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crystallisability and nucleation behaviour of

eicosane in toluene: acetone mixed-solvents†

Influence of solution chemistry on the solubility,

The interplay between the solution chemistry and crystallisability of eicosane in mixed toluene: acetone solutions is examined over the full compositional range from pure toluene to pure acetone, using a combination of polythermal crystallisation experiments and molecular modelling. Enthalpies of dissolution and mixing, as well as metastable zone widths increase with increasing acetone content, up to a mol fraction of 0.85 acetone, followed by a decrease in values to pure acetone solutions. Nucleation is found to occur via an instantaneous pathway for the pure solvent systems and also when toluene is in excess, in contrast to solutions where acetone is in excess, which are found to nucleate progressively. Rationalisation through molecular modelling highlights likely changes in the solution structure, whereby eicosane can be expected to be preferentially solvated by toluene, with this solvated cluster being surrounded by a 'cage' of acetone molecules. This proposed structure is consistent with a model whereby solute diffusion and hence clustering is hindered when acetone is in excess, decreasing the crystallisability of the solution and effecting a change in the mechanism of nucleation. However, above a critical acetone composition, the potential for complete toluene solvation is restricted and easier crystallisation is enabled.

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## 1. Introduction

Solution phase crystallisation is an energy-efficient physical process widely utilised to isolate and/or purify solid-form ingredients. The majority of speciality chemicals are crystallised in multi-substance batch reactors, which provide limited opportunities for specialised reactor designs and for optimising crystal product properties. However, selection of a mixed-solvent crystallisation system, comprising two or more miscible components, provides an alternative opportunity for the control and optimisation of the crystallisation process.

directed, either through cooling of an initial solvent blend<sup>7,16</sup> or through antisolvent crystallisation. 15,17 Both routes require miscible solvents, with the latter requiring significant differences in the solubility of the solute between the solvent and antisolvent, the addition of which lowers solute solubility and hence generates supersaturated conditions. In the design of mixed-solvent crystallisation processes, it is helpful to have developed workflows for the characterisation of the solubility and solution properties over the full range of compositions used and, through this, optimise the process parameters to ensure the desired crystal product characteristics. 18,19

Mixed-solvent crystallisation systems have been intensively studied, due to their compositional variation affecting solubility,2 solid-form produced,3-5 crystal growth,6 habit,7

For the design of antisolvent crystallisation processes knowledge of the solubility-temperature relationship as a function of the composition of solvent mixture is important for control of the solution supersaturation generated by the addition of the lower solubility solvent. Depending upon the magnitude and sign of the enthalpy of mixing, the addition of a second solvent can change the solution temperature 15,17 and hence a detailed knowledge of a solution's mixed-solvent thermodynamics is also needed.

crystal size,8 crystallisability9-12 and nucleation.13-15 Crystallisation from mixed-solvent solution systems can be

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<sup>†</sup> Electronic supplementary information (ESI) available: It includes: the average crystallisation and dissolution temperatures determined for all solutions studied across the full mixed-solvent compositional range, atomic partial charges of eicosane, toluene and acetone used in grid-based molecular modelling studies and the full analysis of solvation sites of eicosane solvated by single toluene and acetone molecules. See DOI: 10.1039/d1ce00322d

Paper CrystEngComm

Changes in solution properties can, in turn, also impact upon solute mass transport within the solution phase and through this, impact upon the overall crystallisability of the material. Hence, characterisation of the metastable zone width (MSZW) as a function of solvent composition is required in order to define the operational conditions for large-scale crystallisation processes.

Variation in mixed-solvent composition can also change the nature of the intermolecular interactions associated with solute desolvation and clustering, which can directly affect the ability of the solute to nucleate, impacting, in turn, on the nucleation kinetics and associated mechanism. 15,17,20 In the latter respect, polynuclear nucleation can occur either through instantaneous (IN) or progressive (PN) nucleation mechanisms,<sup>21</sup> whereby IN describes the process in which all nuclei form instantly and subsequently undergo crystal growth, whereas in PN the nuclei continuously form in the presence of growing crystals. Such mechanistic changes can impact on the crystal product properties, e.g. IN could, in principle, result in a more uniform crystal size distribution of product. Previous studies in other organic solvents have highlighted the fact that changes in solution environment, relating to solution solvent and concentration20,22 as well as the inclusion of nucleation inhibition additives, 23 can alter the mechanism of nucleation. The IN mechanism has been shown to occur when solute clustering is present in solution prior to the nucleation stage,<sup>24</sup> in line with van't Hoff isotherms suggesting strong solute: solute interactions within solution.<sup>20</sup> The relationship between solute clustering, solution structure and nucleation mechanism need to be investigated to provide insight into these cooperative effects.

In this study, the crystallisation of the long-chain hydrocarbon, eicosane (C<sub>20</sub>H<sub>42</sub>), from mixed toluene: acetone solutions is examined. Alkanes, in general, are known to be highly soluble in toluene<sup>25</sup> but can be expected to be much less soluble in more polar solvents, such as acetone. The work presented here investigates this mixed-solvent system, examining the interplay between solvent composition and solution chemistry, in terms of the solutions solubility and crystallisability, thermodynamics and nucleation kinetics and mechanism. In this, the approach adopted encompasses experimental polythermal crystallisation/dissolution measurements<sup>26-28</sup> to characterise the solution solubility and nucleation kinetics and mechanisms coupled with molecularscale grid-based modelling<sup>29</sup> of solute:solvent interactions. The latter seeks to compliment the experimental studies providing an insight into the likely molecular-scale solvation process for the mixed-solvent system and through this,

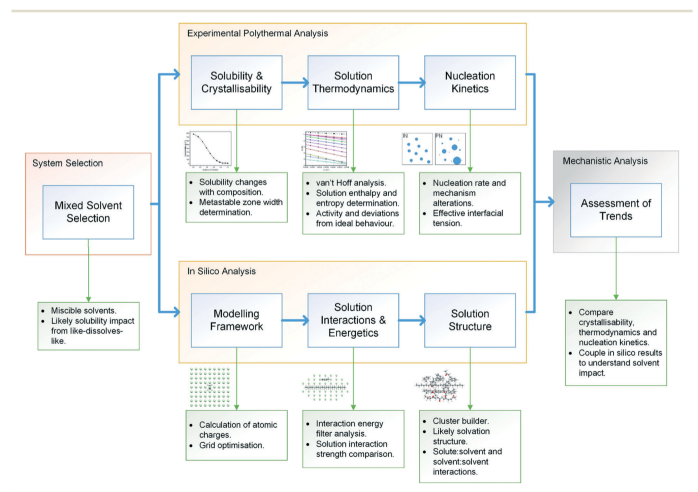


Fig. 1 Methodological workflow incorporating crystallisation analysis with molecular modelling for enhancing process understanding

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more complete understanding intermolecular interactions taking place within the solution phase, together with their concomitant impact upon the crystallisation process.

## 2. Materials & methods

#### 2.1. Workflow

The methodological workflow used in this study is given in Fig. 1, providing a standardised assessment for the characterisation of mixed-solvent solutions and crystallisation, through an integrated approach, coupling experimental and modelling techniques.

#### 2.2. Materials

Eicosane (99% pure) was supplied by Sigma-Aldrich, toluene (≥99.3% pure) was supplied by Honeywell and acetone (99.5+% pure) was supplied by Arcos Organics. All materials were used as received without further purification.

#### 2.3. Experimental procedure

2.3.1. DSC measurements and data analysis. The latent heat of crystallisation and melting temperature of eicosane were measured using a Mettler Toledo differential scanning calorimetry DSC 1 STARe system. 3 mg of eicosane was weighed into 40 µl aluminium standard pans and was subject to a temperature profile initiated by heating to 40 °C, where it was held for an hour to ensure complete homogenization, and then cooled to -15 °C, where it was also held for an hour to allow equilibration. A constant rate (q) of 0.25 °C min<sup>-1</sup> was used in both the heating and cooling segments. The temperature cycle was repeated five times to obtain mean values of the latent heat of crystallisation and melting temperature, which were determined from the integration of the corresponding exothermic and endothermic peaks using Origin software.30

2.3.2. Solution preparation and polythermal measurements. All solutions were prepared at a minimum of four solution concentrations, ranging from 5-850 g L<sup>-1</sup> over the full compositional range of toluene: acetone mixtures, e.g. 250-850 g  $L^{-1}$  for eicosane in pure toluene and 5-35 g  $L^{-1}$ for eicosane in pure acetone solutions, with concentrations varying with solvent composition to take into account changes in solubility. A total of eleven compositions of the toluene: acetone solvent mixtures in 10% volume increments were used.

Solutions were prepared by weighing eicosane into 15 ml prewashed glass vials, using a weighing scale that could measure with ±0.1 mg accuracy. A Fisherbrand 500-5000 μl micropipette was used to add the respective solvent to each concentration sample, with 5 ml total solvent volume used. A magnetic stirrer was placed into the glass vials and the mixtures were placed on a stirrer hot plate and held at 35 °C for 30 minutes under constant agitation, in order to form a homogeneous liquid solution. 1 ml of solution was added to the 1.5 ml glass vials and a standard 7 × 2 mm magnetic stirrer was added to each vial.

Crystallisation experiments were performed using the Technobis Crystal 16® system31 with crystallisation and dissolution onset points being detected by optical turbidity methods. The 1 ml solutions were subject to different cooling and heating cycles. Each temperature cycle began by heating the solutions to 35 °C, before being held at this temperature for one hour to ensure that complete homogenisation had taken place and was the solution was then subsequently cooled at a set rate to -15 °C. This temperature was then held for an hour to allow equilibration, followed by an increase in temperature back to 35 °C at the same specified rate. This temperature profile was performed for each concentration of each composition at rates (q) of 0.25, 1, 2 and 3.2 °C min<sup>-1</sup> for both the cooling and heating segments of the temperature cycles and at a constant stirring rate of 300 rpm. Each temperature cycle was performed five times to obtain mean and standard deviation (STD) values for the ratedependent crystallisation and dissolution temperatures determined.

2.3.3. Solubility and crystallisability assessment. The individual crystallisation  $(T_c)$  and dissolution temperatures  $(T_{\rm diss})$  were determined from transmission (optical turbidity) vs. temperature profiles. The values for sample transmittance were calibrated from 100%, for a homogeneous liquid solution, to 0%, for a fully crystallised solution.  $T_c$  was taken as the onset point where the light transmittance dropped to approximately 95% and  $T_{\rm diss}$  was taken as the point where the light transmittance reached 100% for all measurements. The polythermal data was used to extrapolate both  $T_c(q)$  and  $T_{\rm diss}(q)$  lines back to a cooling rate of 0 °C min<sup>-1</sup> so that the crystallisation temperatures at the kinetic limit  $(T_{c,l})$  and equilibrium saturation temperatures  $(T_e)$  could be determined, respectively. The difference between  $T_{\rm c,l}$  and  $T_{\rm e}$ was taken as a measure of the steady-state metastable zone width (MSZW).

A van't Hoff analysis was performed on the solubility data determined from the saturation temperatures. The solution ideality was determined with respect to the ideal solubility as obtained from DSC analysis, based upon the Hildebrand equation (eqn (1)). Activity coefficients ( $\Upsilon$ ), enthalpy ( $\Delta H_{\rm diss}$ ) and entropy ( $\Delta S_{\rm diss}$ ) of dissolution were calculated based on eqn (2) and (3), respectively:<sup>25</sup>

$$\ln(x_{\rm ideal}) = \frac{\Delta H_{\rm fus}}{R} \left[ \frac{1}{T} - \frac{1}{T_{\rm m}} \right] \tag{1}$$

$$\Upsilon = \frac{x_{\text{ideal}}}{x} \tag{2}$$

$$ln(x) = -\frac{\Delta H_{diss}}{RT} + \frac{\Delta S_{diss}}{R}$$
 (3)

where  $x_{\text{ideal}}$  is the solution concentration, x is experimentally measured molar solubility,  $\Delta H_{\text{fus}}$  is

enthalpy of fusion, R is the ideal gas constant, T is the temperature and  $T_{\rm m}$  is the melting temperature.

The calculated activity coefficients were fitted with respect to their variation in temperature, by:

$$\ln \Upsilon = aT + c \tag{4}$$

where a and c are constants.

The enthalpy  $(\Delta H_{\text{mix}})$  and entropy  $(\Delta S_{\text{mix}})$  of mixing were calculated from the deviation in  $\Delta H_{\rm diss}$  and  $\Delta S_{\rm diss}$  from ideality, through the relationships:

$$\Delta H_{\rm mix} = \Delta H_{\rm diss} - \Delta H_{\rm diss}^{\rm ideal} \tag{5}$$

$$\Delta S_{\text{mix}} = \Delta S_{\text{diss}} - \Delta S_{\text{diss}}^{\text{ideal}}$$
 (6)

2.3.4. Nucleation kinetic analysis. The Kashchiev-Borissova-Hammond-Roberts (KBHR) approach, 26-28 utilised in this study, enables the distinction between instantaneous (IN) and progressive (PN) nucleation in a crystallising system, with the latter referring to continuous birth of new crystallites amongst already growing ones and the former being characterised by all crystallites appearing at once in the solution, followed by only growth, without subsequent nucleation events. In the PN case, the number of crystallites increases as a function of time, with a large distribution of crystallite sizes, whereas in the IN case there are a fixed number of crystallites of equal sizes, assuming a consistent growth rate. IN occurs when a number of strongly active nucleation sites are present in solution, which can occur due to foreign particles within solution, resulting in heterogeneous nucleation, or when solutions are seeded with crystallites, either externally or self-seeded, or when, as previous studies have highlighted, 23,24 solutions are subject to strong solute clustering, resulting in a number of prenucleation clusters that nucleate at once.

Regardless of the mechanism by which nucleation occurs, the solution remains apparently unaffected until a critical level of undercooling is reached corresponding to the limit of the MSZW, at which point the amount of formed crystallites detectable by the monitoring technique. undercooling is dependent upon the nucleation, growth and cooling rates. The KBHR approach utilises the relationship between the relative critical solution undercooling  $(u_c)$  and the cooling rate (q) to enable the determination of crystallisation parameters.  $u_c$  was determined by:

$$u_{\rm c} = \frac{\Delta T_{\rm c}}{T_{\rm e}} \tag{7}$$

where,  $\Delta T_{\rm c} = T_{\rm e} - T_{\rm c}$ .

The mechanism of nucleation was determined through the "rule of three", from ln-ln plots of  $u_c$  vs. q, the slopes of which were fitted by linear regressions, whereby a slope >3 signifies a PN pathway and a slope <3 indicates an IN pathway. 26,27

For the PN case, the final expression for  $u_c(q)$  dependence, when expressed through the number of crystallites at the detection point  $(N_{det})$ , can be defined as:

$$\ln q = \ln q_0 + a_1 \ln u_c - \frac{a_2}{(1 - u_c)u_c^2}$$
 (8)

Plots of  $u_c$  vs.  $\ln q$  fitted with eqn (8) enable the free parameters to be determined for the calculation of the nucleation parameters and the effective interfacial tension  $(\gamma_{\rm eff})$ , eqn (9). This enables the calculation of the critical nucleus radius  $(r^*)$ , assuming spherical nuclei, and the number of molecules in the critical nucleus  $(i^*)$ , through eqn (10) and (11).

$$b = \frac{k_{\rm n} \nu_0^2 \gamma_{\rm eff}^3}{k T_{\rm e} \lambda^2} \tag{9}$$

$$r^* = \frac{2\gamma_{\rm eff}\nu_0}{\lambda u} \tag{10}$$

$$i^* = \frac{2bkT_e}{\lambda u^3} \tag{11}$$

where  $k_{\rm n}$  is the nuclei numerical shape factor,  $\nu_0$  is the volume occupied by a solute molecule in the crystal,  $\lambda$  is the molecular latent heat of crystallisation and k is the Boltzmann constant and the dimensionless thermodynamic parameter b is related to the nucleation rate, by:

$$J(t) = K_{\rm J} e^{-\frac{b}{[(1-u_{\rm c})u_{\rm c}^2]}} \tag{12}$$

where I is the nucleation rate and  $K_{I}$  is the nucleation rate constant, which is related to the Zeldovich factor (Z), the attachment frequency of monomers to the nucleus  $(f^*)$  and the concentration of nucleation sites  $(C_0)$ .

More detailed descriptions of the KBHR approach have been given previously.26-28

#### 2.4. Molecular modelling

2.4.1. Molecular structure building. The molecular structures of eicosane, toluene and acetone, provided in Fig. 2, were built within Biovia Materials Studio, 32 with structural optimisation and atomic fractional charges derived from a calculation of the electrostatic potential using Gaussian09,33 with a grid-based method (ChelpG) scheme,34 a 6-31G\* basis set and the Becke three-parameter<sup>35</sup> Leefunction.36 Yang-Parr exchange-correlation eicosane is likely to exist in a number of conformations within solution, due to its long-chain nature, structural optimisation was performed to generate the lowest energy eicosane molecular structure, representative of a likely structure within the solution environment, with accurate calculated atomic fractional charges.

2.4.2. Molecule-molecule solvation grid-search. Solute: solute and solute: solvent interactions were examined using intermolecular grid search methods, 37,38 with a spatial grid defined around a central target molecule, Fig. 3. A grid

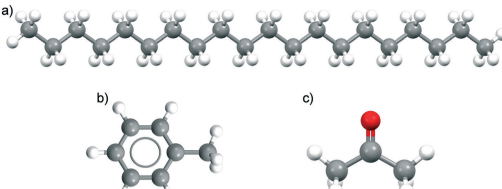


Fig. 2 Chemical structures of a) eicosane, b) toluene and c) acetone.

optimisation was performed and an orthogonal grid shape with size 15, 15 and 35 Å in the X, Y and Z planes, respectively was found to be optimal for energy calculations. In each plane, the number of grid points was defined with 8, 8, 10 steps of grid points on the X, Y and Z planes, respectively. This provided a grid with a total number of 891 points, Fig. 3. During the intermolecular interaction energy calculation procedure, a probe molecule was placed on each point of the grid and rotated through a set of Euler angles, which were set at  $4^{\circ}$  for the x-axis, y-axis and z-axis. This resulted in a heterogeneous six-dimensional search space, and at each point within this the interaction energy between the probe and target molecules was calculated using an atom-atom approach, with parameters from the Momany intermolecular potential. 16,39 Intermolecular interactions in the form of van der Waals, hydrogen bonding and electrostatic interactions were calculated, with the summation of the repulsive and attractive components of each of these interaction types providing the total intermolecular interaction energy.

The solvating power of toluene and acetone to eicosane was assessed by comparison of the intermolecular interaction strengths of the solvents (probes) with eicosane (target). A low-pass energy filter was be applied to remove interactions below given interaction strengths, which were set at -0.5, -1.0, -1.5 and 2.0 kcal mol<sup>-1</sup> for toluene interactions and -0.5, -1.0, -1.2 and 1.4 kcal mol<sup>-1</sup> for acetone interactions. The more negative energy filters highlighted the most favourable interaction sites with an eicosane molecule. A total of 331 986 600 points (location + rotation) were searched for each simulation.

All possible solvent and solute intermolecular interactions within the solution phase (eicosane: eicosane, eicosane: toluene, eicosane: acetone, toluene: toluene, acetone: acetone toluene: acetone) were calculated intermolecular grid search method. A low-pass energy filter of -0.5 kcal mol<sup>-1</sup> was used to screen out the lowest energy interactions with the screen resulting in the calculation of over 4000 possible intermolecular interaction energies for each interaction type within the solution phase.

2.4.3. Solvation shell grid-search. Solvation energies and solution structure were assessed via the grid-search method, using cluster building. In this, a probe molecule was located at the lowest-energy site in the grid around a target, with this molecular-pair becoming a new fixed-target for a subsequent probe molecule, which was then placed on the next available lowest-energy site. This process was repeated until the positions for a shell of molecules had been located. The intermolecular interaction energy between each subsequent probe and target as well as probe-probe interactions were calculated. 20 solvent molecules were used as probes around the target eicosane molecule, in different toluene: acetone mixed solvent ratios of 0:20, 5:15, 10:10, 15:5 and 20:0 molecules. Software procedures required one set of solvent: solute interactions to be calculated prior to the other. A previous study demonstrated that, in solubility mismatched mixed-solvent solutions, such as the system studied here, the local solvation environment of a

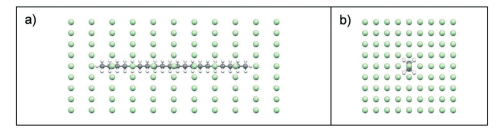


Fig. 3 Spatial grid defined around target eicosane molecule for calculation of intermolecular interaction energy calculations; (a) view along b axis; (b) view along c axis. Grid points shown by green dot symbols.

nucleating solute would be likely to be dominated by the strongest solvating solvent, due to the much higher intermolecular interaction strengths.<sup>17</sup> Therefore, for the current procedure the interactions of toluene molecules with eicosane were calculated first, followed by the acetone molecules. Whilst this molecular modelling technique provides an attractive and representative molecular-scale assessment as to likely solute: solvent ordering within the solution, it is clearly limited by the fixed interaction procedure imposed by the chosen grid size that neither directly allows for the effects of molecular conformational flexibility nor deals with the random solute: solvent interactions, which might be expected within a mixed-solvent/antisolvent system, as studied here.

The overall cluster structures generated were optimised using the Forcite module within Biovia Materials Studio. The SMART algorithm and a very fine tolerance was used to distinguish intra and inter-molecular interactions, ensuring solvation energies could be determined. For this, Gasteiger<sup>40,41</sup> atom point charges were utilised. All structures were visualised using the CCDC's Mercury<sup>42</sup> software package.

# 3. Results

#### 3.1. Assessment of solubility and crystallisability

Examples of polythermal heating-cooling and turbidity profiles used for the temperature cycles are displayed in Fig. 4(a) and (b), highlighting the determination of  $T_{\rm diss}$  and  $T_{\rm c}$ . The equilibrium solubility  $(T_{\rm e})$  and supersolubility  $(T_{\rm c,l})$ temperatures obtained from extrapolation of  $T_{\rm diss}$  and  $T_{\rm c}$ back to 0 °C min<sup>-1</sup> are given in Table 1, together with their MSZW ( $\Delta T_c$ ) for eicosane crystallising from toluene: acetone mixture solutions. Examples of the linear regression plots of  $T_{\rm diss}$  and  $T_{\rm c}$  vs. q together with their standard deviation values from five repeats and the resulting solubility-supersolubility curves are given in Fig. 4(c) and (d), respectively. Experimental data from all of the experiments is provided in

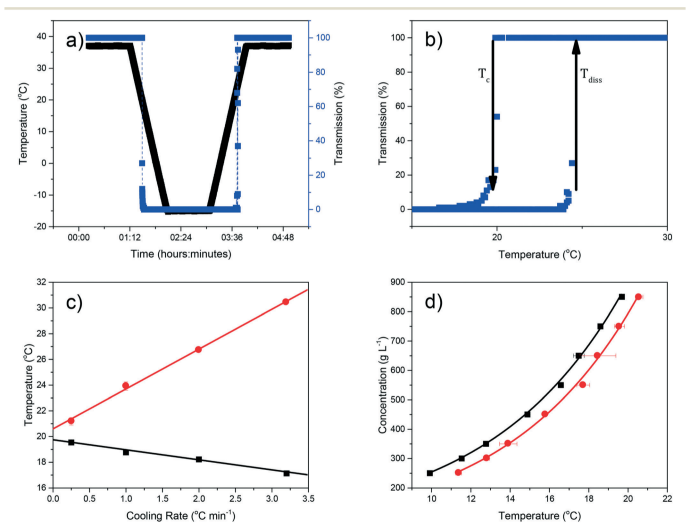


Fig. 4 Representative polythermal data: a) temperature and transmission profiles against the time as used for measuring determination of crystallisation and dissolution temperatures ( $T_c$  and  $T_{diss}$  respectively), for 850 g L<sup>-1</sup> eicosane in toluene solution at cooling rate (q) of 1 °C min<sup>-1</sup>; (b) temperature turbidity plot of a solution highlighting the observation of on-set points for  $T_c$  and  $T_{diss}$ ; (c) plots of  $T_c$  and  $T_{diss}$  as a function of q, with the corresponding best linear fits are also given, for 850 g  $L^{-1}$  eicosane in toluene solution; (d) concentration vs. temperature plot highlighting the equilibrium MSZW between the solubility and supersolubility curves, for eicosane in toluene at all concentrations studied.

Table 1 Solubility ( $T_c$ ), supersolubility ( $T_c$ ) and steady-state MSZW data of eicosane in toluene: acetone solvent mixture solutions, together with values of the slopes of the linear regressions of  $\ln u_c$  vs.  $\ln q$  plots for the range of solvent compositions studied, with the resultant nucleation mechanism determined from the "rule of three" KBHR analysis. The '\*' denotes a low correlation fitting and as such the analysis has a lower confidence value and may be potentially a less reliable result

Conc. $(g L^{-1})$	$T_{\mathrm{c,l}}$ (°C)	<i>T</i> <sub>e</sub> (°C)	MSZW (°C)	$\ln u_{\rm c}$ vs. $\ln q$ slope	Nucleation mechanism	Conc. $(g L^{-1})$	$T_{\mathrm{c,l}}$ (°C)	<i>T</i> <sub>e</sub> (°C)	MSZW (°C)	$\ln u_{\rm c}$ vs. $\ln q$ slope	Nucleation mechanism
v = 0						v = 0.14					
250	9.92	11.37	1.46	2.87	IN	200	10.39	11.85	1.46	2.80	IN
350	12.78	13.90	1.13	2.51	IN	300	13.47	14.46	0.98	2.26	IN
450	14.88	15.79	0.91	2.25	IN	400	15.47	16.48	1.01	2.37	IN
550	16.58	17.72	1.14	2.45	IN	500	16.57	17.82	1.26	3.15	PN*
y = 0.26						y = 0.38					
200	12.81	14.30	1.49	3.50	PN	200	15.05	16.51	1.47	2.87	IN
300	15.45	16.80	1.35	2.87	IN	300	17.61	18.36	0.75	2.05	IN
400	17.18	18.24	1.06	2.37	IN	400	18.87	19.74	0.88	2.41	IN
500	18.70	19.20	0.50	1.64	IN	500	20.20	20.94	0.74	2.01	IN
y = 0.49						y = 0.59					
150	15.42	17.09	1.67	2.82	IN	150	18.39	19.88	1.49	2.70	IN
250	18.69	19.76	1.07	2.32	IN	250	20.64	22.37	1.73	3.08	PN
350	20.26	21.28	1.02	2.53	IN	350	21.94	23.67	1.74	3.41	PN
450	21.31	22.67	1.36	2.67	IN	450	22.97	24.58	1.61	3.32	PN
y = 0.68						y = 0.77					
150	20.80	22.44	1.65	3.16	PN	100	21.59	23.23	1.64	3.18	PN
250	22.63	24.16	1.53	3.58	PN	200	24.29	25.84	1.55	3.27	PN
350	23.80	25.75	1.95	3.82	PN	300	25.45	26.92	1.47	3.04	PN
450	24.58	26.41	1.83	3.25	PN	400	26.02	28.22	2.20	3.91	PN
y = 0.85						y = 0.93					
75	23.17	25.00	1.82	3.61	PN	20	18.80	20.60	1.81	2.84	IN
175	26.27	28.16	1.89	4.00	PN	60	25.09	26.85	1.77	3.74	PN
275	27.29	29.34	2.05	3.84	PN	100	27.29	29.00	1.71	3.70	PN
375	27.90	29.98	2.09	4.04	PN	140	28.36	30.18	1.83	3.92	PN
y = 1											
5	11.92	13.08	1.16	1.68	IN						
15	20.27	21.53	1.26	2.28	IN						
25	23.48	24.76	1.28	4.16	PN						
35	25.81	26.57	0.76	1.95	IN						

ESI† S1. To our knowledge, no current open-literature data is available on the solubility in the ternary system studied. All the solutions measured in this study showed high repeatability from five repeats of cycling experiments, with low standard deviation values.

As expected, the solubility of eicosane was found to be highest in toluene, with increasing acetone composition causing a decrease in solubility over the full range of compositions studied, as shown in Fig. 5. This can be justified in terms of the adage 'like-dissolves-like', with eicosane being a negligibly polar molecule and as such being more likely to dissolve in a non-polar solvent. Toluene has a small dipole moment (0.36 D)<sup>43</sup> in comparison to acetone (2.90 D),44 therefore, eicosane can be expected to have greater affinity towards toluene rather than acetone molecules in solution, consistent with a higher solubility with increased toluene composition. The limited solvation power of acetone perhaps represents an antisolvent, as opposed to a true solvent for eicosane, thus demonstrating the ability of this system to be used both as an initial solvent blend for crystallisation and/or for an antisolvent crystallisation methodology. However, at lower solution temperatures, below around 15 °C, and above an acetone mol fraction (y) of 0.85, the eicosane solubility was found to increase with small additions of acetone, perhaps suggesting alterations in solution structure that could influence solute: solvent interactions and hence eicosane solubility.

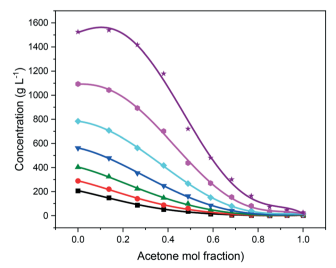


Fig. 5 Solubility of eicosane in toluene: acetone mixtures at different temperatures: 10 °C (black squares), 12.5 °C (red circles), 15 °C (green upward-triangles), 17.5 °C (blue downward-triangles), 20 °C (cyan diamonds), 22.5 °C (pink hexagons) and 25 °C (purple stars).

Previous studies have highlighted that in mixed-solvent solutions solute solubility can increase with small additions of lower solubility solvents,7,15 suggesting that the lower solubility solvent has the ability to solvate parts of the solute molecule that the higher solubility solvent cannot, hence contributing to and enhancing the overall solution solvation power of the mixed solvent system. This behaviour was observed for solution eicosane in toluene: acetone mixedsolvent solutions at 25 °C. However, below this temperature the addition of acetone was found to only reduce eicosane solubility, perhaps demonstrating acetone's comparatively poor ability to solvate the eicosane molecule when compared to toluene. The temperature dependence of this behaviour is interesting in that it demonstrates, perhaps, that a critical level of molecular mobility should be reached in order for the lower solubility solvent to access potential solute solvation sites that are not fully solvated by the higher solubility solvent within the mixed-solvent environment.

MSZW data provides an indication of the ease of which a solution overcomes its inherent nucleation barrier and crystallises, with a narrower MSZW being consistent with a tendency towards easier nucleation. For all concentrations and compositions studied, the MSZWs were found to be around 0.5-2.2 °C. These are in broad agreement with previous work, 25,45-48 but quite small when compared to other typical organic solution phase systems. 20,49 For solution compositions from 0 < y < 0.49 and 0.93 < y < 1 the MSZW was found, in general, to be roughly inversely proportional to the solution concentration. This would be consistent with nucleation being easier with a higher number of solute molecules within solution, which might be expected given that this would offer a greater chance for solute:solute interactions. In contrast, for solutions within compositional range of 0.59 < y < 0.85, the opposite trend was observed, with, in general, an increase in MSZW with increasing concentration. This could indicate changes in solution chemistry and nucleation mechanisms over the compositional ranges studied, perhaps reflecting changing nature of the intermolecular interactions within the solution state.

#### 3.2. Analysis of the solution thermodynamics

van't Hoff analysis provides further indication of the nature of the solution behaviour in terms of the impact of solvent and compositional effects. The van't Hoff plots, as a function of solvent composition, were found to fit well by linear regression for all the solutions studied, as shown in Fig. 6, consistent with no change in solute form. The enthalpy ( $\Delta H_{\rm diss}$ ) and entropy ( $\Delta S_{\rm diss}$ ) of dissolution values, enthalpy ( $\Delta H_{\rm mix}$ ) and entropy ( $\Delta S_{\rm mix}$ ) of mixing values and the activity coefficients ( $\gamma$ ) calculated from van't Hoff analysis are summarised in Table 2.

Less than ideal behaviour was observed for all the solutions with  $\Upsilon$  greater than 1 for the temperature range studied. This behaviour is consistent with a solution structure whereby homogeneous (solute:solute and solvent:

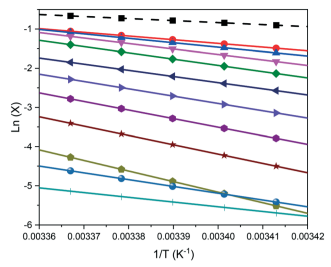


Fig. 6 van't Hoff plots for eicosane in toluene: acetone mixed solvent solutions. Dashed black line represents ideal solubility, and solid lines represent experimental solubilities for different solvent compositions: y = 0 (red circle), y = 0.14 (blue up-pointing triangle), y = 0.26 (pink down-pointing triangle), y = 0.38 (green diamond), y = 0.49 (navy left-pointing triangle), y = 0.59 (violet right-pointing triangle), y = 0.68 (purple hexagon), y = 0.77 (brown star), y = 0.85 (mustard pentagon), y = 0.93 (blue sphere), and y = 1 (turquoise cross).

solvent) intermolecular interactions are preferred with respect to heterogeneous (solute:solvent) intermolecular interactions. In line with the trend observed for the solubility studies, toluene solutions were found to show the most ideal behaviour, with larger deviations from ideal behaviour being observed with increasing acetone composition. This behaviour is consistent with toluene having much stronger intermolecular interactions with eicosane comparative to acetone, with high content acetone solutions not only having likely stronger solute:solute interactions, but also likely stronger solvent:solvent interactions, making for easier eicosane desolvation and hence crystallisation from solution. This is evidenced in the calculated  $\Delta H_{\rm diss}$  values, in Table 2, highlighting that, in general,  $\Delta H_{\rm diss}$  is larger at the higher solution acetone compositions.

However,  $\Delta H_{\rm diss}$  was found to decrease with increasing acetone composition above y=0.85, which, given the low solubility of eicosane in high acetone compositions solutions, would seem to be counter-intuitive. However, the change in van't Hoff plot slopes for these solutions suggest decreasing effects of enthalpy and entropy within solution. This is corroborated with a similar trend in the enthalpy of mixing results, suggesting that the solute:solvent interactions decreased in strength with increasing acetone solution content, until a critical acetone composition created a change in the ratio between solute:solvent:solvent and solute:solvent interactions.

Interestingly, the slopes of the van't Hoff plots were found to be relatively similar for solutions with excess toluene and for higher solution acetone content above y = 0.85. Deviations found for the mixed-solvent solutions with excess acetone

Table 2 Thermodynamic parameters and activity coefficients (1) as a function of temperature calculated from van't Hoff analysis

Acetone mol fraction (y)	Enthalpy of dissolution (kJ mol <sup>-1</sup> )	Entropy of dissolution $(kJ \text{ mol}^{-1} \text{ K}^{-1})$	Enthalpy of mixing (kJ mol <sup>-1</sup> )	Entropy of mixing (kJ mol <sup>-1</sup> K <sup>-1</sup> )	Concentration range (g L <sup>-1</sup> )	$ \ln \Upsilon = aT + c $
0	78.36	0.255	35.50	0.116	250-850	-0.05 T + 1.60
0.14	93.99	0.307	51.13	0.169	200-800	-0.07 T + 2.16
0.26	116.28	0.382	73.42	0.243	200-500	-0.11  T + 3.01
0.38	132.89	0.436	90.02	0.297	200-500	-0.13 T + 3.76
0.49	130.71	0.425	87.85	0.286	150-450	-0.12  T + 4.12
0.59	155.35	0.504	112.48	0.365	150-450	-0.16 T + 5.32
0.68	182.22	0.590	139.36	0.452	150-450	-0.19 T + 6.64
0.77	198.03	0.638	155.17	0.500	100-400	-0.21  T + 7.73
0.85	224.72	0.721	181.86	0.582	75-375	-0.24 T + 9.38
0.93	144.45	0.448	101.58	0.309	20-140	-0.14  T + 7.24
1	100.52	0.296	57.65	0.157	5–35	-0.08 T + 6.41

content demonstrated changing temperature dependence of solution activity, and as such, solution structure, as a function of composition, as outlined by the varying values of aT in Table 2.

Values of  $\Delta H_{\rm diss}$ ,  $\Delta S_{\rm diss}$  and the range of  $\Upsilon$  calculated for eicosane in toluene solution were close to those calculated in a previous study by Tang *et al.*, <sup>25</sup> which focussed on hexadecane and octadecane mixtures crystallising from three fuel representative solvents, including toluene. Values of around 69 kJ mol<sup>-1</sup>, 0.24 kJ mol<sup>-1</sup> K<sup>-1</sup> and the range 1.40–1.59 were calculated in the aforementioned study, for  $\Delta H_{\rm diss}$ ,  $\Delta S_{\rm diss}$  and the range of  $\Upsilon$  over the concentration range studied, respectively.

#### 3.3. Analysis of the nucleation kinetics

For eicosane in mixed toluene: acetone solutions the resultant nucleation mechanisms, detailed in Table 1, were found to vary as a function of solution composition and concentration, in line with the trends observed for the solution thermodynamics and MSZW analysis. For solution compositions from 0 < y < 0.49 and y = 1, the nucleation mechanism was found to be predominantly IN, which is consistent with a previous study that focussed on the nucleation of hexandecane: octadecane mixtures from single solvent solutions, with both alkanes being found to predominantly nucleate *via* IN pathway. In contrast, for solution compositions ranging from 0.59 < y < 0.93, the predominant nucleation mechanism was found to be PN.

For the PN cases, the calculated effective interfacial tensions, critical nuclei radius and number of molecules within the critical nuclei are given in Table 3. Comparatively, low values of effective interfacial tension, ranging from 0.38–1.94, were calculated for all PN systems studied, consistent with data on other long-chain hydrocarbons, <sup>28</sup> and consistent with the nucleation process being predominantly heterogeneous. The latter may also explain the low calculated values for the critical nucleus size and number of molecules within the critical nucleus. Nonetheless, the overall behaviour of interfacial tension as a function of composition was found to closely mirror the variations observed for crystallisability

and solution thermodynamics over the range of compositions studied.

#### 3.4. Compositional trends

Examination of the results of the crystallisability, solution thermodynamics and nucleation kinetics data clearly demonstrate consistent and concurrent trends over the full compositional range for the eicosane in toluene: acetone mixed-solvent system. For all solution compositions, the average MSZWs, over the range of concentrations studied, were found to increase for compositions ranging from 0 < y < 0.85. This was subsequently followed by a decrease in average MSZW with increasing solution acetone content to a solution composition of y = 1. This same trend was observed for values of enthalpies of dissolution and mixing, the average slope

**Table 3** Values of effective interfacial tension ( $\gamma_{\rm eff}$ ), critical nuclei radius ( $r^*$ ) and number of molecules within the critical nuclei ( $i^*$ ) for the PN systems

Conc. $(g L^{-1})$	$\gamma_{eff} \left( mJ \ m^{-2} \right)$	r* (nm)	i*
y = 0.59			
250	0.38	0.28 - 0.12	0.18-0.01
350	0.86	0.62-0.29	1.95-0.20
450	0.87	0.62 - 0.28	2.20-0.19
y = 0.68			
150	0.89	0.51-0.23	1.10-0.10
250	1.19	0.73-0.35	3.26-0.37
350	1.41	0.72 - 0.36	3.14-0.40
450	1.02	0.54 - 0.26	1.33-0.14
y = 0.77			
100	1.15	0.56-0.26	1.44 - 0.14
200	1.16	0.61 - 0.28	1.92-0.19
300	0.96	0.52-0.23	1.20-0.10
400	1.94	0.72 - 0.40	3.06-0.53
y = 0.85			
75	1.54	0.64 - 0.32	2.23-0.28
175	1.72	0.73-0.39	3.19-0.50
275	1.62	0.67 - 0.34	2.48-0.32
375	1.86	0.71-0.38	3.03-0.46
y = 0.93			
60	1.19	0.81 - 0.40	4.46 - 0.55
100	1.00	0.77-0.39	3.82-0.50
140	1.28	0.84 - 0.45	4.89-0.77

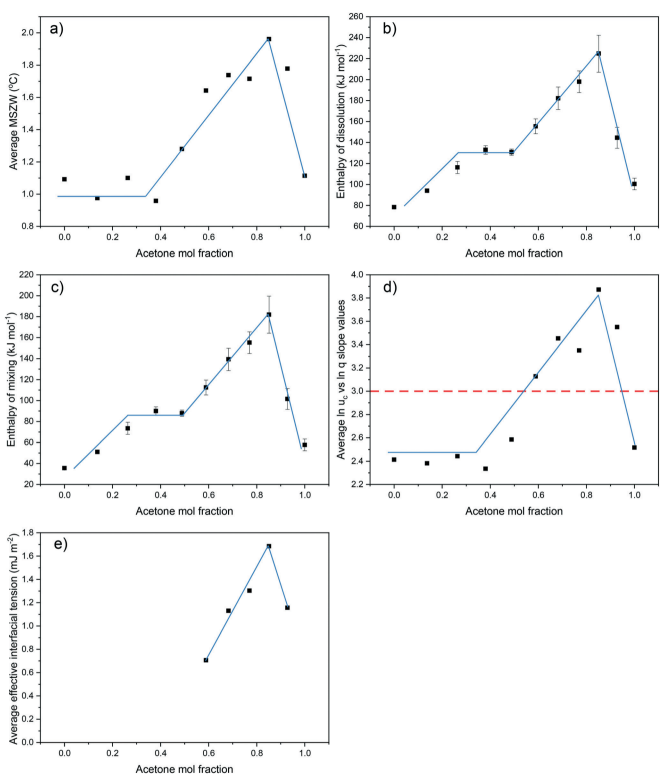


Fig. 7 Trends in values obtained over the compositional range studied for crystallisability, solution thermodynamics, nucleation mechanism and effective interfacial tension. a) Average MSZW values over the concentration ranges studied; b) enthalpy of dissolution; c) enthalpy of mixing; d) average slope values obtained from  $\ln q$  vs.  $\ln u_c$  over the concentration ranges studied; e) average effective interfacial tension values calculated in the range of 0.59 < y < 0.93. Red dotted line in d) represents transition from IN (<3) to PN (>3).

values from linear regressions fitted to plots of  $\ln u_c$  vs.  $\ln q$ , over the full compositional range studied and the values of effective interfacial tension, Fig. 7. This demonstrates a clear

link between the crystallisability, solution thermodynamics and nucleation kinetics and mechanism, which are all dominated by the solution structure. CrystEngComm Paper

#### 3.5. Intermolecular grid-search modelling

Results summarising the calculated solvation sites for a single eicosane molecule solvated with toluene and acetone, using different low-pass energy cut-off filters, are given in Fig. 8. Examination of the energy-filtered data reveals that both toluene and acetone can easily solvate the surfaces of eicosane, with the strongest binding interactions with the carbon backbone of the eicosane molecule, where the number of potential atom-atom van der Waals interactions is highest. However, higher energy cut-off filtering revealed that acetone's solvation propensity diminished quite rapidly. In contrast, toluene was found to maintain strong solvation propensity over the majority of the eicosane molecule, even at the highest energy cut-offs. The full solvation site analysis is given in ESI† S3.

Fig. 9 shows the distribution of binding site energies as a function of the number of calculated interactions. As might be expected on the basis of molecular weight, the strength of intermolecular interaction energies of eicosane with other eicosane molecules were much higher than for other possible solution interaction types, correlating well with the less than ideal solution behaviour observed. Furthermore, in terms of solute: solvent interactions, toluene was found to interact more strongly with eicosane than acetone was, in line with measured relative solubilities, with acetone: eicosane interactions found to be the weakest interaction type calculated, suggesting that in solution acetone may have been more likely to interact with either itself or toluene, instead of eicosane. This is in line with calculated atomic fractional charges for the studied molecules, provided in ESI† S2, which demonstrate increasing molecular polarisability from eicosane < toluene < acetone. Acetone, therefore, would be expected to have stronger electrostatic interactions with itself or toluene than with eicosane in the solution environment.

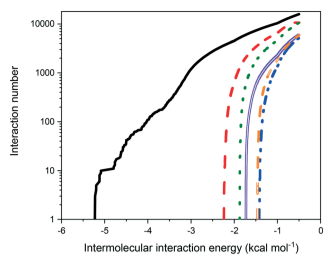
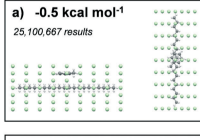
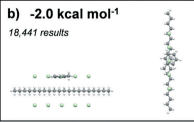


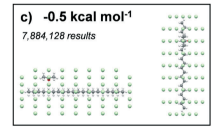
Fig. 9 Calculated intermolecular interaction energies for possible interaction types in eicosane in toluene: acetone solution, over the total interactions calculated. High-energy cut-off set at -0.5 kcal mol<sup>-1</sup>. Coloured lines represent different intermolecular interactions: eicosane: eicosane (black solid line), eicosane: toluene (red dashed line), eicosane: acetone (blue dot-dash line), toluene: toluene (green dotted line), acetone:acetone (purple double line) and toluene: acetone (orange double dashed line).

Fig. 10 summarises the results of the solute:solvent cluster building simulations, highlighting the predicted solvation shell structures based upon different toluene: acetone ratios. The simulation data reveals that for the 10:10 and 15:5 ratios of toluene: acetone, the acetone molecules preferentially bind to the toluene molecules, creating, what is effectively, a 'second shell' around the primary toluene solvation shell around eicosane rather than directly solvating eicosane by filling the potential empty solvation sites.

For the 5:15 ratio of toluene: acetone intermolecular grid search results the acetone molecules were still found to create







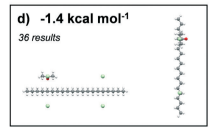


Fig. 8 Interaction energy fields of toluene (a and b) and acetone (c and d) probe molecules, with a central eicosane molecule. Toluene and acetone molecules show most negative interaction energy site. The green dots represent interaction locations of the probes, which passed the set energy filter cut-offs that are highlighted at the top left of images.

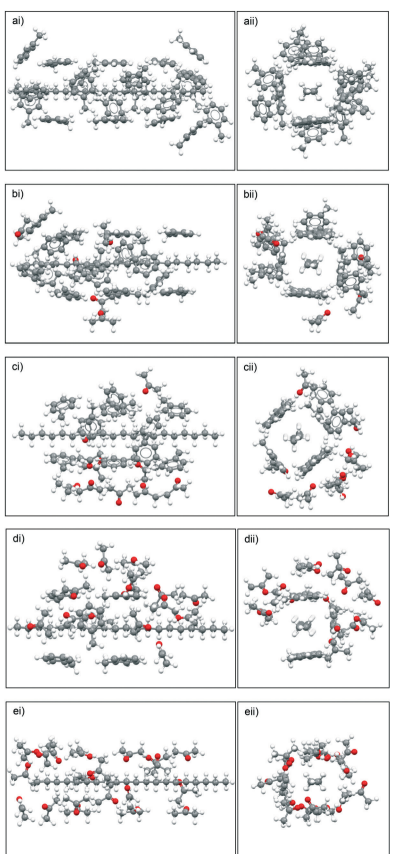


Fig. 10 Solvent clusters built using the intermolecular grid-search method around eicosane molecules, in different toluene: acetone ratios, with toluene molecule interactions determined first, followed by acetone molecules. a) 20:0; b) 15:5; c) 10:10; d) 5:15 and e) 0:20. i) and ii) represent different views of solvation clusters along and down the carbon backbone of eicosane, respectively.

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a 'second' solvation shell around the toluene molecules that were solvating the eicosane molecule, but some free eicosane solvation sites were filled with acetone molecules. This suggests that when the ratio of acetone to toluene was high enough, the acetone molecules would more strongly solvate the eicosane molecule. Therefore, for the studied solutions, above a critical acetone composition the disruptive solvent:solvent interaction effect is likely to be somewhat overcome.

Examination of the calculated solvation energies based on these cluster structures, as well as the intermolecular solvent: solvent interaction energies within the solvation shell are given in Table 4. In general, the total solvation energy was found to decrease with an increase in acetone composition, in line with the trend observed in the measured solubility over the mixed-solvent compositional range studied. However, a slight increase in solvation energy was calculated from a toluene: acetone ratio of 5:15 to 0:20. This correlates well to the observed solubility behaviour at lower solution temperatures, whereby eicosane solubility increased from solutions with a high acetone composition to pure acetone solutions and suggests a changing nature of solvent: solvent interactions affecting solvation power.

The solvent: solvent interactions in the solvation shell were found to be strongest for 10:10 toluene: acetone with decreased total intermolecular interaction energies for 5:15 and 15:5 toluene: acetone ratios. This further suggests that strong toluene: acetone interactions were present in the mixed solvent solutions, with a decrease in interaction intensity when one of the solvents is in excess in solution.

## 4. Discussion

Intermolecular grid-search results suggest that acetone molecules could be more likely to 'solvate' toluene molecules and disrupt the solute: solvent interactions, until a critical acetone composition is reached whereby acetone molecules solvate eicosane more readily. These predictions provide insight into the increasing enthalpies of dissolution and mixing values up to a solution composition of y = 0.85, due to strong interactions between acetone and toluene in the solvation shell, decreasing the solvation power towards eicosane and, as such, decreasing solute: solvent interaction strengths, followed by a decrease in enthalpic values, due to increased acetone solvation.

This behaviour also provides a potential insight into the similar trends in MSZW and changes in the nucleation

Table 4 Solvation energies and intermolecular interaction energies between toluene and acetone molecules in the solvation shell for the solvation of eicosane at different toluene: acetone molecular ratios

Toluene: acetone	Interaction energy (kcal mol <sup>-1</sup> )					
molecular ratio	Total solvation	Toluene : acetone				
20:0	-54.71	_				
15:5	-52.39	-27.80				
10:10	-46.54	-41.96				
5:15	-38.84	-30.69				
0:20	-43.72	_				

mechanism. An IN pathway can represent solutions containing stable prenucleation clusters and at a critical level of supersaturation, there is sufficient driving force for nucleation to occur for all clusters present. Practically, this represents the case whereby all nucleation events within the solution occur at one instance, with only crystal growth occurring after this stage. In contrast, PN suggests that the prenucleation clusters could have a lower stability, therefore only certain clusters would be able to overcome the free energy barrier required for nucleation to occur at one instance of time, resulting in concomitant crystal nucleation growth processes occurring within solution crystallisation progresses in the system.

For the PN systems, the solution environments had an excess of acetone molecules, with respect to toluene. This change in nucleation mechanism perhaps suggests that when acetone is in excess in solution, there was a more complicated solution structure, whereby the solvation layer around eicosane molecules is more disordered and perhaps more polar, containing a high number of heterogeneous and homogeneous solvent: solvent interactions, in line with the grid-search results. Such a structure could disrupt the diffusive transport of eicosane molecules into the prenucleation clusters, seen toluene is in excess, given acetone's limited intermolecular interactions with eicosane. This would have the effect of lowering the attachment frequency of eicosane to a growing cluster, hence reducing the nucleation rate constant,  $K_{\rm I}$ , and, as such, the rate of nucleation (eqn (12)). Thus, only some of the eicosane clusters might reach a critical cluster size and be able to nucleate, before a stable equilibrium in solution was reached. This would result in the case of nucleation events occurring over time, creating concomitant nucleation and growth processes at different locations within solution, in line with the PN mechanism.

In contrast, for the IN cases the solutions either had an excess toluene component with respect to acetone or a single solvent environment, with potentially less disruptive heterogeneous solvent: solvent interactions and hence a lower barrier to the formation of stable prenucleation clusters within solution. Therefore, a distribution of stable prenucleation clusters could be able to form prior to the nucleation stage, once a critical level of supersaturation corresponding to the limit of the MSZW had been reached, with all clusters nucleating at once. Although speculative, at this stage such a model could also explain the increase in MSZW with increasing acetone composition, and as such a lower ease of crystallisation, until a critical acetone composition was reached, in which this behaviour was reversed. Further support for this comes with the calculated values of effective interfacial tension, which mirror the behaviour of the MSZW with mixed-solvent composition, notably in such a case that a higher level of supersaturation would be needed to overcome the energy barrier required to create a solid nucleus surface. This suggests that strong interactions between acetone and toluene molecules might lead to weaker interactions with eicosane clusters, thereby

increasing the barrier to the formation of nuclei surfaces, driving up the effective interfacial tension and lowering the crystallisability. This would be consistent with an increase in the thermodynamic parameter b associated with the nucleation rate expression (eqn (12)), which would reduce the rate of nucleation of eicosane in these solutions. This, coupled with a reduction in the nucleation rate constant, would essentially create a more thermodynamically controlled nucleation process, more consistent with the

The resulting changes in nucleation behaviour and mechanism as a function of mixed-solvent composition, also align well with previous studies that have demonstrated that, in single solvent solutions, the IN mechanism is associated with stable prenucleation clusters in solution<sup>24</sup> and a shift towards a PN pathway can be associated with a disruption of the ability of solutions to form stable clusters prior to nucleation.<sup>23</sup> This demonstrates a wider applicability of the results outlined in this study towards other crystallising systems.

progressive nucleation pathway.

The changing nature of solution behaviour above a critical acetone solution content at around y = 0.85 could also represent increased solvent structuring in solution, as demonstrated by decreasing values of the entropy of dissolution and mixing. Acetone is known to structure through strong carbonyl intermolecular interactions in its crystal structure and liquid state, 50 which could enable free interactions between the methyl groups and solute molecules within solutions, with a high concentration of acetone molecules. This could provide an additive solubilising effect towards eicosane in solution alongside the more strongly interacting toluene molecules, which could be responsible for the observed increases in solubility at lower solution temperatures, for solutions with a high acetone content above y = 0.85. However, further study is needed to investigate this possible behaviour.

## 5. Conclusions

Eicosane in mixed toluene: acetone solvent solutions was studied over a range of solution compositions from pure toluene to pure acetone, revealing comparatively small MSZWs for all solution compositions in comparison to other organic crystallising systems, highlighting their ease of crystallisability. A broadly inverse relationship between solution concentration and MSZW was found for solution compositions at acetone mol fractions (y) of 0 < y < 0.49 and 0.93 < y < 1. In contrast, the opposite relationship was observed for solution compositions of 0.53 < y < 0.85.

The solubility of eicosane was found, in general, to decrease with increasing solution acetone content. At higher solution temperatures, small additions of acetone to pure toluene solutions were found to increase solubility, whereas at lower solution temperatures this behaviour was not observed and when acetone was in excess, further additions of acetone increased solubility above a solution composition of y = 0.85. All solutions revealed less than ideal behaviour,

with activity coefficient temperature dependency altering as a function of mixed-solvent composition. Both the enthalpies of dissolution and mixing were found to increase over the compositional range of 0 < y < 0.85, followed by a decrease with higher acetone content, demonstrating a change in solution chemistry above a critical solution acetone composition.

Intermolecular grid search results from energy cut-off calculations were in good agreement with the solubility findings, with toluene molecules being found to interact much more strongly with eicosane than did acetone molecules. The strongest solvation sites were found to be along the carbon backbone of the eicosane molecule, where closer atom-atom contacts and strong van der Waals interactions were readily available. Intermolecular interaction energies revealed eicosane: eicosane interactions to be the strongest, in line with the less than ideal behaviour observed for all solutions. Acetone was found to interact more strongly with itself and toluene than with eicosane, suggesting that it would have limited interaction with eicosane in a mixed-solvent solution environment.

Analysis of nucleation kinetic data revealed that solution compositions in the range 0 < y < 0.49 and y = 1, nucleated via an IN mechanism, whilst in contrast, solution compositions ranging from 0.59 < y < 0.85 were found to predominantly nucleate via the PN pathway, the latter suggesting a more thermodynamically-controlled system. The kinetic and thermodynamic dependencies of the nucleation of these solutions were found to correlate to the dependency of the MSZW vs. the solution concentration behaviour observed. PN solutions crystallised with low effective interfacial tensions, in line with previous studies on alkane systems, but were found to increase with increasing acetone composition up to y = 0.85, above which, the effective interfacial was found to decrease again.

The key crystallisation process metrics were all found to follow the same consistent behaviour as a function of mixedsolvent composition, notably: MSZW, enthalpies dissolution/mixing, interfacial tension and the balance between the thermodynamic and kinetic factors on the nucleation process, calculated by the IN/PN determination slopes. Analysis of solvation energies and cluster structures was found to provide an insight into the potential intermolecular clustering behaviour within this mixed-solvent solution, notably acetone interacts more strongly with toluene than with eicosane, with the latter not being effectively solvated by acetone molecules in the mixed-solvent solutions. A 'cage' of acetone molecules created due the stronger toluene: acetone interactions limited the ability of eicosane to form stable prenucleation clusters throughout solution, altering the nucleation mechanism from IN to PN when acetone was in excess. Above a critical acetone content this effect was overcome.

Overall, the solution chemistry and structure were found to directly relate to crystallisation behaviour and this was integrated within a methodological workflow, developed to Abbroviations

aid future research into mixed solvent solution crystallisation processes.

#### 5.1. Future work and current limitations

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The molecular grid-search based modelling of indicative solution structure could be enhanced in future work through its integration with more atomistically detailed modelling molecular dynamic (MD) simulations, which would have the potential to provide a more complete and extended structural insight into the solution environment, associated with solvation by mixed solvent systems. Through this, the longerrange solution structure and its relationship, as well as its relationship with the solute mass transfer, could be directly related to the subsequent nucleation kinetics.

Furthermore, small angle neutron scattering (SANS) of mixed-solvent systems using deuteration to confirm specific solvent species within the mixed-solvent environments, could potentially validate the solvation structural ordering proposed in this work.

## Nomenclature

KBHR PN parameter  $a_1$ KBHR PN parameter  $a_2$ Dimensionless thermodynamic parameter Molar enthalpy of dissolution (kJ mol<sup>-1</sup>)  $\Delta H_{\rm diss}$  $\Delta H_{\rm f}$ Enthalpy of fusion (kJ mol<sup>-1</sup>)  $\Delta H_{\rm mix}$ Molar enthalpy of mixing (kJ mol<sup>-1</sup>)  $i^*$ Number of molecules in the critical nucleus Nucleation rate (nuclei m<sup>-3</sup> s<sup>-1</sup>) Boltzmann constant (m<sup>2</sup> kg s<sup>-2</sup> K<sup>-1</sup>) k $K_{\rm J}$ Nucleation rate constant (m<sup>-3</sup> s<sup>-1</sup>) Nuclei numerical shape factor  $k_{\rm n}$ λ Molecular latent heat of crystallisation (J) Number of crystallites at the detection point  $N_{\rm det}$ Cooling rate (°C min<sup>-1</sup>) qKBHR PN parameter  $q_0$ Ideal gas constant (J K<sup>-1</sup> mol<sup>-1</sup>) R Critical nucleus radius (m) Molar entropy of dissolution (kJ K<sup>-1</sup> mol<sup>-1</sup>)  $\Delta S_{\rm diss}$ Molar entropy of mixing (kJ K<sup>-1</sup> mol<sup>-1</sup>)  $\Delta S_{\rm mix}$ TSolution temperature (°C)  $\Delta T_{\rm c}$ Undercooling (°C) Crystallisation temperature (°C)  $T_{\rm c}$ Crystallisation temperature at the kinetic  $T_{\rm c,l}$ limit/equilibrium crystallisation temperature (°C)  $T_{
m diss}$ Dissolution temperature (°C) Equilibrium saturation temperature (°C)  $T_{\rm e}$ Melting point (K)  $T_{\rm m}$ Relative critical undercooling  $u_{\rm c}$  $v_0$ Volume occupied by a solute molecule in the crystal (m<sup>3</sup>) Molar solubility  $\chi$ Molar solubility in ideal state  $x_{ideal}$ 

## **Abbreviations**

DSC Differential scanning calorimetry

IN Instantaneous nucleation

KBHR Kashchiev-Borissova-Hammond-Roberts

MD Molecular dynamics
MSZW Metastable zone width
PN Progressive nucleation

SANS Small angle neutron scattering

SE Standard error

## Conflicts of interest

The authors declare no competing financial interest.

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Effective interfacial tension (mJ m<sup>-2</sup>)

Mol fraction acetone

Activity coefficient

γ

γeff

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