Morphology and Growth of Methyl Stearate as a Function of Crystallization Environment

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ABSTRACT: In situ studies of methyl stearate growing from supersaturated n-dodecane, kerosene, and toluene solutions reveal strong evidence that solvent choice influences the crystal morphology and crystal growth kinetics. Crystals with similar habit are observed in all solvents, with the exception of lower supersaturations in kerosene, where a less symmetric morphology was observed. BFDH analysis based on the monoclinic C2 crystal structure of methyl stearate yielded the morphological indexation to be (110), (1−10), (−110), and (−1−10) for the dominant observed habit and (110) (1−10) (−1−10) (−240) (−3−10) for the less symmetric habit observed in kerosene solvent. Measurements of the growth rate for the (110) and (1−10) faces are similar for all solutions ranging from 0.02 to 1.13 μm/s, for significantly lower values of supersaturation in the case of toluene. The tendency of the growth rate dependence on σ was consistent with the Burton-Cabrera-Frank (BCF) growth mechanism in n-dodecane, the Birth and Spread (B&S) mechanism in kerosene and diffusion controlled in toluene solvent.

1. INTRODUCTION

Wax formation in diesel and biodiesel fuels at low temperatures is one of the major problems faced by the fuels industry as crystallization of the saturated compounds present in these solutions can plug up filters and obstruct pipelines. Preventing wax formation in these multicomponent mixtures requires a good understanding of the crystallization behavior of saturated compounds such as fatty acid methyl esters (FAMEs) present in biodiesel fuel. Notably, growth kinetics and morphological studies can provide the basis to develop adequate technology, such as additives, that can modify the crystals’ habit to allow for the fuel’s adequate flow and settlement properties.

Table 1. Composition of Kerosene from 2D Gas Chromatography Analysis Performed by Infineum UK

| hydrocarbon | mass % |
|-------------|--------|
| paraffins   | n-alkanes | 16.29 |
|             | iso-paraffins | 23.04 |
| cycloalkanes| naphthenes | 42.40 |
| aromatics   | alkyl benzenes | 7.60 |
|             | benzylocycloparaffins | 6.80 |
|             | naphthalenes | 3.43 |
|             | biphenyls/acenaphthenes | 0.30 |
|             | fluorenes | 0.15 |

Relevant progress has been made in this area including: the use of molecular dynamics (MD) to study crystal nucleation and growth and the prediction of crystal’s growth from the assessment of the solid/solution interface molecular structure. The prediction of the crystal’s habit, using the extended interface structural analysis (EISA) shows that interfacial molecular events can be efficiently incorporated to assess crystal growth and morphology when cosolvents, impurities, and additives are used. Similarly, molecularly based simulations have been used to control and predict the crystal shapes of organic compounds based on energetic and crystal structure features, e.g., the control of AmB crystal shapes using tailor-made additives; the effect of hydroxyl-propyl methylcellulose (HPMC) concentration and solvent on the crystal habit of Nifedipine (Nif); the prediction of the shape of para-amino benzoic acid (PABA) crystals, and 6-methyl-2-thiouracil. Methodologies for the study of face specific growth kinetics include novel techniques such as the use of a growth cell coupled with phase contrast microscopy and stereovision imaging to analyze the effect of solution flow on crystal growth. The publication by Van Driessche A.E.S. et al. provides a thorough review of measuring techniques for the study of crystal growth kinetics.

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Despite the developments in this area, thus far there have been no relevant fundamental studies for crystallization of saturated methyl esters, such as methyl palmitate and stearate, due to the complex nature of crystallization of these materials. Morphological and growth kinetic studies in saturated methyl esters are scarce, not only due to the difficulty of obtaining reliable crystallographic information for these compounds, but also due to the difficulty of growing observable faceted crystals.

Limited structural information, obtained through powder X-ray diffraction on methyl stearate, suggests this compound crystallizes in two polymorphic forms: monoclinic and orthorhombic. The unit cell parameters of the stable monoclinic form are thought to be $a = 5.61$ Å, $b = 7.33$ Å, $C = 106.6$ Å, $\beta = 116.47^\circ$ with crystal space group $A2/\alpha$. Similarly the unit cell parameters of the orthorhombic form are $a = 5.61$ Å, $b = 7.35$ Å, $C = 95.15$ Å with crystal space group $Pnab$. Observations of methyl stearate crystals have shown that this compound crystallizes in a plate-like morphology in which the expected dominant (001) face is believed to grow via a screw dislocation mechanism because of spiral growth on this face.\(^{17}\)

Given the lack of studies on the crystallization of saturated methyl esters, it is the aim of this study to deliver fundamental information on the morphology and crystal growth kinetics of methyl stearate as a function of solution environment. These findings are complemented and cross-correlated in section 3.3 with unpublished results,\(^{29}\) where a fundamental analysis of methyl stearate nucleation was carried out.

2. MATERIALS AND METHODS

2.1. Materials. The solute methyl stearate and solvents $n$-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 Infinium Ltd. Its hydrocarbon composition and $n$-alkanes chain length distribution is summarized in Table 1 and Figure 1 respectively.

2.2. Equipment and Experimental Procedure. In situ crystal growth studies were carried out using an experimental setup comprising an optical microscope (Olympus BX51), operated in differential...
Table 2. Parameters Used for the Collection of Crystals Micrographs for Methyl Stearate Crystallizing in n-Dodecane, Kerosene, and Toluene Solutions

|        | C(methyl stearate) (g/L) | MSZW (°C) | temperature (°C) range | supersaturation (σ) range |
|--------|--------------------------|-----------|-------------------------|--------------------------|
| n-dodecane | 350                      | 4.3       | 19.8–24.1               | 20.2 to 21.0             | 0.30 to 0.39               |
| kerosene   | 350                      | 4.3       | 17.7–22.0               | 17.8 to 18.3             | 0.45 to 0.52               |
| toluene    | 538                      | 4.3       | 10.2–14.5               | 13.5 to 13.9             | 0.04 to 0.08               |

The supersaturation required for crystallization was created by reducing the solution temperature from the equilibrium temperature (T_eq) to different chosen temperatures within the metastable zone. Although the supersaturation is set by decreasing the solution temperature, circulating water through the cell, the growth of the crystals is only measured once the targeted temperature has been established. The supersaturation level at each temperature is calculated using expression 1:

\[ \sigma = \frac{x}{x} - 1 \]  

where \( x \) is the solution concentration and \( x \) is the molar fraction of the solute in the solution at equilibrium, obtained from the van’t Hoff equation at the temperature of measurement.

The crystal morphology and subsequent growth of the observed crystals was followed by recording images at equal time-intervals, every 5–20 s depending on the speed of the crystal growth in each system. The growth rates of the individual faces (G) were obtained by following the increase with time of the normal distance from the center of the projected two-dimensional (2D) crystal to the faces as shown in Figure 3. The crystal centers were defined by drawing lines that connected the crystal corners defined by the two most important observed faces. Ten to thirteen measurements of the normal distance increase were recorded.

Due to the experimental challenges involved, notably due to these crystallization systems having a very small metastable zone width (MSZW), it was not feasible at this stage of experimental methodology and data acquisition to make a statistically significant number of measurements for these systems (only two growth measurements were possible at each of the selected supersaturations). However, for a less challenging system such as ibuprofen,\(^1\) this has been assessed previously where the standard deviation between measurements was found to be quite low. Similarly, the methodology has shown to be reproducible as it has been previously applied in the assessment of the growth kinetics of other organic molecules such as \( n \)-docosane\(^12\) and \( \alpha \)-para-aminobenzoic acid (PABA).\(^3\)

2.3. Data Analysis. 2.3.1. Morphological Predictions. The morphological analysis was carried out making use of a methodology presented elsewhere.\(^1\) This methodology relies on the iteratively predicted Bravais-Friedel-Donnay-Harker (BFDH) morphology using pairwise Miller indices and comparing these predictions with the micrographs obtained experimentally. A summary of this methodology is given in Figure 1 in the Supporting Information (SI) to this paper.

2.3.2. Crystal Growth Kinetics. Given the experimental method used to collect crystal growth rates, the measured growth rates are influenced not only by the incorporation of growth units into the crystal surface, but also by the diffusion of the growth units within the bulk of the solution. Thus, growth models that combine these two effects acting in series have been derived as part of this work.\(^18\) This uses an analogy to a circuit as shown in Figure 2 of the SI delivering specific kinetics models for the dependence of growth rate (G) on supersaturation as described by a power law,\(^19\) the Birth and Spread (B&S) and Burton-Cabrera-Frank (BCF) models.\(^20\) These models are given by expressions 2, 3, and 4, respectively. A value of \( r = 1 \) in expression 4 corresponds to the case of rough interface growth (RIG).\(^3\,2\) The complete derivation of these models is presented in section 3 of the SI.
Table 3. List of the Predicted Zone Axis Defining the Different Zone Groups, Representative (hkl) Planes and \(d_{hkl}\) Spacing within Each Group

| Orthorhombic (Pnab) group | zone axis [uvw] | representative plane (hkl) | interplanar distance \(d_{hkl}\) |
|---------------------------|----------------|-----------------------------|-----------------|
| 1                         | (100)          | (011)                        | 7.33            |
|                           |                | (020)                        | 3.68            |
|                           |                | (031)                        | 2.45            |
|                           |                | (042)                        | 1.84            |
|                           |                | (064)                        | 1.22            |
| 2                         | [1–10]         | (111)                        | 4.46            |
|                           |                | (220)                        | 2.23            |
|                           |                | (331)                        | 1.49            |
| 3                         | [2–10]         | (120)                        | 3.08            |
| 4                         | [3–10]         | (131)                        | 2.25            |
|                           |                | (260)                        | 1.12            |
| 5                         | [0–10]         | (200)                        | 2.81            |
|                           |                | (602)                        | 0.94            |
| 6                         | [1–20]         | (211)                        | 2.62            |
|                           |                | (420)                        | 1.31            |
| 7                         | [3–20]         | (231)                        | 1.85            |
|                           |                | (460)                        | 0.92            |
| 8                         | [1–30]         | (311)                        | 1.81            |
|                           |                | (620)                        | 0.91            |

| Monoclinic A2/a group | zone axis [uvw] | representative plane (hkl) | interplanar distance \(d_{hkl}\) |
|-----------------------|----------------|-----------------------------|-----------------|
| 1                     | (100)          | (011)                        | 7.33            |
| 2                     | [1–10]         | (111)                        | 4.46            |
| 3                     | [2–10]         | (120)                        | 3.08            |
| 4                     | [3–10]         | (131)                        | 2.25            |
| 5                     | [0–10]         | (200)                        | 2.81            |

Figure 5. Example of initial designation of crystal faces for the two morphologies observed for methyl stearate.

\[
G' \left( \frac{m}{s} \right) = \frac{1}{k_m} + \frac{1}{k_f(\sigma)m^{\frac{1}{2}}(\sigma)}
\]  

(4)

where \(\sigma\) is the solution’s relative supersaturation, \(k_f\) is the growth rate constant, \(r\) is the growth exponent in the RIG interface growth kinetic model, and \(A_1\) and \(A_2\) are thermodynamic parameters in the B&S and BCF interface growth kinetic models, respectively. \(k_m^{\prime}\) is related the coefficient of mass transfer within the bulk of the solution, \(k_M T\) through expression 5

\[
k_M T \left( \frac{m}{s} \right) = \frac{k_M T C MW}{\rho_s}
\]

(5)

In this expression \(\rho_s\) is the solute density, \(MW\) the solute molecular weight, and \(C\) the equilibrium concentration (solubility).

3. RESULTS AND DISCUSSION

3.1. Morphological Observations as a Function of Solution Environment and Crystal Indexing. Using the setup for the measurements of crystal growth rates, images of C18:0 crystals growing from three solvents were obtained. Depending on the initial assessment of the solution behavior, different ranges of supersaturations within the MSZW were chosen for this analysis. In the case of \(n\)-dodecane solutions, growth was difficult to observe above 21 \(^\circ\)C (\(\sigma < 0.3\)) and would occur too fast to be recorded below 20.2 \(^\circ\)C (\(\sigma > 0.39\)). Similarly for kerosene
solutions growth was difficult to observe above 18.3 °C (σ < 0.45) and would occur too fast below 17.8 °C (σ > 0.52). For toluene systems the growth would occur very fast even at temperatures close to the solubility line and therefore the range of assessed supersaturations was limited. A summary of the width of the metastable zone at the corresponding solution concentration, together with the parameters used during the growth measurements for each system, is presented in Table 2.

Selected micrographs of the crystals observed in the range of supersaturations studied for the three solvents are shown in Figure 4.

In general the crystals observed in all cases were very thin plate-like crystals. The assessment of C18:0 crystal morphology reveals that the crystal habit does not change with solvent except for the crystal habits observed in kerosene at the lowest supersaturations (σ). Although the habit observed at higher supersaturation in kerosene solutions is more elongated when compared to those obtained in the other two solvents, the internal angles of the observed crystals are equal in all cases. The lozenge shape of these crystals could suggest that their morphology is based on an orthorhombic crystal system where these pairs of faces are coincidental. However, due to the geometry of the space group the prediction for the crystal habit does not change with solvent except for the crystal habits observed in kerosene at the lowest supersaturations (σ). Although the habit observed at higher supersaturation in kerosene solutions is more elongated when compared to those obtained in the other two solvents, the internal angles of the observed crystals are equal in all cases. The lozenge shape of these crystals could suggest that their morphology is based on an orthorhombic crystal system where these pairs of faces are coincidental.

The prediction of the morphology for C18:0 crystals was carried out following the procedure presented in Figure 1 of the SI, using three different crystal structures that have been suggested for methyl stearate. The first two crystal structures correspond to monoclinic and orthorhombic crystal systems with space group A2/a and Pnab respectively and the third one to the monoclinic crystal system with space group C2. In the latter the unit cell parameters are \( a = 5.60 \text{ Å}, b = 7.39 \text{ Å}, c = 47.96 \text{ Å}, \) and \( \beta = 91.15^\circ \) (I. More/Infinium UK, personal communication, July 25, 2014).

BFDH analysis revealed 49, 81, and 147 habit planes for orthorhombic, monoclinic (A2/a), and monoclinic (C2) crystal systems, respectively. The expected (001) plane is obtained in all predictions for the indexation of the larger crystals face perpendicular to the c axis. According to the methodology (Figure 1 SI), all other resulting indices were arranged by grouping the lattice planes into 9 zones for the first two crystal systems and into 17 zones for the latter. These groups with their corresponding zone axes indexation, including the complete set of planes delivered by the BFDH prediction are given in section 2 of the SI. Table 3 shows a summary of this information featuring some representative planes and corresponding \( d \) spacing within each group.

The interplanar angle correlation for the plane pairs was carried out using the annotation in Figure 5 both for the morphology observed at the lowest supersaturation in kerosene and for the dominant morphology observed in all solvents.

As for the case of orthorhombic Pnab and monoclinic A2/a crystal systems, the list of Miller indices obtained belong to the same zone axes groups, and thus a single analysis was carried out. This delivered only two pairs of Miller indices matching AB interplanar angles (111) and (1−11) or (011) and (311). However, due to the geometry of the space group the prediction for the pairwise (111) (1−11) is the only one delivering the expected crystal lozenge shape. The same analysis using the monoclinic C2 crystal system delivered similar results with the pairs (110) and (1−10) or (240) and (2−40) matching the internal angle between A and B. This information is presented in Table 4.

In the case of the indexation for the morphology observed at the lowest supersaturation of kerosene solutions, the set of matches for the pairs of Miller indices of the remaining interplanar angles between BC/CD and BD faces are summarized also in Table 4. Miller indices for the BC and CD faces are presented within the same column as the interplanar angles between these pairs of faces are coincidental.

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Table 4. Likely Combinations of Miller Indices for the Four Faces of the Experimental Crystals Observed at the Lowest Supersaturation σ < 0.45 in Kerosene Solvent

| A       | B       | C       | D       |
|---------|---------|---------|---------|
| (110)   | (1−10)  | (3−10)  | (460)   |
| (110)   | (1−10)  | (4−20)  | (460)   |
| (110)   | (1−10)  | (6−40)  | (460)   |

The row highlighted in the table represents the chosen final unique solution using monoclinic C2 crystal system.
Within the morphology delivered by the orthorhombic/monoclinic \( A_2/a \) crystal systems there is only a pair of Miller indices that match the interplanar angles between BD faces, and one pair for AB faces, that also satisfies the faces present in the dominant lozenge shape morphology. This facilitates obtaining the solution for the entire morphology after cross-checking with

Figure 6. Predicted BFDH morphology of methyl stearate crystals using the Miller indices in the obtained unique solutions and comparison with the crystal micrographs: (a) dominant morphology observed in three solvents, (b) predicted morphology using orthorhombic/monoclinic \( A_2/a \) crystal system for kerosene crystals observed above 18.3 °C (\( \sigma < 0.45 \)), and (c) predicted morphology using monoclinic \( C_2 \) crystal system for kerosene crystals observed above 18.3 °C (\( \sigma < 0.45 \)).
the indices allocated for BC/CD pairs. If AB faces are allocated the pair (111) (1\textsuperscript{−}11) and BD faces the pair (011) (211) then the solution for BC/CD pairs should contain either the Miller indices (011) or (211) together with either the (111) or (1\textsuperscript{−}11) indices. Therefore, the solution for the entire morphology is given by the indices (111) (1\textsuperscript{−}11) (011) (211). Further modification of the perpendicular distance from the center of the crystal to the faces, produce the morphology prediction with lengths proportional to those observed experimentally.

Among the Miller indices pairs delivered by the prediction using the monoclinic C\textsubscript{2} crystal system, only three of these pairs matched BD face interplanar angles and two pairs matched those angles between AB faces. Using the same principle as in the previous analysis, if AB pairs are given by the (110) (1\textsuperscript{−}10) indices, then three options would be available for BC/CD faces as they have to contain either the (110) or (1\textsuperscript{−}10) indices: (110) (3\textsuperscript{−}10), (110) (4\textsuperscript{−}20), and (110) (6\textsuperscript{−}40) and two pairs would likely satisfy BD faces (110) (4\textsuperscript{−}60) and (240) (3\textsuperscript{−}10). This is so because by cross-checking, these pairs share either of their two indices. Similarly, if AB pairs are given by (240) (2\textsuperscript{−}40), then BD would be given by the pair (240) (3\textsuperscript{−}10) and BC/CD faces would be likely indexed by the (110) (3\textsuperscript{−}10) pair.

A summary of the unique potential morphology solutions using these indices combinations is given in Table 5.

After performing the morphological predictions using the combinations in Table 5, the analysis suggests the indexation (110) (1\textsuperscript{−}10) (3\textsuperscript{−}10) (240) for the morphology observed at the lowest supersaturations in kerosene. Further modification of the perpendicular distance from the center of the crystal to the faces, in the corresponding cif file, produce the morphology prediction with lengths proportional to those observed experimentally.

| solvent | σ | no. crystals | mean growth rate G (μm/s) |
|---------|---|--------------|--------------------------|
| n-dodecane | 0.39 | 2 | (110) 1.13 | (1\textsuperscript{−}10) 1.05 |
|           | 0.38 | 2 | 0.66 | 0.70 |
|           | 0.37 | 2 | 0.64 | 0.63 |
|           | 0.36 | 2 | 0.34 | 0.34 |
|           | 0.34 | 2 | 0.31 | 0.34 |
|           | 0.33 | 2 | 0.25 | 0.26 |
|           | 0.32 | 2 | 0.14 | 0.14 |
|           | 0.31 | 2 | 0.13 | 0.14 |
|           | 0.30 | 2 | 0.09 | 0.11 |
| kerosene | 0.52 | 2 | 0.15 | 0.35 |
|           | 0.50 | 2 | 0.11 | 0.25 |
|           | 0.49 | 2 | 0.12 | 0.27 |
|           | 0.48 | 2 | 0.08 | 0.15 |
|           | 0.46 | 2 | 0.07 | 0.14 |
|           | 0.45 | 2 | 0.02 | 0.01 |
| toluene | 0.08 | 2 | 0.37 | 0.34 |
|           | 0.06 | 2 | 0.27 | 0.27 |
|           | 0.05 | 2 | 0.16 | 0.17 |
|           | 0.04 | 2 | 0.02 | 0.03 |

Table 6. Mean Growth Rates of (110) and (1\textsuperscript{−}10) Faces of Methyl Stearate Crystals Growing in n-Dodecane, Kerosene, and Toluene

Figure 7. C18:0 growing from n-dodecane. For each set of four plots, above: G(σ) experimental data fitted to the power law and BCF models; below: trend of the total resistance to mass transfer as a function of ΔC using the parameters obtained from the data fitting to these models. Left refers to the (110) and right to the (1\textsuperscript{−}10) faces, respectively.
Figure 6 compares the most likely morphology prediction, using either the orthorhombic/monoclinic A\(2/a\) or monoclinic C\(_2\) unit cell parameters, with a micrograph taken from a selected experimental crystal. The selected micrographs feature both the dominant morphology observed in all solvents and that observed at the lowest supersaturations in kerosene systems. In the case of the final chosen morphology using the monoclinic crystal system with space group A\(2/a\), the occasional face (011) is predicted for the habit observed at lower supersaturation in kerosene. This is in agreement with the morphology of the B monoclinic polymorph reported for stearic acid crystals.\(^ {24−26} \)

Given the asymmetrical nature of the crystals’ morphology observed at the lowest supersaturations in kerosene, the morphology predicted by the monoclinic crystal system C\(_2\) seems to deliver the best match to those crystals observed experimentally. Thus, the final chosen indexation is (110) (1−10) (−1−10) (−240) (−3−10).

### 3.2. Mean Growth Rates and Growth Rate Mechanism of the (110) and (1−10) Faces of Methyl Stearate Crystals As a Function of Growth Environment.

A sequence of images of methyl stearate crystals in a 0.5 mL cuvette crystallization cell in three different solvents is shown in section 4 of the SI. The mean growth rates of the (110) and (1−10) faces of single crystals of methyl stearate growing in \(n\)-dodecane, kerosene, and toluene are presented in Table 6.

Up to now no information has been reported in the literature on the growth and morphology of methyl ester crystals. However, the growth rates of stearic acid crystals growing from different solvent systems have been studied.\(^ {24−26} \) The values reported for the (110) face of these crystals are within the same order of magnitude as those obtained in this work (Table 6).

In general growth occurs faster for crystals growing from toluene. In this case growth rates are within the same order of magnitude as those observed in the other two solvents but take place at supersaturations which are a much lower order of magnitude, e.g., 0.04 < \(\sigma\) < 0.08 for toluene compared to 0.30 < \(\sigma\) < 0.39 for \(n\)-dodecane.

Using the indices predicted for the monoclinic C\(_2\) crystal system the crystal faces were identified for the asymmetrical morphology produced for kerosene at lower supersaturation. These revealed the presence of the (−3−11) face, which grows at a rate of 0.16 \(\mu\)m/s.

To assess the mechanistic regime of crystal growth the data collected for the (110) and (1−10) faces were fitted to the models described by eq 2 to 4. However, given that experimental \(G(\sigma)\) observations showed there is a critical supersaturation (\(\sigma_{\text{crit}}\)) below which growth does not proceed, this parameter was introduced within the models to account for its effect by subtracting it from the term \(\sigma\).

Figures 7, 8, and 9 show the best fits of these models to the experimental data for both the (110) and (1−10) faces for \(n\)-dodecane, kerosene, and toluene solvents, respectively. In addition to this, using the bulk and interface transfer coefficients obtained from the experimental data fitting, they also present the trend of the total resistance to mass transfer of growth units as a function of driving force. The driving force is defined as the difference between the actual solution’s concentration and the
equilibrium concentration (solubility) \((\Delta C = C - C_e)\), \(\frac{1}{k_{MT}}\) is defined by the denominator of the \(G(\sigma)\) expressions given by the corresponding mechanistic model assessed.

All relevant parameters obtained through this analysis are presented in Table 7. For comparative assessment, when more than one model fitted well to the experimental data all fitting lines were drawn and the corresponding modeled parameters were also given. For illustration, although a more accurate assessment of the rate limiting step can be done through Figures 7–9, an estimation of both the resistance to transfer within the bulk and at the interface are given in Table 7 using average values of \(\sigma\) and \(C_e\) within the range of study.

Although the amount of experimental data collected is probably not enough to accurately determine the values of \(k_{MT}\) and \(k_G\), the fitting of the models presented to these data can nonetheless deliver relevant mechanistic information.

While there is no direct visual evidence of the presence of screw dislocations for the (110) and (1\(-10\)) faces of C18:0 crystals, the best fittings to the experimental data for C18:0 growing from \(n\)-dodecane solvent were obtained for the models given by both the power law and the BCF mechanism. Given that in the power law a value of \(r\) equal to 2 is associated with the BCF mechanism, this strongly suggests that growth in this case proceeds via screw dislocations. Since a linear dependence of \(K_1\) on \(\Delta C\) is relevant to growth limited by bulk diffusion, Figure 7 indicates that growth of C18:0 from \(n\)-dodecane is hindered by transfer of growth units at the crystal/solution interface. In addition, the shape of the \(\frac{1}{k_{MT}}(\Delta C)\) lines is coincident with that of the hyperbolic form given by the BCF mechanism but in opposite direction showing that as expected higher interfacial resistance would be placed at lower levels of driving force. The resistance to incorporation at the crystal/solution interface \(\left(\frac{1}{k_{s}}\right)\) (calculated using average values of \(\sigma\) within the range of analysis) is significantly higher than the resistance to mass transfer within the bulk \(\left(\frac{1}{k_{MT}}\right)\) (Table 7), this supporting surface integration as the rate limiting step.

The best fittings to the experimental data for C18:0 growing from kerosene solvent were obtained for the models given by both the power law and the B&S mechanism. However, since the fitting to the power law delivered \(r\) values of 0.51 and 0.68, and a value of \(r\) equal to 0.83 could be associated with the B&S mechanism, the parameters obtained via the fitting to this model were taken for this analysis. Figure 8 shows that if this is the case the resistance to mass transfer within the bulk would play a more important role on the face growth and would be rate limiting. This representing between 44.8% and 100% of the total resistance to transfer of growth units for the (110) face and 100% of the resistance for the whole range of \(\sigma\) studied for the (1\(-10\)) face. The differences in these proportions can be explained by the fact that the resistance to incorporation of growth units at the interface \(\left(\frac{1}{k_{s}}\right)\) (calculated using average values of \(\sigma\) within the range of analysis) for the
(110) face is 9 orders of magnitude higher than that of the (1−10) face (Table 7). Considering that $k_2$ values are of the same order of magnitude, this significant difference is due to the value obtained for the parameter $A_1$ which is inversely related to the 2D nuclei interfacial tension. Therefore, the more elongated habit of the crystals observed in this case is likely to be the result of differences between the molecular structures of the (110) and (1−10) faces in which the former would provide a less suitable surface for the formation of 2D nuclei, imposing a higher proportion of resistance to the face growth at the interface.

A fitting to the experimental data for C18:0 growing from toluene solvent was obtained only for the model given by the power law, and similarly to the case of growth in kerosene the resistance to mass transfer within the bulk is rate limiting. This effect is shown in Figure 9 where the proportion of $1/k_{MT}$ to the total resistance to transfer $1/k_{MTOT}$ for both faces is above 50% all over the range of $(\sigma)$ studied. Interestingly, this analysis delivered values of $r$ equal to 0.4 and 0 for the (110) and (1−10) faces respectively, which cannot be associated with any of the interface kinetic models given by expression (1) to (4) of the SI, indicating that growth from toluene would be entirely controlled by the diffusion of growth units toward the crystal.

A comparison between the values obtained for the resistance to mass transfer within the bulk, using the best model fitting parameters, shows an interesting trend in which this resistance is 1 order of magnitude lower in toluene than in kerosene and 4 orders of magnitude lower in n-dodecane than in toluene. This result is likely to be due to the aromatic nature of toluene and the presence of both aromatic and cyclo-alkyl compounds in kerosene that would provide stronger solute–solvent interactions due to the more polar nature of solvent molecules. This condition would make difficult the desolvation process and decrease the rate of transfer of the solute molecules.

The values obtained for the resistance to transfer at the interface are much more similar in magnitude, being only significantly lower for the (1−10) face of crystals growing from kerosene solvent, where their values are 8 and 9 orders of magnitude lower than those observed in the other solutions, likely due to very low resistance to the formation of 2D nuclei. This resistance is 1 order of magnitude higher in n-dodecane than in toluene solvents which could only be the case if the faces of crystals growing from toluene provide sufficient and larger number of kink sites.
3.3. Combined Assessment of Methyl Stearate Crystallization as a Function of Solvent. Table 8 presents an overview of results together with those obtained from the assessment of nucleation in the same solutions’ systems.29 If a comprehensive assessment is carried out drawing on the findings within each section of this paper together with those presented in ref 29, it can be concluded that the crystallization process in toluene systems will deliver a lower number of crystals, with larger sizes in comparison to those crystals expected to be obtained from n-dodecane and kerosene solvents. This is consistent with lower nucleation rates and significantly higher crystal growth rates in this solution system. These findings are supported by observation of the crystallization process under microscope in which fewer larger faceted crystals were observed growing at such significant rate that their growth could only be recorded at very low supersaturation σ below 8%. In this case the highest critical undercooling relates to the highest solubility in toluene and presumably to the highest solution density, hence to the lowest solute diffusivity and mass transfer. Similarly, it suggested that integration of growth units to the crystal surface would proceed via a BCF and a B&S mechanism for growth in n-dodecane and kerosene solvents respectively. Integration of growth units for growth in toluene would be facilitated by the presence of sufficient kinks sites.

Table 8. Summary of Parameters Obtained through the Combined Assessment of Solubility, Morphology and Nucleation and Crystal Growth Kinetics of Methyl Stearate in Three Different Solvents

| Solubility | n-Dodecane | Kerosene | Toluene |
|------------|------------|----------|---------|
| Y (20°C)   | 2.01       | 1.85     | 1.14    |
| ΔH_{diss}  | 63.55      | 60.81    | 59.84   |
| ΔS_{diss}  | 0.20       | 0.22     | 0.19    |
| ΔH_{mix}   | 9.61       | 15.87    | 5.90    |
| ΔS_{mix}   | 0.03       | 0.05     | 0.02    |

Conclusion

All systems behave less than ideal as activities are higher than one. However, higher values in n-dodecane followed by kerosene solutions indicate either solute-solute or solvent-solvent interactions are favoured in these systems. Solute-solute interactions are similar to those of solute-solute in toluene solutions as activities are 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 solutions evidence strong dependence of activities on temperature.

Nucleation

| Range σ  | 0.32-0.44 | 0.50-0.66 | 0.49-0.58 |
| Range γ  | 0.94-1.55 | 1.21-1.52 | 1.18-1.88 |
| Range r* | 0.84-1.00 | 0.74-0.94 | 0.73-1.17 |
| Range J  | 6.70 x 10^{11} - 1.27 x 10^{12} | (1.24 - 1.79) x 10^{17} | 4.56 x 10^{16} - 1.04 x 10^{17} |

Conclusion

Relatively higher values of interfacial tensions in toluene seem to hinder nucleation in this solvent. Although interfacial tension values are close in n-dodecane and kerosene solutions, nucleation in the former is one order of magnitude lower. This could indicate less available nucleation sites in these solutions, due to the low solubility of methyl stearate in n-dodecane.

Morphology

The morphology observed in all cases was the same except for the lowest supersaturation levels in kerosene in which there was a less symmetrical habit. The indexation of the latter allowed for the identification of the crystal system that would predict the most likely morphology - the monoclinic C2 crystal system.

Crystal growth

| Range σ  | 0.30-0.39 | 0.45-0.52 | 0.04-0.08 |
| ΔS = C - C_{av} | 174-215 | 233-259 | 34-61 |
| Average C_{av} | 632.86 | 581.33 | 1053.16 |

G (μm)

| (110) | (1-10) | (110) | (1-10) | (110) | (1-10) |
| 0.09-1.13 | 0.11-1.05 | 0.02-0.15 | 0.01-0.35 | 0.02-0.37 | 0.03-0.34 |

Growth mechanism

| BCF | BCF | B&S | B&S | Diffusion controlled | Diffusion controlled |
| 9.00 x 10^5 | 9.00 x 10^5 | 2.47 x 10^{11} | 2.43 x 10^{11} | 6.02 x 10^{10} | 6.02 x 10^{10} |
| 1.95 x 10^{11} | 2.08 x 10^{11} | 1.53 x 10^{11} | 1.33 x 10^{12} | 3.05 x 10^{10} | 2.74 x 10^{10} |

Conclusions

Growth is favoured in toluene solvent where similar range of growth rates were obtained for significantly lower levels of supersaturation. While growth in this case can occur at σ levels above 4%, σ levels above 27 and 44% are required for growth to proceed in n-dodecane and kerosene solvent respectively. The analysis of G(σ) data revealed that growth is limited by transfer of growth units at the interface in n-dodecane solvent and within the bulk of kerosene and toluene solvents. Similarly, it suggested that integration of growth units to the crystal surface would proceed via a BCF and a B&S mechanism for growth in n-dodecane and kerosene solvents respectively. Integration of growth units for growth in toluene would be facilitated by the presence of sufficient kinks sites.

“Y 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.”
same order of magnitude than those observed in n-dodecane solvent then it could be expected that crystallization process in the former solvent will produce a larger population of crystals of smaller sizes.

We know from crystal growth theory\textsuperscript{10,27} that impurities (solvent) can disrupt the nucleation and growth processes, through their impact on molecular recognition and self-assembly associated with the growth process. Presumably such effects can only be remedied by increasing supersaturation, through which the critical cluster size becomes less than the separation between impurity molecules on the crystal surfaces, thus enabling the initiation of nucleation/growth. With toluene solute/solvent binding is much stronger when compared to the other solvents and hence higher supersaturation might be needed.

Conversely for crystal growth in toluene solutions, the process is rate-limited by mass transfer effect rather than by interface kinetics. This contrasts with growth from n-dodecane and kerosene solvents where surface integration is the rate limiting process. In the latter case the molecular similarity between alkyl solvent and methyl stearate may effect slower desolvation, compared to the aromatic solvent toluene at the growth interface, where the surface integration takes place in a much more confined environment when compared to that occurring within the bulk solution. In the latter case the solvent would be much more likely to be retained temporally at the interface thus acting as an “impurity”, which would impede growth until the supersaturation is high enough and surface nuclei radius small enough to allow growth to proceed.

4. CONCLUSIONS

A detailed analysis of the crystal morphology for methyl stearate, using a methodology based on a combined BFDH and zone axis analysis\textsuperscript{12} yielded the morphological indexation to be (110) and (1–10) based on the monoclinic C2 crystal structure. Crystals growing from supersaturated n-dodecane, kerosene, and toluene solutions, as study using in situ optical microscope, at supersaturation (\(\sigma\)) levels of 0.30–0.39, 0.45–0.52, and 0.04–0.08, respectively, showed that the crystal morphology changes only in the case of methyl stearate crystallizing from kerosene at the lowest supersaturation values (\(\sigma < 0.45\)). The growth rates measured in toluene are of the same order of magnitude to those observed in kerosene, ranging from 0.02 to 0.37 \(\mu m/s\) and are around half those observed in n-dodecane solutions, in which these values range from 0.09 to 1.13 \(\mu m/s\). Measurements of the growth for the (110) face in the case of kerosene solutions are much lower in magnitude compared to those for the (1–10) face, ranging from 0.02 to 0.15 \(\mu m/s\), which gives these crystals an elongated lozenge shape. Similar growth rates values are observed between the (110) face and the (1–10) face for crystals growing from n-dodecane and toluene solvents. The tendency of the growth rate dependence on \(\sigma\) was consistent with the BCF growth mechanism in n-dodecane, the B&S mechanism in kerosene, and diffusion controlled in toluene solvent.

### ASSOCIATED CONTENT

1 Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.cgd.6b01436.

Description of the methodology used to solve crystal morphology; the complete set of planes delivered through the prediction of the BFDH morphology for methyl stearate crystals using three different sets of unit cell parameters; the derivation of models expressions for the assessment of the dependence of single faces growth rates (\(G\)) on supersaturation (\(\sigma\)) and a sequence of images of methyl stearate crystals growing with time in three different representative diesel type solvents (PDF)

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**Notes**
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### SYMBOLS

\(A_s\), Growth parameter in B&S model
\(A_p\), Growth parameter in the BCF model
\(C_s\), Solution concentration (\(m^{-3}\))
\(C_p\), Equilibrium concentration (\(m^{-3}\))
\(d_{hkl}\), Interplanar distances within morphological (\(hkl\)) forms
\(f\), Nucleation rate (nuclei mL\(^{-1}\) s\(^{-1}\))
\(k_b\), Boltzmann constant (\(J K^{-1}\))
\(k_d\), Docking rate (mol m\(^{-2}\) s\(^{-1}\))
\(k_{G}\), Growth rate constant (m\(^{1/(m/s)}\) s\(^{-1}\))
\(k_{MT}\), Mass transfer coefficient (m s\(^{-1}\))
\(r\), Growth exponent
\(r^*\), Critical nucleus radius
\(R\), Single face growth rate (ms\(^{-1}\))
\(S\), Supersaturation ratio
\(T\), Solution temperature (K)
\(T_{eq}\), Solution saturation (or equilibrium) temperature (K)
\(V\), Volume of solution (m\(^3\))
\(x_s\), Mole fraction of solute in solution
\(x_m\), Equilibrium mole fraction
\(\sigma\), Relative supersaturation
\(\sigma_{crit}\), Critical relative supersaturation
\(\rho_s\), Solute density (kg m\(^{-3}\))
\(\Delta H_{diss}\), Enthalpy of dissolution (kJ mol\(^{-1}\))
\(\Delta H_{mix}\), Enthalpy of mixing (kJ mol\(^{-1}\))
\(\Delta S_{diss}\), Entropy of dissolution (kJ mol\(^{-1}\) K\(^{-1}\))
\(\Delta S_{mix}\), Entropy of mixing (kJ mol\(^{-1}\) K\(^{-1}\))
\(\gamma\), Interfacial tension (m J\(^{-2}\))
\(Y\), Activity coefficient
ABBREVIATIONS

BCF, Burton-Cabrera-Frank model
BFDH, Bravais-Friedel-Donnay-Harker model
B&S, Birth and Spread model
MSZW, Metastable zone width
MW, Molecular weight
RIG, Rough Interface Growth model
SD, Standard deviation
2D, Two dimensional
3D, Three dimensional

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