaturated methyl esters from which the former represent an important proportion of these solutions and commonly contain methyl palmitate (C16:0) and methyl stearate (C18:0). The cold-flow behavior of biodiesel is determined to a great extent by the amount of saturated compounds present in its composition.

Some studies^{16–19} have been carried out in order to characterize the cold-flow behavior of biodiesel fuel produced from different bioresources, measuring certain properties such as cloud point (CP), pour point (PP), and cold filter plugging point (CFPP) and the effect of cold-flow improvers on these properties. The fractionation of methyl esters crystallizing from biodiesel fuel mixtures produced from different bioresources has also been addressed.^{20–25} However, up to now there has been a lack of fundamental studies on the nucleation of saturated methyl esters, such as methyl palmitate and methyl stearate.

As a baseline case, it is the aim of this study to deliver fundamental information on the solubility and nucleation of methyl stearate measured as a function of solution environment. In this the solubility is assessed using the van't Hoff analysis.²⁶ This was combined with a rigorous analysis of the associated nucleation data through the recently developed Kashchiev–Borissova–Hammond–Roberts (KBHR) theory.^{1–3} This theory, outlined in section 2, permits a first-principles analysis of nucleation

kinetics and enables the derivation of key kinetic parameters and the characterization of the nucleation mechanism.

Because in real-world operation biodiesel fuel is commonly mixed with traditional diesel fuel, three different model solvents were chosen representing the variation in solvation environment, viz., dodecane, toluene, and kerosene.

2. NUCLEATION KINETICS AND MECHANISM

2.1. Classical Nucleation Theory (CNT) and Its Assessment. The classical 3D nucleation theory (CNT) provides a model for the prediction of the rate at which nuclei of a new crystalline phase are formed. This model given by expression 1 describes the dependence of the nucleation rate (J) on two terms: a thermodynamic (exponential) component which accounts mainly for the effect of temperature, supersaturation ratio (S), and effective interfacial tension (γ_{eff}) on the formation of nuclei and a kinetic (pre-exponential) component which describes the frequency with which molecules will attach to the nucleus. The relative balance between these two terms determines a system's nucleation behavior.

$$J = K_J \exp\{-[k_n v_0^2 \gamma_{\text{eff}}^3 / (kT)^3 (\ln S)^2]\} \quad (1)$$

where K_J is the nucleation rate constant, k_n is the nuclei numerical shape factor, i.e., $16\pi/3$ for spherical nuclei and 32 for cubic nuclei, v_0 is the volume occupied by a solute molecule in the crystal, and k is the Boltzmann constant. S , the supersaturation, is given by:

$$S = \frac{x}{x_e} \quad (2)$$

Received: October 19, 2017

Revised: February 9, 2018

Published: February 20, 2018

Here x is the solution's molar concentration and x_c is the equilibrium concentration.

Supersaturation can also be expressed as the relative supersaturation (σ) as given by [expression 3](#):

$$\sigma = S - 1 \quad (3)$$

The analysis of nucleation kinetics can be performed by either the isothermal methodology or the polythermal methodology, which both use the concept of solution state metastability to create the supersaturation needed to promote nucleation. (see [Figure 1](#)). The isothermal method makes use of the kinetic expressions derived from classical nucleation theory, in particular, utilizing the assumption that the induction time (τ) can be taken as being inversely related to the nucleation rate (J). In contrast, the polythermal method assesses nucleation through establishing the effect of cooling rate (q) on crystallization temperature (T_c).

2.1.1. The Isothermal Method. In this method, one can calculate key nucleation parameters including the interfacial tension (γ) and the critical nucleation cluster size (r^*), as a function of solution supersaturation and temperature.

In this case, many experiments ideally should be carried out, i.e., typically 5–10 different supersaturations with ca. 80 repeats at each chosen supersaturation.²⁷ The analysis of this data is relatively simple, though, as nucleation rates can be directly extracted by fitting a probability distribution of the measured induction times. A comprehensive assessment that quantifies the uncertainty associated with the parameters estimated using this methodology is provided in the work presented by Xiao et al.²⁸

2.1.2. The Polythermal Method. The polythermal approach continuously varies the solution supersaturation, and hence induction time, upon cooling and assesses the balance between excess concentration generation via the cooling rate and the material's intrinsic nucleation rate. The effect that supersaturations has on nucleation is implicitly evaluated over the whole width of the metastable zone. This is defined, for the work presented here, as the difference between the equilibrium concentration derived from van't Hoff analysis and the solution concentration pertinent to the temperature at which spontaneous crystallization occurs.

The process involves driving the solution by cooling until the induction time is effectively zero, i.e., the point at which spontaneous nucleation takes place. Under these conditions mass transfer due to molecular diffusion can be expected to be rapid and hence the degree of statistical variation would be much less than that for induction time measurements. The latter is typically recorded for lower supersaturations, concomitantly larger cluster sizes, and lower levels of molecular diffusion.

In the polythermal case, experimental data is comparatively easy to collect using automated temperature controlled solution turbidimetric methods. As nucleation is clearly of a stochastic nature, in our analysis all the determined parameters are presented with their corresponding standard deviations using the most reliable experimental methodology developed in our previous work.³ In this, we have shown that typically five repeats at each cooling rate are sufficient (section 2 of the Supporting Information provided in ref 3) to obtain reliable data to perform an assessment using this approach.

The analysis of polythermal data can be much more complex than the isothermal case and could be performed using a number of different available models, both empirical^{29,30} and first principles.^{31–33} The most widely used interpretation of $\Delta T_c(q)$ data is using the empirical Nyvlt expressions.^{29,30} However, given the empirical nature of the Nyvlt approach, Kubota³¹ and Sangwal^{32,33}

have reinterpreted these equations in order to derive more physically meaningful parameters. An example of the application of these approaches is provided by Mitchell et al.,³⁴ where key kinetic parameters were derived for paracetamol–ethanol solutions.

2.1.3. Comparison between Isothermal and Polythermal Methods. [Figure 1](#) compares the different approaches used in

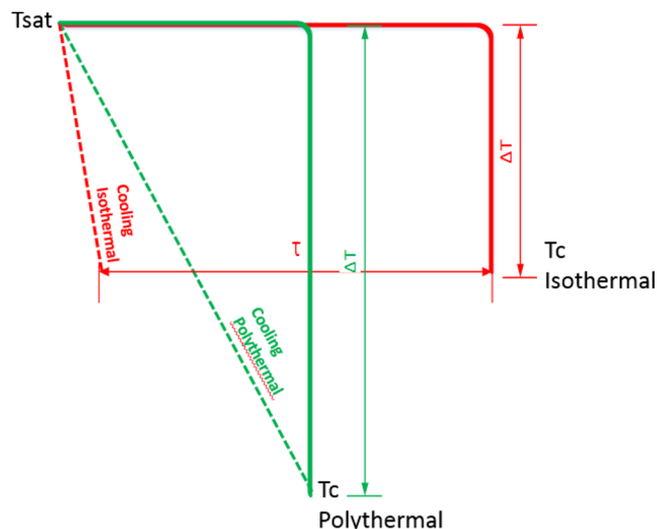


Figure 1. Comparative scheme of the different approaches used to collect experimental crystallization temperatures using both the isothermal and the polythermal methods.

each of the methodologies to collect experimental data. Using a turbidimetric technique, in the isothermal method a clear homogeneous solution is rapidly cooled to a given supersaturation where the solution is kept until crystallization is detected through the increase in the solution's turbidity, after a given induction time (τ). On the other hand, in the polythermal method a solution is cooled continuously at a given rate, until crystallization is detected through the increase in the solution's turbidity. Although in the isothermal method the onset of crystallization is monitored at the same temperature (or supersaturation), this method is to a certain extent inherently polythermal as the solution has already gone through a number of different temperatures (or saturations) during the initial rapid cooling to the chosen supersaturation. Although the two methodologies are apparently quite different, interestingly, they reach the same end point; e.g., as shown in Table 9 in Camacho et al.,³ nucleation parameters calculated via the two methodologies are broadly equivalent. For clarity to the reader on how to establish the corresponding supersaturations using either of the two methodologies, [Figure S1](#) in the [Supporting Information](#) (SI) provides also a comparison based on a plot of concentration vs temperature.

Due to the nature of crystallization in methyl ester solutions, characterized by very narrow metastable zones and very short induction times, a polythermal method was used to collect experimental data. Details of this methodology are provided in [section 4](#).

2.2. The Kashchiev–Borissova–Hammond–Roberts (KBHR) Approach. A first-principles analytical polythermal approach, the Kashchiev–Borissova–Hammond–Roberts (KBHR) approach, comprises a set of model equations analytically derived starting from the Kolmogorov–Johnson–Mehl–Avrami (KJMA) expression. Such a model is analogous to “chemical reaction progress kinetic analysis,”³⁵ in which reaction progress is monitored as a function of a dynamically changing

reactant concentration, which is akin to carrying out hundreds of separate initial rate experiments.³⁵ A key outcome of the KBHR model is the so-called “rule of three”^{1,2} which can be used to discriminate between two case nucleation mechanisms, i.e., progressive nucleation and instantaneous nucleation.

In progressive nucleation (PN),² new crystal nuclei are continuously formed in the presence of the already growing ones. In this case the measured induction times are associated with both the nucleation and the growth processes until the crystals reach a size where they are detectable optically via the turbidimetric technique.

In instantaneous nucleation (IN),¹ all nuclei emerge at once at the beginning of the crystallization process to subsequently grow and develop into crystals. In this case, strong nucleation sites will favor the nucleation process and therefore inductions times are only associated with the time for these crystals to grow to a detectable size.

The expressions related to both the PN and IN mechanisms, analytically derived in the KBHR approach, are presented below.²

2.2.1. Progressive Nucleation. The general expression relating critical undercooling and cooling rate for this mechanism is given by expression 4:

$$\ln q = \ln q_0 + a_1 \ln u_c - \frac{a_2}{(1 - u_c)u_c^2} \quad (4)$$

The relative critical undercooling (u_c) is a dimensionless quantity associated with the critical undercooling (ΔT_c) given by

$$u_c = \frac{\Delta T_c}{T_e} \quad (5)$$

Similarly, the critical undercooling ΔT_c , which represents the solution's metastability limit in terms of temperature, is defined as

$$\Delta T_c = T_e - T_c \quad (6)$$

where T_e and T_c are the solution equilibrium and crystallization temperatures, respectively.

Expression 4 describes the dependence of the number of crystals at the detection point (N_{det}) on the cooling rate (q) when $d = 0$; i.e., the volume of single crystals is unaccounted for (see derivation in the SI) and thus the free parameters a_1 , a_2 , and q_0 are given by²

$$a_1 = 3 \quad (7)$$

$$a_2 = b \quad (8)$$

$$q_0 = \frac{VK_j T_e}{N_{\text{det}} 2b} \quad (9)$$

where V is the volume of the solution and b is given by²

$$b = \frac{k_n v_0^2 \gamma_{\text{eff}}^3}{k T_e \lambda^2} \quad (10)$$

In this expression λ is the molecular latent heat of crystallization.

When eq 4 is derived by means of the relative volume of crystals (α), the parameters q_0 , a_1 , and a_2 are defined by

$$a_1 = 3 + \frac{3nmd}{md + 1} \quad (11)$$

$$a_2 = \frac{b}{md + 1} \quad (12)$$

$$q_0 = T_e \left\{ \frac{\Gamma[(n + 1)md + 1] K_v a^{nmd} K_j K_G^{md}}{(n + 1)^d (2b)^{(n+1)md+1} \alpha_{\text{det}}} \right\}^{1/(md+1)} \quad (13)$$

Here, d is the dimensionality of crystallite growth, i.e., 3 for spheres or cubes, 2 for disks or plates, and 1 for needles. n and $m > 0$ are the crystallite growth exponents which are related to the different growth mechanisms.³⁶ The $n = 1$ case corresponds to growth mediated by diffusion of solute toward the crystallite or transfer of solute across the crystal/solution interface. The $n = 2$ case characterizes growth controlled by the presence of screw dislocations in the crystallite. The parameter m ranges between 1/2 and 1: $m = 1/2$ is for growth controlled by undisturbed diffusion of solute, and $m = 1$ is for growth by diffusion of solute through a stagnant layer around the crystallite or for normal or spiral growth limited by transfer of solute across the crystal/solution interface. At $m = 1$ the crystallite radius increases linearly with time.^{1,2,37} $k_v(m^{3-d})$ is the crystallite's growth shape factor, i.e., $4\pi/3$ for spheres, 8 for cubes, πH_0 for disks, $4H_0$ for square plates (H_0 is the fixed disk or plate thickness), and $2A_0$ for needles (A_0 is the fixed needle cross-sectional area). K_G is the crystal growth rate constant, Γ is the gamma function, and α_{det} is the relative volume of crystals at the detection point.

2.2.2. Instantaneous Nucleation. In the case of IN, the expression for the dependence of relative critical undercooling on cooling rate is given by

$$\ln q = \ln q_0 + \left(\frac{1}{m} \right) \ln [u_c^{(n+1)m} - u_0^{(n+1)m}] \quad (14)$$

In this expression $u_0 \geq 0$, $u_c > u_0$, and the parameter q_0 is given by

$$q_0 = \left[\frac{k_v C_0}{(n + 1)^d \alpha_{\text{det}}} \right]^{1/md} a^n K_G T_e \quad (15)$$

where C_0 is the concentration of nuclei at the time at which the instantaneous nucleated crystallites are formed and a is given by expression 16:

$$a = \frac{\lambda}{k T_e} \quad (16)$$

If, additionally, the undercooling at which all nuclei spontaneously appear is small enough so that

$$u_0^{(n+1)m} \ll u_c^{(n+1)m} \quad (17)$$

eq 14 takes the form of a straight line given by

$$\ln q = \ln q_0 + (n + 1) \ln u_c \quad (18)$$

The $\ln q$ vs u_c line corresponding to expression 4 is only slightly curved; thus in a not too wide experimental q range it can be approximated to a straight line.^{1,2} This linear relationship can be analytically derived using an arbitrarily critical undercooling as shown in ref 2 and is given by expression 19. When comparing this expression with eq 18, then from the slope of a line of the dependence of relative critical undercooling for crystallization (u_c) on the cooling rate (q), the nucleation mechanism can be established using the “rule of three”:^{1,2} slope $> 3 = \text{PN}$ or slope $< 3 = \text{IN}$.

$$\ln q = \ln Q + \left(3 + \frac{3nmd}{md + 1} + \omega a_2 \right) \ln u_c \quad (19)$$

In expression 19 ω is a positive number and Q is a parameter related to q_0 .¹

Both expressions 4 and 14 are subjected to the inequalities (20) as shown in the analytical derivation in section 2 of the SI. This means that this analysis is restricted to small enough values of the critical undercooling (u) for which the inequalities are satisfied:

$$u < 0.1, \quad au < 1 \quad (20)$$

The critical radius of the nucleus (r^*) and the number (i^*) of molecules in the critical nucleus can be calculated from expressions 21 and 22 given in terms of the relative undercooling u .

$$r^* = \frac{2\gamma_{\text{eff}}v_0}{\lambda u} \quad (21)$$

$$i^* = \frac{2bkT_c}{\lambda u^3} \quad (22)$$

The classical 3D nucleation rate model, given in terms of the parameters defined by the KBHR approach, is presented in eq 23:

$$J(t) = K_j e^{-b/[(1-u)u^2]} \quad (23)$$

Here K_j is related to the attachment frequency of monomers to the nucleus (f^*), the concentration of nucleation sites (C_0), and the Zeldovich factor Z . The latter accounts for the probability that a critical nucleus would become a crystal and not redissolve.

The attachment frequency (f^*) is given by either expression 24 or 25 for attachment of monomers controlled by volume diffusion or interface transfer, respectively.³⁷

$$f^* = 4\pi r^* \xi D X_1 \quad (24)$$

$$f^* = \xi \omega^* d_0 A^* X_1 \quad (25)$$

where ξ is the sticking coefficient, D is the diffusion coefficient of colliding building units, X_1 is the concentration of colliding building units, ω^* is the transfer frequency of building units from adsorbed to integrated, d_0 is the thickness of the adsorbed surface layer, and A^* is the surface area of the nucleus.

The attachment of building units to the cluster is quite sensitive to changes in the temperature, the effect being mainly due to the viscosity. This is particularly relevant when the attachment of monomers is controlled by volume diffusion and nucleation occurs within a temperature range in which the solution viscosity varies strongly with T .

It is important to highlight here that, due to the nature of the derivation of the KBHR approach, the assessment of polythermal data using this theory is subject to the following assumptions:

1. The supersaturation at which crystallization is detected (MSZW) has to be sufficiently small, and the solution viscosity does not change significantly within the assessed range.
2. The former will allow the assumption that the nucleation mechanism does not change within the range of concentrations measured on cooling.

A fuller description of this theory is provided in the SI.

3. MATERIALS AND METHODS

3.1. Materials. Methyl stearate, dodecane, and toluene were purchased from Sigma-Aldrich. The purity of the methyl stearate used was 96%, and that of the two solvents was higher than 99%. No further purification was carried out. Kerosene was supplied by Infineum Ltd. (Milton Hill, Oxfordshire, U.K.). Its hydrocarbon composition is summarized in Table 1. Its n -alkane chain length distribution is given in Figure S2 of the SI.

3.2. Equipment and Experimental Procedure for Polythermal Data Collection. Crystallization experiments were carried out using the Technobis Crystal 16 system <https://www.crystallizationsystems.com/crystal16>. This provides a multiple reactor facility with four separate

Table 1. Composition of Kerosene from Two-Dimensional Gas Chromatography Analysis Performed by Infineum UK

	hydrocarbon	mass %
paraffins	unbranched alkanes	16.29
	isoparaffins	23.04
cycloalkanes	naphthenes	42.40
aromatics	alkylbenzenes	7.60
	benzocycloparaffins	6.80
	naphthalenes	3.43
	biphenyls/acenaphthenes	0.30
	fluorenes	0.15

Peltier heated aluminum blocks, each of which has a capacity to hold four magnetically agitated 1 mL solution vials. Each block can be individually programmed to follow a given temperature profile during which the variations in the solution turbidity are followed as a function of temperature.

Solutions of methyl stearate in three different solvents, dodecane, kerosene, and toluene, were prepared at solution concentrations of 200, 250, 300, and 350 g of solute/L of solvent for the first two solvents and 154, 192, 231, and 269 g of solute/L of solvent for toluene.

In order to ensure accurate measurement of temperatures, calibration of the Crystal 16 unit was required. Four vials containing each of the solvents were placed in each of the blocks which were programmed to a specific temperature in the range 20 to -8 °C. While each block was kept at a chosen temperature, measurements of the actual temperature with ± 0.5 °C accuracy were carried out by positioning a thermocouple within each of the vials. The average of the four temperature readings obtained in each block was plotted against the programmed temperature and fitted by a straight line represented by the expressions $y = 0.87x + 2.20$, $y = 0.96x + 1.34$, and $y = 0.95x + 0.99$ for dodecane, kerosene, and toluene, respectively. These expressions were then used to correct the experimentally measured temperature values. The temperature-calibration lines obtained are given in Figure S3 in the SI.

The solutions were subjected to heating and cooling cycles, with each cycle initiated by heating the solutions to 40 °C, where they were held for 1 h to ensure complete homogenization, and then cooled to -15 °C, where they were also held for 1 h to allow equilibration. This temperature profile was applied at each solution's concentration using four different rates of 0.25, 1.0, 3.2, and 9.0 °C/min for the dodecane and kerosene systems and 0.25, 0.5, 1.0, and 1.5 °C/min for toluene solutions. The ranges of both concentrations and cooling rates for solutions of a given solvent were chosen to ensure accurate temperature profiles. This was achieved by setting temperatures profiles in which crystallization was detected above -15 °C (lowest working temperature of the Crystal 16) and a sufficiently wide range of cooling rates in compliance with both the equipment cooling power capacity and the methodology applied. For the solutions where crystallization was detected at lower temperatures, a narrower cooling range had to be used due to the decrease in the equipment cooling capacity observed at these temperature levels.

At each rate the temperature cycle was repeated five times to obtain average values for the crystallization and dissolution temperatures T_c and T_{diss} . These were estimated based upon the points in the turbidity profile at which sudden changes in light transmittance are detected, T_c . Figure 2 shows a typical experimental profile together with a representative raw data set for one of the experimental runs.

To assess the influence of the solute and solvent molecule's polarity on the solubility of methyl stearate, its dipole moment was calculated in a vacuum using three different methods: semiempirical:AM1, Hartree-Fock:3-21G, and DFT:B3LYP/6-31G*, delivering values of 1.67, 1.54, and 1.52 D, respectively (Ken Lewtas, private communication, Sept 4, 2015).

3.3. Data Analysis. 3.3.1. Solubility. The polythermal data were used to establish the solubility for methyl stearate in dodecane, kerosene, and toluene solutions by extrapolation of the $T_{\text{diss}}(q)$ lines to 0 °C/min. The solubility was modeled according to the van't Hoff equation given by expression 26. This expression is derived from the general expression of solid-liquid equilibrium assuming that the specific heat capacity

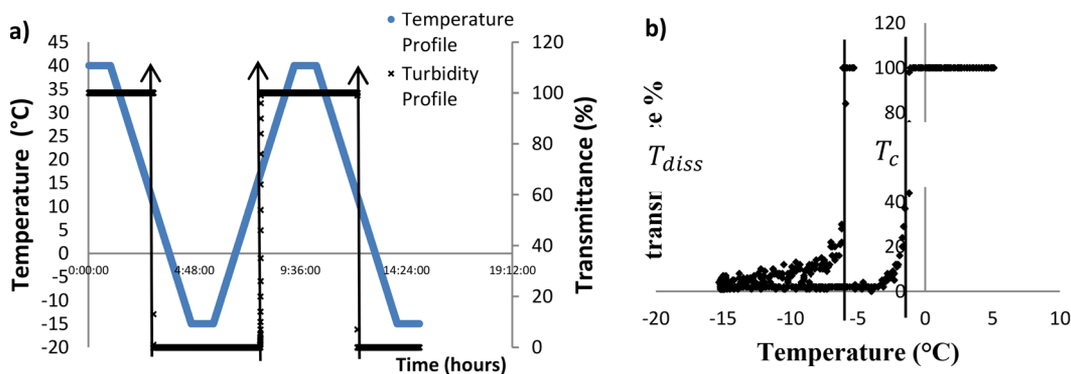


Figure 2. (a) Typical experimental profile using Crystal 16 by applying the polythermal method. (b) Representative turbidity profile in transmittance vs temperature coordinates obtained by the application of a polythermal method.

(ΔC_p) can be neglected. This assumption can be applied to the analysis of methyl stearate solubility as according to previous work³⁸ ΔC_p values do not change significantly within the temperature range studied here.

$$\ln(x_e) = -\frac{\Delta H_{\text{diss}}}{RT} + \frac{\Delta S_{\text{diss}}}{R} \quad (26)$$

The strength of the solutions' chemical interactions was assessed by comparing the solubility with the ideal model solid–liquid equilibrium given by expression 27:

$$\ln(x_e) = -\frac{\Delta H_{\text{fus}}}{RT} + \frac{\Delta S_{\text{fus}}}{R} \quad (27)$$

In these expressions x_e is the mole fraction of the solute in the solution at saturation; T is the solution temperature; ΔH_{diss} (J/mol) and ΔH_{fus} (J/mol) are the molal enthalpies of dissolution and fusion, respectively; ΔS_{diss} (J/mol·K) and ΔS_{fus} (J/mol·K) are the molal entropies of dissolution and fusion, respectively; and R (8.314 J/mol·K) is the gas constant.

A compound's molar solubility can be related to the solubility of its ideal state through its activity coefficient (Υ) which is given by expression 28 obtained by equating the activity a_e at the solution's equilibrium and that of its ideal state:

$$\Upsilon = \frac{x_{\text{ideal}}}{x_e} \quad (28)$$

An activity coefficient equal to 1 indicates the solution behaves ideally; i.e., the enthalpy of dissolution is equal to 0, as the energy needed to break solute–solute interactions added to that of breaking solvent–solvent interactions is equal to the energy released when solute–solvent bonds are formed. On the other hand, if the activity coefficient is either lower or higher than 1, this would indicate a solution will dissolve more or less of the expected solute concentration at equilibrium, respectively. For $\Upsilon > 1$ forces of attraction between like molecules would be favored over those of unlike molecules. For $\Upsilon < 1$ forces of attraction between unlike molecules would be favored over those of unlike molecules. Deviations from a solution's ideal behavior can be due to either enthalpic or entropic factors, and this can be established by comparing the solubility of the van't Hoff model line with that of the ideal solution under the same temperature range. If the slopes of the lines are different, dissolution would be both enthalpic and entropic driven. If the lines are parallel, this would indicate that dissolution is only entropically driven.

3.3.2. Nucleation kinetics. Using the KBHR approach, the analysis of nucleation kinetics, from polythermal experimental data, can be performed following the procedure in the flowchart given in Figure 4 of the SI.

4. RESULTS AND DISCUSSION

4.1. Solubility. The average values for the collected crystallization T_c and dissolution T_{diss} temperatures together with the corresponding standard deviations (SDs) as a function of cooling rate q and concentration are presented in the SI. An example of the linear dependence of T_c and T_{diss} on q is given in Figure 3.

The saturation temperature T_e , obtained from the extrapolation to 0 °C/min of $T_{\text{diss}}(q)$ lines at each of the solution's concentrations, are also shown in Table S1 of the SI.

The specific data points used to model the solubility according to the van't Hoff equation are given in Table 2, together with the corresponding enthalpy and entropy of dissolution and mixing (ΔH_{mix} and ΔS_{mix}). The activity coefficients and their modeled dependence on temperature are also provided. Methyl stearate enthalpy (ΔH_m) and entropy (ΔS_m) of melting are 53.94 kJ/mol and 0.17 kJ/mol·K, respectively. The subtraction of these values from those of enthalpy and entropy of dissolution delivered the corresponding values of enthalpy and entropy of mixing. The comparative van't Hoff lines are given in Figure 4.

van't Hoff plots fit well to a linear model for all solutions studied. This indicates consistency of the structure of a material, in the corresponding range of temperature. Activity coefficients are all higher than 1, indicating that solute–solute interactions are stronger in all cases. However, the significantly lower values obtained in toluene together with the lowest enthalpy of mixing suggest that in these solutions solute–solvent interactions are stronger than in the other two solvent systems.

The highest solubility is observed in toluene and decreases as a function of solvent in the following order: toluene > kerosene > dodecane. This is in the same order as polarity/aromaticity. Fatty acid methyl esters are organic compounds with medium polarity due to the presence in their chemical structure of both a COO–alkyl group and a nonpolar long hydrocarbon chain. The solubility trend could be justified in terms of the solvent polarity and the like-dissolves-like rule of thumb; i.e., a nonpolar compound will be dissolved by nonpolar solvents and vice versa. Straight chain n -alkanes such as dodecane with a dipole moment of around 0.07 D³⁹ are essentially nonpolar molecules which will have lower affinity with methyl stearate, whose dipole moment calculated in a vacuum ranges between 1.52 and 1.67 D. On the other hand, toluene will show better affinity due to a higher dipole moment equal to 0.36 D.³⁹ Kerosene is expected to deliver a solubility higher than that of dodecane but lower than that observed in toluene due to its composition that comprises different types of hydrocarbon molecules including aromatic species.

According to this, it could be expected that dipole–dipole interactions between methyl stearate and toluene molecules are stronger. Additionally, it has been hypothesized^{39,40} that the low activity coefficients obtained in the case of toluene are likely to be, to a great extent, due to the polarizing effect created by the delocalized electron cloud around the benzene ring.⁴⁰ This could cause the distortion of the electron cloud around the solvent molecule inducing temporary dipoles among solute–solvent

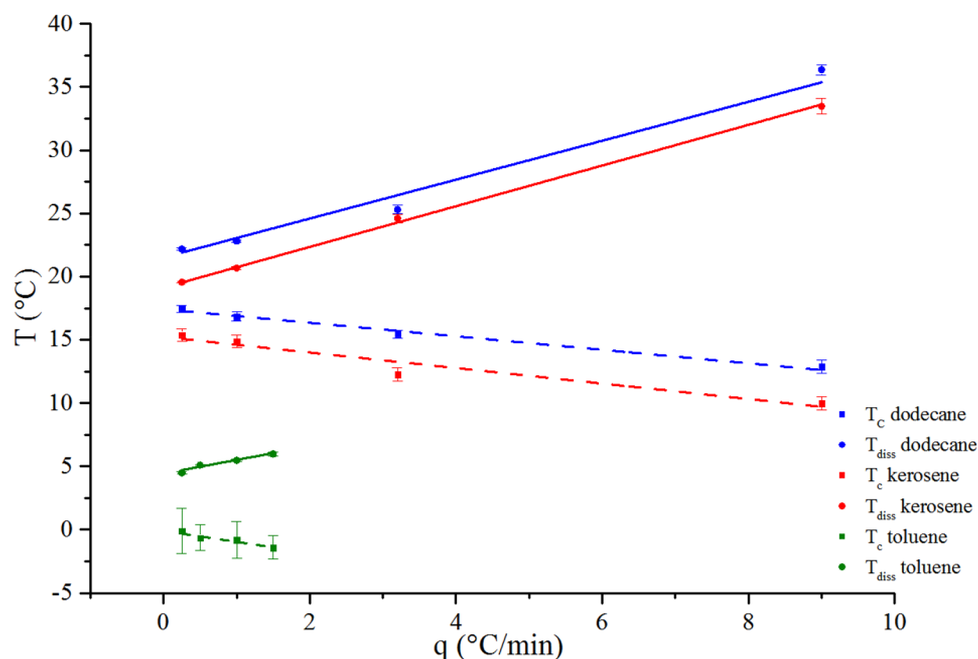


Figure 3. Crystallization T_c and dissolution T_{diss} temperatures as a function of cooling rate q for solution concentrations of 250 g/L for methyl stearate crystallizing from dodecane and kerosene solvents and of 192 g/L for methyl stearate crystallizing from toluene solvent.

Table 2. Solubility, Enthalpy, and Entropy of Dissolution and Mixing for Methyl Stearate in Three Different Solvents, Together with Corresponding Activity Coefficients^a

solvent	T (°C)	x_e	ΔH_{diss} (kJ/mol)	ΔS_{diss} (kJ/mol·K)	ΔH_{mix} (kJ/mol)	ΔS_{mix} (kJ/mol·K)	Y (20 °C)	$\ln Y = aT + b$
dodecane	18.70	0.132	63.55 ± 10.860	0.20 ± 0.037	9.61	0.03	2.01	$-0.013T + 0.96$
	21.09	0.160						
	22.18	0.186						
	24.12	0.210						
kerosene	17.25	0.123	69.80 ± 2.874	0.22 ± 0.098	15.87	0.05	1.85	$-0.022 T + 1.06$
	19.21	0.149						
	20.86	0.173						
	22.01	0.196						
toluene	1.90	0.052	59.84 ± 3.785	0.19 ± 0.014	5.90	0.02	1.14	$-0.009 T + 0.31$
	4.36	0.064						
	6.04	0.076						
	7.50	0.088						

^aParameter values obtained by modeling solubility data according to the van't Hoff plot. The errors of the slope and the intercept for enthalpy and entropy of dissolution refer to the 95% confidence interval.

molecules which can be quite strong interactions in the case of the benzene ring, due to the effect of London dispersion forces.³⁹ This effect appears to be manifested through the very low ΔH_{mix} in toluene solvent, consistent with the high released of energy associated with the formation of these solute–solvent bonds.

The noticeable dependence of activities on temperature for kerosene solutions can be evident on the steep slope of the van't Hoff line. Higher enthalpy and entropy of dissolution in this case are likely to be due to the range of different compounds present in kerosene, especially the aromatic molecules, which not only differ in size but will also complicate chemical interaction with the solute.

4.2. Nucleation Kinetics. Values of the relative critical undercooling u_c at the corresponding concentrations and cooling rates are presented in Table S1 of the SI. For each of the four solution concentrations within each solvent system, a plot of the cooling rate q (K/s) vs relative critical undercooling u_c in ln–ln coordinates was then constructed to obtain the slope of the straight line fitting these data points according to expressions 18 and 19.

Figure 5 presents an example of the plot obtained for a concentration of 250 g/L in dodecane and kerosene solutions and for a concentration of 192 g/L in toluene solutions. The best linear fittings to these data are given by $y = 4.21x + 13.41$, $y = 3.65x + 10.76$, and $y = 7.16x + 24.13$, respectively. The slope and the correlation coefficient R^2 of the best-fit straight line to the data for each concentration within the three solvent systems are presented in Table 3.

In all cases the slopes of the lines are higher than 3, suggesting that crystallization of methyl stearate proceeds via the PN mechanism. Thus, according to the KBHR approach, eq 4 should describe the experimental data plotted in $\ln q$ vs u_c coordinates. The values of a_1 , a_2 , and $\ln q_0$ parameters obtained, using OriginPro 8.5.1, are presented in Table 3. For each of the solution concentrations, within each solvent system these values along with the correlation coefficients for the fitting of eq 4 to the experimental data are given.

The best-fit curves between the experimental $u_c(q)$ values and those calculated from eq 4 were obtained by setting $a_1 = 3$.

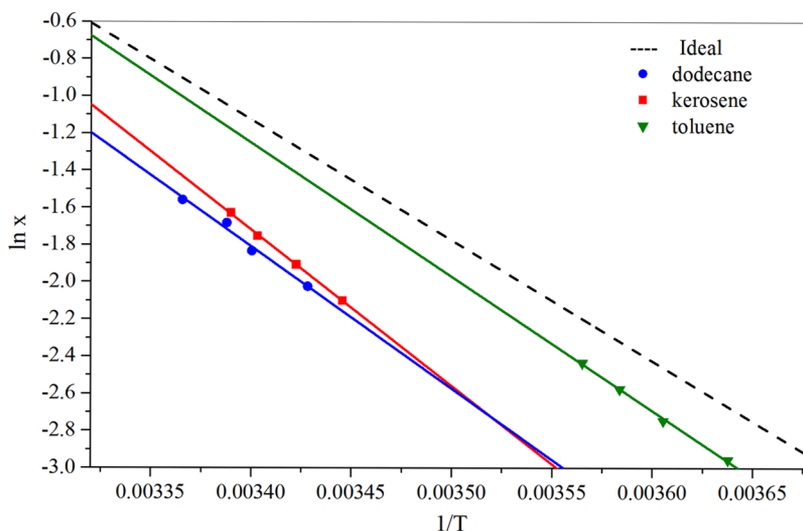


Figure 4. van't Hoff plot for methyl stearate in three different diesel type solvents. Solid lines represent experimental solubilities, and the dashed line represents the ideal solubility. Experimental solubilities were obtained by extrapolation to 0 °C of $T_{\text{diss}}(q)$ lines at four different solution concentrations.

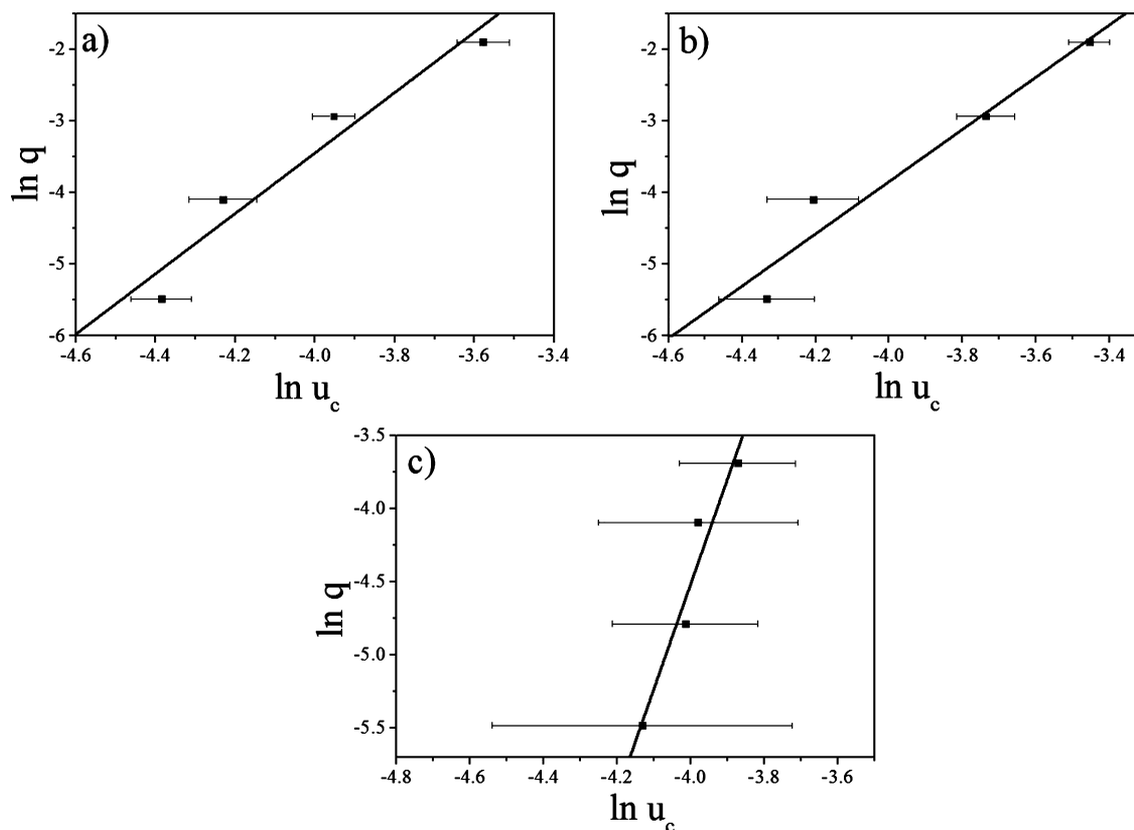


Figure 5. Experimental polythermal data in $\ln q$ vs $\ln u_c$ coordinates (a, b) for solution concentration of 250 g/L for methyl stearate crystallizing from (a) dodecane and (b) kerosene solvents and (c) for solution concentration of 192 g/L for methyl stearate crystallizing from toluene solvent.

Examples of such curves for the concentration of 250 g/L in dodecane and kerosene solutions and for a concentration of 192 g/L in toluene solution are presented in Figure 6.

Even though a much lower range of cooling rates was used for the toluene solvent (0.25–1.5 °C/min compared to 0.25–9 °C/min for the other two solvents), the low values of R^2 suggest that both relationships given by eqs 4 and 18 are not followed. Thus, in this particular case further analysis could be undertaken by solving numerically eq 8 in the SI. The numerical solution of this equation would allow nucleation kinetic parameters to be obtained

that would give a better fit regarding the dependence of the relative volume of crystals on the relative critical undercooling. Comparison of this trend line with the corresponding experimental data would provide an insight into whether this system can be better analyzed using this more rigorous approach when compared to the constraints inherent in the analytical solution. In particular, such constraints describe only the early stages of crystallization over which concentration can be assumed to remain virtually unchanged and hence the dependence of K_j and b on C and T can be effectively ignored. If this is the case, the solution

Table 3. Slopes of the Best Linear Fit to Data Points in $\ln q$ vs $\ln u_c$ Coordinates and Correlation Coefficients; Values of the Free Parameters a_1 , a_2 , and $\ln q_0$ Obtained from the Data Fitting in $\ln q$ vs u_c Coordinates According to eq 4 and Correlation Coefficients^a

concn (g/L)	slope $\ln u_c$ vs $\ln q$	R^{2b}	nucl mech	a_1	$a_2 = b$	$\ln q_0$	q_0 (K/s)	R^{2c}
dodecane								
200	3.50	0.96	PN	3	$7.44 \times 10^{-5} \pm 4.15 \times 10^{-5}$	9.29 ± 0.26	10812.30	0.98
250	4.21	0.94	PN	3	$2.19 \times 10^{-4} \pm 8.71 \times 10^{-5}$	9.35 ± 0.37	11548.60	0.97
300	3.89	0.89	PN	3	$1.39 \times 10^{-4} \pm 1.09 \times 10^{-4}$	9.54 ± 0.60	13936.82	0.92
350	4.30	0.90	PN	3	$3.30 \times 10^{-4} \pm 1.69 \times 10^{-4}$	9.10 ± 0.55	8917.04	0.94
kerosene								
200	4.92	0.95	PN	3	$5.21 \times 10^{-4} \pm 1.46 \times 10^{-4}$	8.98 ± 0.37	7967.05	0.97
250	3.65	0.94	PN	3	$1.32 \times 10^{-4} \pm 1.09 \times 10^{-4}$	8.64 ± 0.42	5639.21	0.95
300	4.22	0.99	PN	3	$2.62 \times 10^{-4} \pm 2.94 \times 10^{-5}$	8.70 ± 0.09	5998.83	0.99
350	3.92	0.95	PN	3	$2.39 \times 10^{-4} \pm 1.03 \times 10^{-4}$	8.67 ± 0.33	5847.80	0.97
toluene								
154	3.98	0.56	PN	3	$1.76 \times 10^{-4} \pm 4.08 \times 10^{-4}$	7.94 ± 1.26	2815.71	0.57
192	7.16	0.94	PN	3	$6.83 \times 10^{-4} \pm 2.04 \times 10^{-4}$	9.59 ± 0.64	14662.96	0.94
231	6.65	0.54	PN	3	$7.09 \times 10^{-4} \pm 8.21 \times 10^{-4}$	9.38 ± 2.35	11823.57	0.55
269	6.42	0.79	PN	3	$5.32 \times 10^{-4} \pm 3.72 \times 10^{-4}$	9.38 ± 1.26	11852.21	0.78

^aThe errors of the slope and the free parameters refer to the 95% confidence interval. ^bLinear fitting. ^cFitting eq 4.

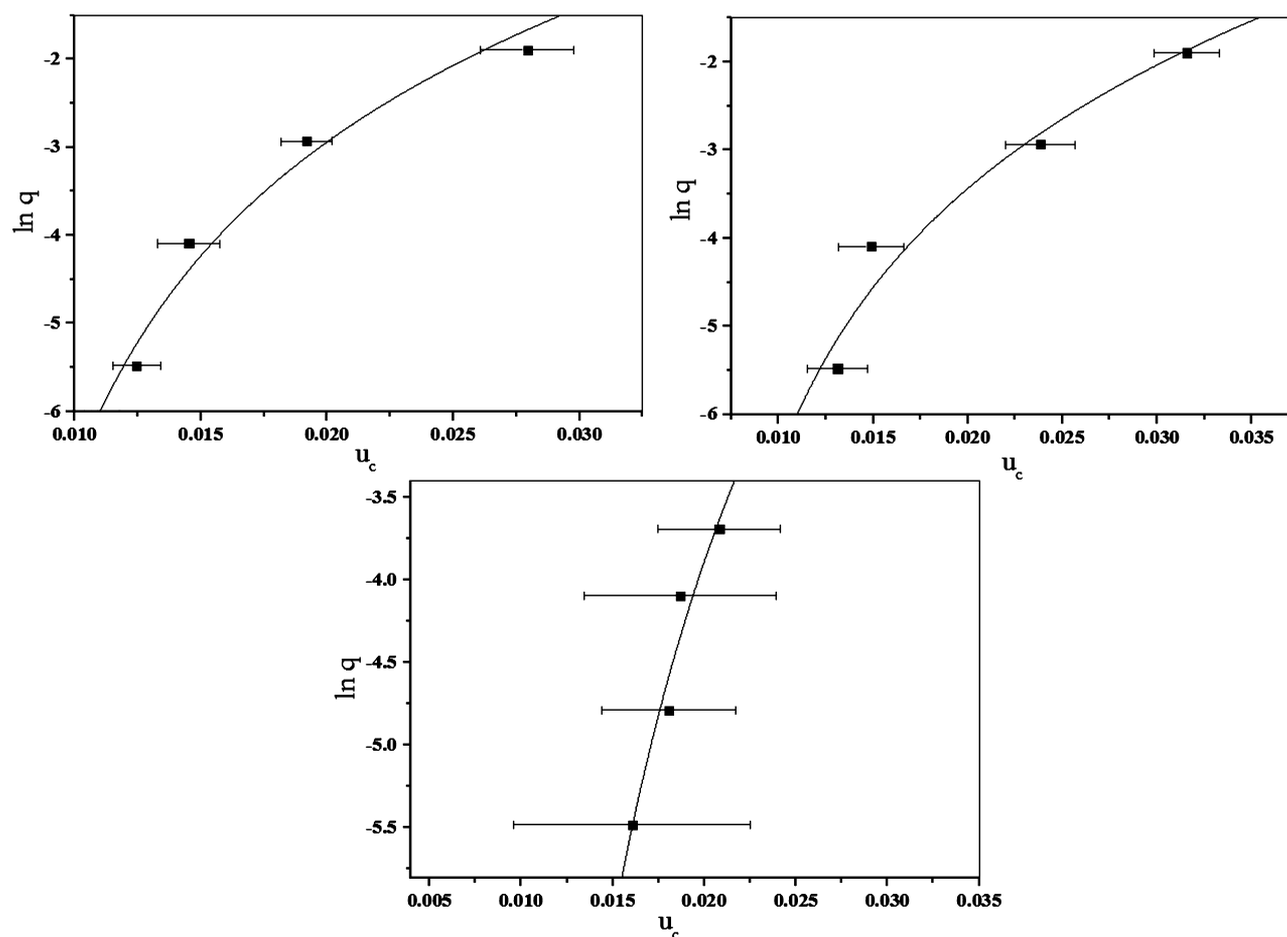


Figure 6. Increase in relative critical undercooling with the natural logarithm of the cooling rate. The points represent the data for crystallization of methyl stearate in solution with (top left) 250 g/L dodecane, (top right) 250 g/L kerosene, and (bottom) 192 g/L toluene; the lines illustrate the best fits according to eq 4.

viscosity could also be assumed to remain virtually unchanged on cooling.

Nonetheless, the analysis was still performed for toluene solutions for comparison with the other two solvents.

According to eq 8 $a_2 = b$, a dimensionless thermodynamic parameter defined by eq 10 from which γ_{eff} can be calculated. The

results obtained for $\ln q_0$ yield the values of q_0 , a parameter related through eq 9 to the nucleation rate constant K_j and the number N_{det} of crystallites at the detection point.

The effective interfacial tension γ_{eff} was evaluated from eq 10, using $v_0 = 0.491 \text{ nm}^3$,⁴¹ the calculated equilibrium temperature T_e , the shape factor $k_n = 16\pi/3$ for spherical nuclei, and the

Table 4. Nucleation Kinetics Parameters and Nucleation Rates for Methyl Stearate Crystallizing from Three Different Solvents at Four Different Solution Concentrations^a

solvent	x_c ($\frac{\text{mol of solute}}{\text{mol of solution}}$)	u_c	x_f	σ	γ (mJ/m ²)	r^* (nm)	i^*	N_{det}	J (nuclei/mL·s)
dodecane	0.132	0.010	0.10	0.32	0.94	0.84	5	2.43×10^{19}	6.70×10^{16}
	0.160	0.013	0.12	0.37	1.35	0.99	8	2.07×10^{19}	9.04×10^{16}
	0.186	0.011	0.13	0.38	1.16	0.98	8	2.57×10^{19}	1.06×10^{17}
	0.210	0.015	0.15	0.44	1.55	1.00	8	3.16×10^{19}	1.27×10^{17}
kerosene	0.123	0.017	0.07	0.66	1.91	0.94	7	2.64×10^{19}	1.24×10^{17}
	0.149	0.014	0.10	0.50	1.21	0.74	4	5.83×10^{19}	1.48×10^{17}
	0.173	0.015	0.11	0.55	1.52	0.84	5	5.11×10^{19}	1.79×10^{17}
	0.196	0.015	0.13	0.54	1.48	0.85	5	5.47×10^{19}	1.68×10^{17}
toluene	0.052	0.016	0.03	0.55	1.18	0.73	3	3.93×10^{19}	7.08×10^{16}
	0.064	0.016	0.04	0.49	1.86	1.17	14	1.08×10^{19}	4.56×10^{16}
	0.076	0.017	0.05	0.58	1.88	1.07	10	1.87×10^{19}	1.04×10^{17}
	0.088	0.015	0.06	0.49	1.71	1.13	12	1.64×10^{19}	6.53×10^{16}

^aThe critical radius, number of crystals at the detection point, and nucleation rates are calculated at u_c corresponding to T_c values obtained by the extrapolation to 0 °C/min of ($T_c(q)$) lines. x_c is the equilibrium solubility, u_c is the relative critical undercooling, x_f is the fraction of methyl stearate that remains in solution at the corresponding crystallization temperatures, σ is relative supersaturation, γ is interfacial tension, r^* is the critical nucleus radius, i^* is the number of molecules in the critical radius, N_{det} is the number of crystals at the detection point, and J is the nucleation rate.

molecular latent heat λ of crystallization estimated to be 1.06×10^{-19} , 1.16×10^{-19} , and 9.94×10^{-20} J for methyl stearate crystallizing from dodecane, kerosene, and toluene, respectively. These values were calculated assuming the enthalpy of dissolution obtained from the solubility data can be equated to the enthalpy of crystallization. In addition to this, the critical nucleus radius r^* and number i^* of molecules were obtained from eqs 21 and 22, respectively, calculated at u_c levels obtained using T_c values corresponding to the extrapolation to 0 °C/min of $T_c(q)$ lines, at each of the four concentrations within each solvent system. The results are given in Table 4.

The low values of the effective interfacial tension are an indication of a prevalence of the heterogeneous nucleation (HEN) mechanism for the nucleation of the methyl stearate crystallites regardless of the solution environment and are within the same order of magnitude of values reported earlier for *n*-alkanes.^{42–45}

The number of crystallites formed at the detection point N_{det} at a given crystallization temperature, can be obtained by performing a mass balance using the van't Hoff models derived for each solvent system. Thus, using the corresponding values of the mole fraction, the mass of C18:0 per unit volume in solution with the solvent can be obtained by solving the mass parameter in the molar fraction relationship. The mass of C18:0 in the solid phase per unit volume is therefore the difference between the mass of C18:0 in the initial solution and the mass in solution at the corresponding crystallization temperature. This value can be converted to volume by dividing the mass of the solute in the solid phase by the corresponding density.

Finally, the number of nuclei per unit volume, i.e., N_{det} as a function of solvent and solution concentration can be estimated from dividing the total volume of solid by the volume of a single nucleus. The latter can be obtained using the values of the critical radius assuming spherical nuclei.

Following this, the nucleation rate constants K_j and nucleation rates J can be calculated from eqs 9 and 23 respectively using $V = 1$, the corresponding equilibrium temperature T_e , values of the dimensionless thermodynamic parameter b , and values of the parameter q_0 for C18:0 as a function of solvent and concentration. A summary of these results is given in Table 4. The trend of these parameters is also shown in Figure S5 of the SI.

Figure 7 provides a comparison of the trend of nucleation rates (J) together with the critical radius (r^*) on an iso-supersaturation

basis. r^* values were calculated using the interfacial tension obtained for each of the solution concentrations studied at the corresponding σ levels.

4.2.1. Nucleation at the Detection Point. Nucleation rates are observed to be higher in kerosene where the supersaturation levels at which nucleation is detected are greater than in the other two solvents (Table 5). The lowest values of the rates obtained for toluene solvent can be associated with the higher interfacial tensions observed in this case, where the flatter nature of the slopes of the $T_c(q)$ lines shows that nucleation proceeds more progressively or is a more thermodynamically controlled process. This is likely to be the result of the highest solubility of C18:0 in toluene which would favor solute–solvent interactions over solute–solute ones. In addition to this, in toluene systems the fraction (x_c) of methyl stearate that remains in solution at the corresponding crystallization temperatures is lower in comparison to those in the other two solvent systems (Table 4), as crystallization occurs at lower temperatures in this case. Thus, given that interfacial tensions are inversely related to x_c , higher resistance to nucleation should be expected. In these solutions a sufficient level of supersaturation is then required to overcome the free energy for nucleus formation evidenced in the high levels of σ , which compares to those observed in kerosene where nucleation rates are at least 1 order of magnitude higher than in toluene.

Given that lower levels of interfacial tensions were observed in kerosene followed by dodecane solutions, it is likely that in these solvents nucleation is controlled, either by the attachment frequency f^* or the concentration of nucleation sites C_0 . This could indicate that either volume diffusion or interface transfer of building units (molecules) would be rate limiting according to expressions 24 and 25. In the case of kerosene solvent, these conclusions are in line with a higher range of σ at which higher x_c values were observed in comparison to dodecane solutions, allowing for both sufficient driving force nucleation and solute availability in solution that would reduce interfacial tension.

Interestingly, although values of interfacial tension similar to those of kerosene systems were obtained in dodecane solutions, the nucleation rates in the latter case were significantly lower by 1 order of magnitude. Given the expected lower strength of both solvent–solvent and solvent–solute intermolecular interaction, due to the nonpolar nature of dodecane molecules, the diffusion of solute molecules should not be rate limiting and therefore

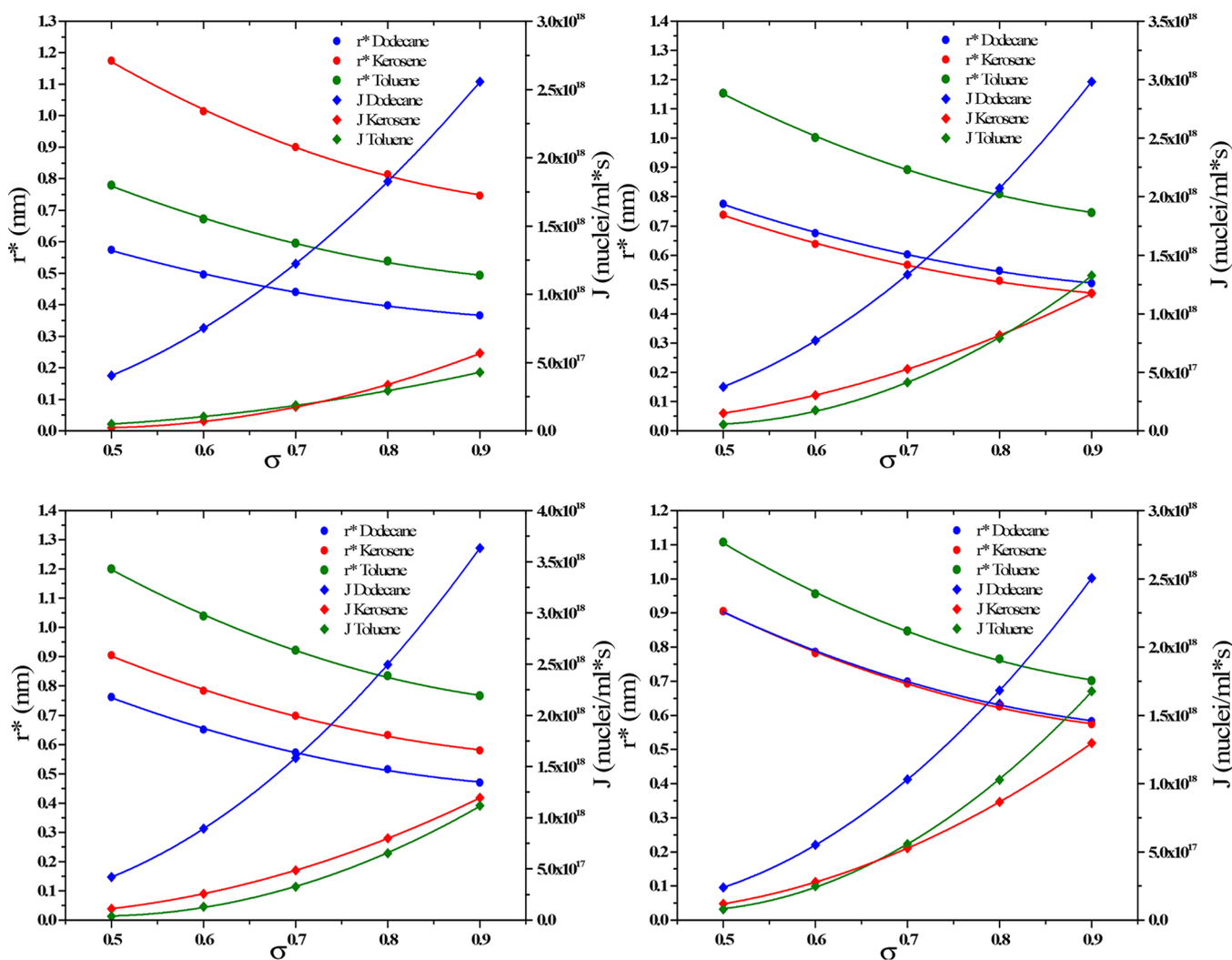


Figure 7. Tendency of critical radius (r^*) and nucleation rates (J) as a function of supersaturation (σ) for C18:0 growing from dodecane, kerosene, and toluene solvents using parameters derived for the range of concentrations studied. Concentration increases from left to right and from top to bottom.

sufficient levels of attachment frequency should be possible to reach. Based on these observations and accounting for the definition of nucleation rate given by eq 23, it is likely that in these systems nucleation rates are then hindered by the availability of nucleation sites C_0 , due to the lower solubility of methyl stearate in dodecane that would allow a lower amount of solute per unit volume.

4.2.2. Nucleation at the Same Levels of Supersaturation (σ).

If the analysis is performed comparing nucleation parameters at equal levels of solution supersaturation (Figure 7), nucleation rates are higher in dodecane followed by kerosene and toluene solvent. The differences between dodecane and both kerosene and toluene solution rates increase with an increase in σ up to even 1 order of magnitude at supersaturations higher than 80%. In general, the critical nucleus radius is higher in toluene followed by kerosene and dodecane solutions, although this trend changes at the lowest and highest solution concentrations studied in which kerosene solutions show the highest and lowest values of this parameter, respectively.

Given that the critical nucleus radius is directly related to interfacial tension, these observations confirm that interfacial tensions are rate limiting in toluene solvent over all the range of supersaturations chosen. Interfacial tensions in kerosene are closer in magnitude to those observed in dodecane solvent, while delivering

nucleation rates closer in magnitude to those observed in toluene solutions. The latter complement the discussion presented in section 4.2.1 in the sense that this effect could be associated with the complex composition of kerosene, which is comprised of molecules of different types including paraffins, cycloalkanes, and aromatics, that could hinder the diffusion processes associated with molecular attachment into the nucleus and therefore decrease the attachment frequency (f^*).

Interfacial tensions were observed to be the highest in kerosene at the lowest solution concentration becoming closer to those observed in dodecane and below those of toluene solutions as solution concentration increases. This could be associated with an increase in the thermodynamic barrier to nucleation at lower concentrations where solute–solvent interactions would be favored.

In the case of dodecane solutions the lowest values of interfacial tensions are in line with a more kinetically controlled process as discussed in section 4.2.1. The lower values of this parameter can be associated with the lower solubility of C18:0 in dodecane that favor solute–solute interactions. Likewise, attachment frequencies should be favored by an easy desolvation process due to the low solute–solvent strength interactions indicating that nucleation is likely to be hindered by the low values of C_0 .

The differences in the tendency observed when nucleation is analyzed at the detection points in comparison to the same σ

Table 5. Summary of Parameters Obtained through the Combined Assessment of Solubility and Nucleation Kinetics of Methyl Stearate in Three Different Solvents^{a,b}

assessed criteria	assessed parameters	dodecane			kerosene		toluene		conclusion
		lowest	intermediate	higher	intermediate	higher	intermediate	higher	
solubility	solubility level								All systems behave less than ideal as activities are higher than 1. However, higher values in dodecane followed by kerosene solutions indicate either solute–solvent or solvent–solvent interactions are favored in these systems. Solute–solvent interactions are similar to those of solute–solvent in toluene solutions with activities close to 1; this is further supported by the lowest enthalpy of mixing. The highest values of both enthalpy of dissolution and mixing in kerosene solution evidence strong dependence of activities on temperature.
	γ (20 °C)	2.01	1.85	1.14	1.85	1.14	1.14	1.14	
	ΔH_{diss} (kJ/mol)	63.55	69.81	59.84	69.81	59.84	59.84	59.84	
	ΔS_{diss} (kJ/mol·K)	0.20	0.22	0.19	0.22	0.19	0.19	0.19	
	ΔH_{mix} (kJ/mol)	9.61	15.87	5.90	15.87	5.90	5.90	5.90	
nucleation	ΔS_{mix} (kJ/mol·K)	0.03	0.05	0.02	0.05	0.02	0.02	0.02	Relatively higher values of interfacial tensions in toluene seem to hinder nucleation in this solvent. Although interfacial tension values are close in dodecane and kerosene solutions, nucleation in the former is 1 order of magnitude lower. This could indicate less available nucleation sites in these solutions, due to the low solubility of methyl stearate in dodecane.
	σ range	0.32–0.44	0.50–0.66	0.49–0.58	0.50–0.66	0.49–0.58	0.49–0.58	0.49–0.58	
	γ (mJ/m ²) range	0.94–1.55	1.21–1.52	1.18–1.88	1.21–1.52	1.18–1.88	1.18–1.88	1.18–1.88	
	r^* (nm) range	0.84–1.00	0.74–0.94	0.73–1.17	0.74–0.94	0.73–1.17	0.73–1.17	0.73–1.17	
	J (nuclei/mL·s) range	6.70×10^{16} – 1.27×10^{17}	$(1.24$ – $1.79) \times 10^{17}$	4.56×10^{16} – 1.04×10^{17}	$(1.24$ – $1.79) \times 10^{17}$	4.56×10^{16} – 1.04×10^{17}	4.56×10^{16} – 1.04×10^{17}	4.56×10^{16} – 1.04×10^{17}	

^a γ is the activity coefficient; ΔH_{diss} and ΔS_{diss} are the enthalpy and entropy of dissolution, respectively; ΔH_{mix} and ΔS_{mix} are the enthalpy and entropy of mixing, respectively; γ is the interfacial tension; r^* is the nucleus critical radius; and J is the nucleation rate. ^bThese results are presented together with those obtained for the assessment of morphology and crystal growth kinetics, for the same solutions' systems, in ref 46.

levels suggests that the complex composition of kerosene solvent significantly influences the nucleation process. In this case the roles of thermodynamics and kinetics are more interconnected due to the different types of intermolecular interactions that would affect desolvation and diffusion process as well as solubility.

A summary of the obtained solubility and nucleation kinetics parameters as well as some conclusions related to each section are given in Table 5.

5. CONCLUSIONS

The solubility and nucleation of methyl stearate crystallizing from dodecane, kerosene, and toluene were studied. Solutions of C18:0 in all cases show lower solubility than that of an ideal solution with the highest solubility obtained in toluene followed by kerosene and dodecane solvents. The analysis showed in all cases a progressive nucleation mechanism and crystallite interfacial tension (γ_{eff}) values between 0.94 and 1.55 mJ/m², between 1.21 and 1.52 mJ/m², and between 1.18 and 1.88 mJ/m² for methyl stearate crystallizing from dodecane, kerosene, and toluene, respectively. Nucleation rates calculated using the obtained values of γ_{eff} and the number of crystals at the detection point (N_{det}) ranged between 4.56×10^{16} and 1.79×10^{17} nuclei/mL·s, with the highest rates predicted for methyl stearate crystallizing from kerosene solutions. This trend changes when the analysis is performed at the same σ levels where nucleation rates were the highest in dodecane solvent. This is effect is thought to be due to the complex nature of kerosene in which the interplay between desolvation, diffusion process, and solubility plays a more important role.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.energyfuels.7b03212.

Comparative schemes for the isothermal or polythermal method based on a concentration vs temperature profile (section 1), full derivation of the set of expressions that comprise the KBHR approach (section 2), kerosene *n*-alkane chain length distribution (section 3), temperature calibration lines for the Crystal 16 unit (section 4), flowchart describing how to apply the KBHR approach for the analysis of nucleation kinetics from polythermal experimental data (section 5), experimental crystallization T_c and dissolution T_{diss} temperatures as a function of cooling rate q (section 6), and comparative figures of nucleation kinetics parameters for methyl stearate crystallizing from dodecane, kerosene, and toluene solvents at experimental crystallization temperatures (section 7) (PDF) The research data supporting this publication can be accessed at: <https://doi.org/10.5518/336>.

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The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors gratefully acknowledge Infineum Ltd. for the funding of this research which forms part of the doctoral studies of one of us (D.M.C.). We also gratefully acknowledge the U.K.'s EPSRC for the support of nucleation and crystal growth research at Leeds and Manchester through funding the Critical Mass Project: Molecules, Clusters and Crystals (Grants EP/IO14446/1 and EP/IO13563/1). The research on the analysis of nucleation data using the polythermal method was inspired by Prof. Dimo Kashchiev (Institute of Physical Chemistry, Bulgarian Academy of Sciences) during his sabbatical leave in Leeds (Leverhulme Trust, Grant F10100A). We are most grateful to him for his insightful contribution to this research area.

NOMENCLATURE

Symbols

a = dimensionless molecular latent heat of crystallization
 a_c = solution activity
 A^* = surface area of nucleus (m^2)
 b = dimensionless thermodynamic parameter
 C_{nuc}^* = equilibrium nucleus concentration (m^{-3})
 C_0 = concentration of nucleation sites or instantaneously nucleated crystallites (m^{-3})
 f^* = attachment frequency of monomers to the nucleus
 J = nucleation rate ($\text{m}^{-3} \text{s}^{-1}$)
 K_J = nucleation rate constant ($\text{m}^{-3} \text{s}^{-1}$)
 k_n = nucleus numerical shape factor
 n = crystallite growth exponent
 N_{det} = detectable number of crystallites
 q = cooling rate (K s^{-1})
 q_0 = parameter in $u_c(q)$ dependence for both PN and IN (K s^{-1})
 Q = parameter in eq 19, related to q_0
 T = solution temperature (K)
 T_c = crystallization temperature (K)
 T_{diss} = equilibrium dissolution temperature (K)
 T_e = solution saturation (or equilibrium) temperature (K)
 ΔT_c = critical undercooling for crystallization (K)
 v_0 = volume of solute molecule in crystal (m^3)
 V = volume of solution (m^3)
 x = mole fraction of solute in solution
 x_e = equilibrium mole fraction
 x_{ideal} = ideal equilibrium mole fraction
 X_1 = concentration of colliding building units
 Z = Zeldovich factor
 ΔH_m = molal enthalpy of melting (J mol^{-1})
 ΔS_m = molal entropy of melting ($\text{J mol}^{-1} \text{K}^{-1}$)
 ΔH_{diss} = molal enthalpy of dissolution (J mol^{-1})
 ΔS_{diss} = molal entropy of dissolution ($\text{J mol}^{-1} \text{K}^{-1}$)
 ΔH_{mix} = molal enthalpy of mixing (J mol^{-1})
 ΔS_{mix} = molal entropy of mixing ($\text{J mol}^{-1} \text{K}^{-1}$)
 i^* = number of molecules in critical nucleus
 r^* = critical nucleus radius (m)
 σ = relative supersaturation
 σ_{crit} = critical relative supersaturation
 u_c = relative critical undercooling for crystallization
 Υ = activity coefficient
 γ_{eff} = effective interfacial tension of crystal nucleus in 3D HEN (mJ m^{-2})
 λ = molecular latent heat of crystallization (J)
 ρ = density
 ξ = sticking coefficient

ω = positive number in expression 19

ω^* = transfer frequency of building units from adsorbed to integrated

Abbreviations

HEN = heterogeneous nucleation
 HON = homogeneous nucleation
 IN = instantaneous nucleation
 KBHR = Kashchiev–Borissova–Hammond–Roberts
 MSZW = metastable zone width
 PN = progressive nucleation
 SD = standard deviation
 3D = three-dimensional

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