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Multiscale analysis of triglycerides using X-ray scattering: implementing a shape-dependent model for CNP characterization[†]

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In the last decade, research has focused on examining the fundamental interactions occurring in triglycerides, aiming to comprehend the self-assembly of crystalline nanoplatelets (CNPs) and their role in forming larger hierarchical structures essential for fat functionality. Microscopy research on CNPs frequently requires disruptive preparatory techniques, such as deoiling and sonication, to achieve quantitative outcomes. Conversely, X-ray scattering has proven to be an advantageous method for studying triglycerides, as little sample is needed to quantify the system's hierarchical structures. Specifically, ultra-small-angle X-ray scattering (USAXS) has emerged as a fitting technique for studying CNPs, owing to its length scale range falling between 25 nm and 3.49 µm. In this study, we characterized four different 30% fat dilutions of stearic acid-based fats in triolein, with various purities and preparation protocols. Samples were characterized by combining diverse microscopy techniques (cryo-SEM, TEM, polarized light and phase contrast microscopy) with synchrotron-radiation X-ray scattering (WAXS, SAXS, and USAXS). A shape-dependent model for the interpretation of USAXS data is proposed, overcoming some of the drawbacks linked to previously utilized models. CNPs are modeled as polydisperse parallelepipeds, and the aggregates are characterized by fractal dimensionality. This model offers novel insights into CNP cross-section, as well as aggregation. In the long run, we hope that the model will increase our understanding of CNP conformation and interactions, helping us design new fat systems on the mesoscale.

1. Introduction

Over the past few decades, attention has been focused on studying the primary building blocks of triglycerides.^{1–8} The core goal is to understand how edible fat molecules self-assemble and how their aggregation plays a role in the formation of macroscale structures responsible for fat functionality.⁹

Edible fats are complex food systems with intricate geometrical arrangements.² During the crystallization process of triglycerides (TAG), TAG molecules self-assemble into distinct structures at different length scales. A graphical representation is provided in Fig. 1. At a nanoscopic level, inter- and intramolecular interactions between TAG molecules result in packing arrangements (polymorphs - α , β' , and β) and specific configurations (2L or 3L).¹⁰⁻¹⁴ TAG molecules then selfassemble into lamellar phases, which stack epitaxially due to van der Waals forces forming anisotropic crystalline nanoplatelets (CNPs).^{9,13,15,16} CNPs then aggregate into larger agglomerates, forming different domain sizes, including crystallites and crystal clusters.¹ Eventually, polycrystalline spherulites, also known as flocs, will interact with each other, determining the macrostructure and functionality of soft matter fat systems. Each level greatly influences the functionality and applicability of lipids, as the overall size and morphology of the different aggregation levels determine key macroscopic properties (i.e., texture, solid fat content, and oil binding capacity).^{9,13}

The first systematic study to identify the primary unit was done by Acevedo and Marangoni in 2010.² The authors set up a series of transmission electron microscopy (TEM) experiments that allowed the identification of the primary building blocks,

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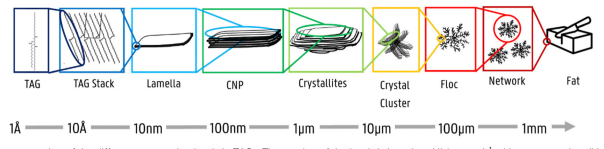


Fig. 1 Representation of the different aggregation levels in TAGs. The naming of the levels is based on Mishra *et al.*,¹ with one exception: "domain" is here replaced by crystalline nanoplatelet ("CNP").¹⁷

so-called CNPs. These CNPs, formed by the stacking of lamellae, are considered to be the basis of the complex supramolecular structural hierarchy in TAG crystal networks.¹⁸ A considerable number of experiments were performed to describe and quantify these CNPs, seeking to understand how composition and processing parameters would alter their nature.²⁻⁸ Alternatively, a more recent study conducted by Mishra et al.1 successfully utilized atomic force microscopy (AFM) to characterize crystallites and their corresponding lamellar domains. Their findings suggest that within 1 nm and 10 µm, there are at least three different domain sizes, namely lamellae (at the nanometre level), domains (here named CNPs, of about hundred nanometres) and crystallites (at the micrometre level). These methodologies, however, require intense disruption of the fat crystal network, as preparatory methods such as deoiling and sonication are necessary to obtain quantitative results.

In order to overcome the burdens of sample preparation, in 2013, X-ray scattering was first used to study CNPs. X-ray scattering is a powerful technique to study the submicroscopic structural organization of materials. It allows the determination of the dimensionality at different length scales by analysing the measured scattering intensities using appropriate structural models.¹⁹ X-ray scattering is a well-established technique that has helped the understanding of several multi-level hierarchical systems.²⁰ It is a particularly well-suited technique for investigating concentrated and opaque samples, such as TAGs, which are prone to structural damage when performing microscopic experiments (*i.e.* deoiling).^{9,21}

Peyronel *et al.* pioneered the use of USAXS for the study of CNPs. The first USAXS experiments reported for fat crystal networks were conducted in the Advanced Photon Source, Argonne National Laboratory, utilizing a Bonse-Hart instrument that allowed data collection in the range of 30 nm to 10 μ m.³

In the context of their research, two analytical models were utilized to process USAXS data. The first model utilized is the unified fit model.³ Although helpful, the unified fit model fails to account for the shape of the particle. In fact, in the calculation of the CNP equivalent radius (R_{CNP}), the model assumes that the CNPs are monodisperse spheres, an assumption known to be false and that can yield an overestimation of the actual scattering particle.^{21,22} Therefore, this model disregards the anisotropic nature of the primary scatterer. In addition, the unified fit model has certain drawbacks, as it remains relatively unstable at the

crossover point from the Guinier region to the power law regime (which occurs at a rather arbitrary value of q).^{21,23} The interpretation when analysing polydisperse multi-level systems, such as triglycerides, must be done with utmost care.

To compensate for the lack of shape-dependency of the unified fit model, a second model is suggested. In the context of triglycerides, the Guinier–Porod model has only been utilized to obtain the *s*-factor, indicative of the shape of the primary scattering units.³

Although the efforts so far have been considerable, little to no attention has been put on the morphological features of the CNPs. In fact, many of these morphological features have not been taken into account when studying fat crystal networks and, therefore, have not been implemented in the modelling, interpretation or design of new fat systems.

In this article, we propose a new model for the interpretation of USAXS data. The proposed model responds to the need for modelling the data based on assumptions in real space rather than curve fitting. In fact, unlike previous models, this is a shape-dependent model, and the output contains information on the cross-section of CNPs, as well as their aggregation. Thus, by modelling the USAXS range, we were able to fully characterize the mesoscale of triglyceride systems, from the primary building blocks (CNPs) to their aggregation.

To engage in the model discussion, we presented certain measurements as examples. We combine wide-angle X-ray scattering (WAXS), small angle X-ray scattering (SAXS) and ultra-small angle X-ray scattering (USAXS) to study, at various length scales, four different stearic acid based systems (from the angstrom level up to the micrometre level). We characterize the SAXS and USAXS ranges with different available models and compare the output. Whenever possible (in the case of length scale overlap), the findings were compared to *ex situ* microscopic techniques, such as transmission electron microscopy (TEM) or cryo-SEM (cryogenic scanning electron microscopy), which allowed us to validate the findings of the scattering models.

2. Materials and methods

2.1 Materials

Triolein (OOO) (>99%, CAS 122-32-7) and tristearin (SSS) (>99%, CAS 555-43-1) were acquired from Nu-Chek-Prep Inc

(Elysian, MN, USA). Fully hydrogenated rapeseed oil (FHRO) and high oleic sunflower oil (HOSO) were kindly provided by Vandemoortele R&D (Izegem, Belgium). The predominant fatty acid in FHRO was stearic acid (18:0) at around ~90.2%. Small amounts of palmitic acid (16:0) and arachidic acid (20:0) were present at ~5.4% and ~1.8% respectively. Acetone (>99%, CAS 67-64-1), *n*-hexane (>99%, CAS 110-54-3) and isobutanol (>99%, CAS 78-83-1) were acquired from ChemLab (Zedelgem, Belgium). Quartz glass capillaries (diameter <1.5 mm, 80 mm length, 0.01 mm thickness, Q-15-001-80) were acquired from WJM-Glas Müller GMBH (Berlin, Germany). Polypropylene disposable syringes (1 ml INJEKT[®] Solo, B. Braun, Melsungen, Germany) and needles (STERICAN[®] needles 21 G 0.8 mm × 120 mm) were utilized to transfer the TAG dilutions into the quartz capillaries.

2.2 Statically crystallized sample preparation

A 30% (w/w) dilution of tristearin (SSS) in triolein (OOO) was meticulously weighed (1 g) in a 5 ml beaker. The sample was melted to 80 $^{\circ}$ C for at least 15 minutes. Once molten, the sample was transferred into a quartz capillary using a preheated disposable polypropylene syringe. Each capillary was measured to make sure the diameter did not exceed 1.3 mm. The procedure was then repeated for a second dilution of 30% (w/w) FHRO in HOSO (200 g in a 250 ml beaker).

Samples were crystallized four weeks before synchrotron experiments. For this, an altered HFSX350 Linkam stage that allows precise temperature control of samples in capillaries was utilized (Xenocs, Grenoble, France). Samples were melted to 80 $^{\circ}$ C for 15 minutes to erase crystal memory, after which they were crystallized following the appropriate cooling protocol. Samples were stored in a thermostatic cabinet until transported to the ESRF in a portable mini car refrigerator set at 15 $^{\circ}$ C (Vevor, USA).

2.3 Pilot scale sample preparation

Pilot scale samples were prepared with a benchtop scraped surface heat exchanger (SSHE) (Het Stempel B.V., Zwijndrecht, The Netherlands). The process was carried out using a micropump and a micro-SSHE (a volume of 29 ml), equipped with a set of scraping knives on the rotor axis. This device allowed for shear crystallization, mimicking industrially relevant conditions. For further specifications on the device, the reader is referred elsewhere.²⁴ For the trial, 4 kg of 30% dilution of FHRO in HOSO were used as the fat base. The inlet temperature of the melt was set to 77 °C, the flow rate was set to 5 kg h^{-1} and the SSHE rotation had a value of 500 rpm. For the X-ray scattering experiments, a small amount of sample was gathered immediately after production (in the outlet point) and placed in-between a galvanized metallic washer (5.3 \times 10 mm). The sample was sealed in between two 9 mm Kapton tape dots (0.05 mm thickness) (PrintTec B.V., Geldermalsen, The Netherlands). For the microscopy experiments, the crystallized fat system was stored in 200 ml containers. Samples were then stored in a thermostatic cabinet set at 15 °C until transported to the ESRF in a portable mini car refrigerator (also set at 15 °C) (Vevor, San Bernardino, CA, USA).

2.4 Cryo-SEM microscopy

Visualization of CNPs was performed utilizing a cryogenic electron scanning microscope (cryo-SEM). For this, a threestep methodology was utilized. To ensure replicability of the procedure, visualizations were replicated at least twice.

2.4.1 Sample preparation. Statically crystallized fat systems were prepared in an X-ray-altered Linkam shear cell utilizing a 1500 μ m gap (CSS450, Linkam, Salfords, UK) (Xenocs, Grenoble, France). Samples were first heated to 80 °C for 15 minutes to erase any crystal memory. The system was then cooled at 1 °C min⁻¹ or 10 °C min⁻¹ to 15 °C, where it was kept isothermally at 15 °C for at least 15 minutes and transferred to a pre-tempered glass vial. Samples were then stored in a thermostatic cabinet at 15 °C until deoiling.

2.4.2 Sample deoiling. A small amount of the crystallized fat sample was dissolved in an excess of pre-tempered (15 $^{\circ}$ C) solvent solution of 80/20% v/v hexane/acetone.²⁵ Every 12 h, the solvent was renewed, and the exchange continued. The procedure was repeated twice to ensure the removal of the liquid oil and disruption of the supra-CNP network. One drop of suspension was placed on top of a double-sided adhesive conductive carbon disc (SPI supplies, West Chester, USA) attached to a SEM aluminium stub. The excess solvent was allowed to evaporate for at least 4 hours.

2.4.3 Visualization. The stub was vitrified in a liquid nitrogen slush (-210 °C) and transferred into a PP30100T cryo-transfer system (Quorum Technologies Ltd, East Sussex, UK) conditioned at -140 °C. The sample was then sublimated (-70 °C for 90 min) and sputter-coated with platinum using argon gas. The visualization was performed in a Jeol JSM-7100F TTLS LV TFEG-SEM (Jeol Europe BV, Zaventem, Belgium) under high-vacuum conditions (1×10^{-6} mbar) at -140 °C and at an accelerated voltage of 3 keV. At least 50 different images were taken at different sample locations to ensure that results were representative of the sample.

The software Image J Fiji was utilized for size measurements. At least 50 different measurements were performed per sample.

2.5 Transmission electron microscopy (TEM)

For the TEM visualization, sample preparation and sample deoiling followed the same procedure as described in the cryo-SEM section. Prior to the visualization, negative staining was necessary. For this, solutions with the lipid crystals were diluted in 1 M trisbuffered saline (TBS). C-FlatTM carbon film gold 300 mesh EM grids (Jena Bioscience, Jena, Germany) were placed in a glow-discharger. Then, 4 μ l of the sample was deposited onto the EM grids and left to incubate for 1 min. Excess solution was removed by quickly touching the grid edge with filter paper. Subsequently, the EM grids were washed twice with tris-buffer and aqua dest before transferring the grids to a drop of 2% uranyl acetate. After 1 min, the excess solution was removed, and the EM grids were air-dried before imaging. EM grids were evaluated in a transmission electron microscope, Tecnai G2 Spirit BioTWIN (FEI, Eindhoven, The Netherlands) at 120 kV.

2.6 Polarized light microscopy

Polarized light micrographs were captured employing a Leica DM2500 microscope (Wetzlar, Germany) equipped with a Leica MC170 HD colour camera. A small amount of sample was placed between a glass slip and a coverslip and then placed under the microscope. For the statically crystallized samples, a Linkam PE120 Peltier system equipped with a T95 system controller was used. Each sample was brought to 80 °C for 15 minutes to erase crystal memory. Crystallization then occurred as the sample was cooled down from 80 °C to 15 °C at a rate of 1 °C min⁻¹ or 10 °C min⁻¹ (slow *vs.* fast cooling). The sample was kept at 15 °C for 15 minutes isothermally. For the pilot scale sample, a small amount of sample was placed between a glass slip and a coverslip and then placed under the microscope at 15 °C. The acquisition was performed with a Leica Microscope Lens HCX PL Fluotar $20 \times /0.50$ utilizing the software Leica Application Suite (LAS) V4.3. At least ten different images were acquired each time.

2.7 Phase contrast microscopy

For phase contrast microscopy, a Leica DM2500 LED microscope (Wetzlar, Germany), equipped with a Leica K3C colour camera, was utilized. Samples were prepared as explained in Section 2.6. The acquisition was performed with an HC PL FLUOTAR $40 \times /0.80$ PH2 objective, utilizing the software Leica LAS X. At least five different images were acquired each time.

2.8 X-ray scattering experiments

X-ray scattering experiments were performed using a TRUSAXS instrument of the European Synchrotron Radiation Facility (ESRF), Grenoble, France (beamline ID02). The wavelength was 1.013 Å (12.230 keV).²⁶ The setup and data reduction are described in a great level of detail in Appendix 1 (ESI[†]).

2.9 Short spacings

WAXS profiles were utilized to identify polymorphisms. For this, a Python script was utilized to find the relevant peaks. Eqn (1) was used to convert from inverse scattering vector q (nm⁻¹) to real space spacings d (nm).

$$d = \frac{2\pi}{q} \tag{1}$$

Polymorphisms were identified following the AOCS Official method C_j 2–95. An α polymorph was identified as a single peak at d = 0.415 nm, a β' polymorph was identified as two main peaks at d = 0.38 nm and d = 0.42 nm, and a β polymorph was identified as a strong peak at around d = 0.46 nm combined with several other peaks. The details of the 3D cell structure in the β polymorph can be found in Appendix 2 (ESI[†]).

2.10 Long spacings

Similarly, a Python script was utilized to find the peak positions in the SAXS area. After background subtraction, the peaks were fitted to a Gaussian curve, and the full width at half maximum (FWHM) was calculated for the three available peak reflections (001, 002 and 003). In each case, the equivalent *d*-value was calculated (eqn (1)), and the *d*-spacing was obtained from the linear interpolation of all recorded positions (Appendix 3, ESI†).²⁷ An alternative method for calculating long spacings has been previously described by Acevedo (2018),¹⁷ where the *d*-spacing is calculated utilizing only the first small angle reflection (d_{001}). The differences between calculation methods were found to be negligible for the systems reported here.

2.11 CNP thickness and micro-strain determination

The average crystallite size (D_{Schr}) was calculated from the SAXS profiles. For this, three alternate methods were utilized. First, the well-known Scherrer equation was used (eqn (2)),

$$D_{\rm Schr} = \frac{2\pi K}{\rm FWHM_{001}} \tag{2}$$

where FWHM₀₀₁ is the full width at half maximum of the 001 SAXS peak (in the inverse length) and *K* is the Scherrer constant, here assumed to be 0.9. The reader is recommended to review the article published by Langford and Wilson (1978) for more information on the physical meaning of the *K* factor, as a deeper review is out of the scope of this article.²⁸

Alternatively, the first-order SAXS peak (001) was utilized to obtain crystallite thickness distribution, following the procedure described by Rondou *et al.*^{29–31} with some minor modifications (Appendix 4, ESI†). From the distribution, the corresponding volume weighted average crystallite size was also calculated.

Finally, a Williamson–Hall plot was utilized for determining the impact of microstrain (noted as ε) and crystallite size (noted as D_{WH}). For this, eqn (3) was utilized.

$$FWHM_{hkl}\cos(\theta_{hkl}) = \frac{K\lambda}{D_{WH}} + 4\varepsilon \cdot \sin(\theta_{hkl}).$$
(3)

Employing a linear interpolation between $\sin(\theta_{hkl})$ and FWHM_{*hkl*} $\cos(\theta_{hkl})$ of all recorded SAXS peaks (001, 002, 003), ε and *D* were calculated (Appendix 5, ESI[†]). Instrumental peak broadening was calculated, and the resulting differences were found to be negligible (Appendix 6, ESI[†]).

2.12 Area per chain determination

The area per chain can be calculated differently depending on the polymorphic form. Each polymorphic form has a different unit cell arrangement; consequently, the calculation method for the area per chain is different. Whenever a beta polymorphic form is identified, the area per chain can be calculated following the methods described by Tenorio-Garcia *et al.*³² (Appendix 2, ESI[†]).

Results

3.1 Microscopy results

Given the hierarchical nature of fats, a comprehensive imaging strategy is necessary to capture the diverse length scales encompassed. Here, we combine cryogenic scanning electron microscopy (cryo-SEM), transmission electron microscopy (TEM), polarized light microscopy (PLM) and phase contrast microscopy (PCM). To begin with, cryo-SEM and TEM were employed to visualize the lamellae structures (Fig. 2). From the microscopy results, it was possible to corroborate the lamellar stacking (white arrows) for a slow-cooled and fast-cooled 30% SSS dilution in OOO. Although measuring the lamellae through microscopy remains impossible at the given scale, the length scale seems to correspond to the expected order of magnitude (of around 4.5 nm – see 3.2 X-ray scattering results). As seen in Fig. 2C and D, upon stacking, the lamellae formed a secondary domain size, here called CNPs. The thickness of such a structure was measured *via* the TEM microscopies (n = 49), and the resulting kernel density estimators are presented in Fig. 2E. The median thickness was found to be around 31 nm for the 30% SSS cooled at 10 °C min⁻¹ to 15 °C.

Furthermore, cryo-SEM was used to elucidate the morphology of the CNPs and their aggregation behaviour. For this, two different magnification levels were used (Fig. 3). A higher magnification was utilized to visualize the smallest building blocks (Fig. 3A–D). These are visualized as platelets with relatively smooth surfaces and are assumed to be the previously reported CNPs. These results align, in terms of conformation, with the TEM microscopy results presented by Acevedo for SSS systems.² The primary units are, therefore, identified as platelet-like structures characterized by length, width and thickness. When comparing the results in terms of shape and homogeneity, three main conclusions can be made. First, pure systems, composed of SSS, display bigger platelets with more consistent shapes when compared to their commercial counterparts (FHRO systems). Second, shear crystallization modifies the general platelet configuration, as corners seem to be eroded by shear crystallization. Third, slow cooled systems seem to have a high size polydispersity, as CNPs formed vary broadly in size.

Interestingly, regardless of the size differences, all measured samples seemed to keep a quasi-constant aspect ratio (as presented in Appendix 7 (ESI[†]), through similar regression

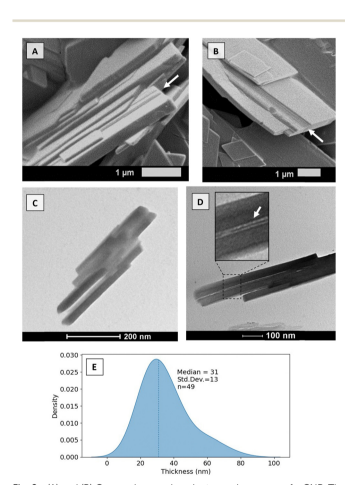


Fig. 2 (A) and (B) Cryogenic scanning electron microscopy of a CNP. The sample was slow-cooled 30% dilution of SSS in OOO (cooled at 1 °C min⁻¹ to 15 °C) and visualized after deoiling. (C) and (D) TEM images of CNPs. The inset in (D) presents a magnified view of the lamellar structures. The sample was a deoiled 30% dilution of SSS in OOO, cooled at 10 °C min⁻¹ to 15 °C. In all cases, white arrows indicate layering structures attributed to lamellar stacking. (E) Kernel density estimators of the thickness of CNPs measured from TEM images (*n* = 49). Median and standard deviation values are reported in nanometers.

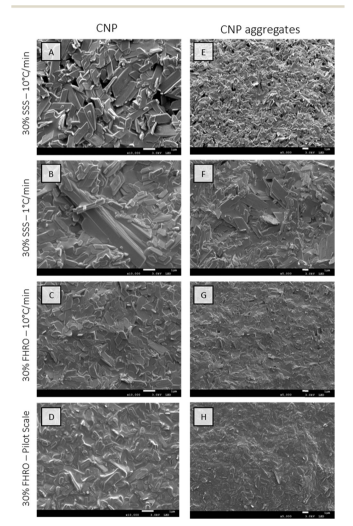


Fig. 3 Cryo-SEM images of all measured systems. (A) and (E) A statically crystallized 30% SSS in OOO cooled at 10 °C min⁻¹ to 15 °C. (B) and (F) A 30% SSS in OOO statically cooled at 1 °C min⁻¹ to 15 °C. (C) and (G) A statically produced 30% FHRO in HOSO crystallized at 10 °C min⁻¹ to 15 °C. Finally, (D) and (H) were produced on a pilot scale SSHE with the outlet temperature set to 15 °C (30% FHRO in HOSO).

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slopes). Lower magnifications were also utilized to visualize the population behaviour of the CNPs (Fig. 3E-H), as aggregates are probably destroyed during the deoiling procedure. In all cases, CNPs are represented in a disordered manner. The acquired images served as crucial inputs for quantifying their dimensions, specifically the width and length, as presented in the kernel density estimators in Fig. 4. The size of CNPs follows the order 30% FHRO-pilot scale < 30% FHRO – 10 °C min⁻¹ <30% SSS - 10 °C min⁻¹ < 30% SSS - 1 °C min⁻¹. The reader must interpret these results comparatively and within their length scale, rather than their absolute value, as measurement of cryo-SEM images can be subject to depth artifacts (due to the limited imaging depth). In addition, results are based on a limited number of platelets (n > 50).

Unsurprisingly, the size of the CNPs depends highly on the processing conditions, as fast-cooled systems appear to be smaller than the slow-cooled ones. This result is expected, as Acevedo et al. had previously reported that, for a 50% FHCO in HOSO, fast cooling led to smaller platelet dimensions than slow cooling.⁶ Later studies also confirmed the findings, as slow cooling created longer and wider CNPs at several different concentration levels.^{33,34} In addition, the application of shear resulted in smaller CNPs, with a homogeneous size distribution. These results also align with the work of Acevedo, where shear decreased the CNP length and width. The output is explained as shear enhances nucleation kinetics and mass and heat transfer in the melt.³³ Although the trend remained consistent, the overall size range of the SSS system seemed to differ from the previous literature, as bigger values were found in this study.² Differences were attributed to modifications in the deoiling

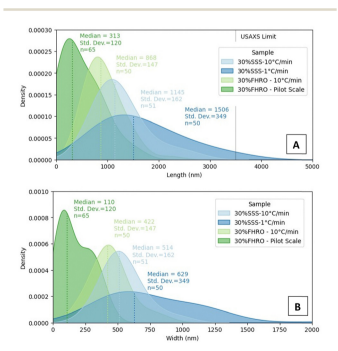
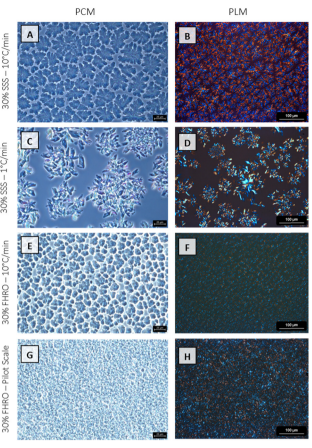


Fig. 4 Qualitative results were obtained by measuring the width and length of CNPs in the cryo-SEM images. In (A) kernel density estimators of the length of CNPs and in (B) kernel density estimators of the width of CNPs. Median and standard deviation values are reported in nanometers.



30% SSS - 1°C/min

Phase contrast microscopy (PCM) and polarized light microscopy Fia. 5 (PLM) images of the four different stearic acid samples.

procedure, as in this study, no shear or sonication was applied to the sample before measurement.

In the microscopic range, fat systems showed a clear tendency to form clusters. The floc formation was studied utilizing both PCM and PLM. Fig. 5 reveals pronounced differences in the microstructural features. The discrepancies between the four treatments become markedly noticeable when examined under a PCM, where the contrasting sizes and arrangements of the microstructures are more vividly apparent. FHRO formed the smallest flocs. Specifically, the pilot scale sample depicts the smaller units, followed by the fast cooled treatment. In contrast, 30% SSS forms expansive spherulites, and the slow cooled sample is characterized by larger and more open structures.

3.2 X-ray scattering results

3.2.1 WAXS. The WAXS measurements encompassed a qrange between 0.0018 and 0.2453 nm⁻¹. Fig. 6A presents the WAXS results for the four measured systems, with Table 1 documenting all identified spacings. Across all samples, three principal Bragg spacings are consistently reported, with a prominent peak at around d = 0.46 nm, indicative of a triclinic (β) arrangement. This result is somewhat unsurprising, as it is the most commonly attributed polymorphic form for saturated monoacid triglycerides. In addition, the result aligns with similar

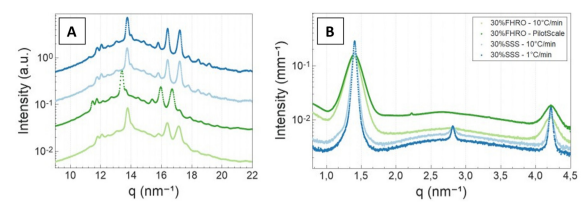


Fig. 6 Wide angle X-ray scattering (A) and small angle X-ray scattering (B) results. A vertical shift (factor of 5) was performed in (A) for better visualization.

Table 1	Wide angle V rea	v conttoring rocul	a an a a man a sain	a chart chaoinge	idontified n	aluma araba anadar	an nar chain values
Table 1	vvide andle A-ra	v scattering resul	S. ENCOMDASSI	a short spacings	. idenunea p	olvmorphs and an	ea per chain values

Sample	30% SSS				30% FHRO						
Cooling rate	10 °C min ⁻¹		1 °C min ⁻¹		10 °C min ⁻¹		Pilot scale				
Short spacing	$q (\mathrm{nm}^{-1})$	<i>d</i> (nm)	$q (\mathrm{nm}^{-1})$	<i>d</i> (nm)	$q (\mathrm{nm}^{-1})$	<i>d</i> (nm)	$q (\mathrm{nm}^{-1})$	<i>d</i> (nm)			
d_{10}	17.20	0.37	17.20	0.37	17.14	0.37	16.70	0.38			
d_{-11}	16.41	0.38	16.41	0.38	16.39	0.38	15.96	0.39			
d_{01}	13.75	0.46	13.76	0.46	13.78	0.46	13.40	0.47			
Identified polymorph	β		β		β		β				
Area per chain (nm ²)	0.1876		0.1875		0.1877		0.1981				

findings reported by other authors for SSS systems.^{35–39} Table 1 also presents the calculated area per chain, which was calculated following eqn (A2) (ESI \dagger).³² This parameter gives a quantitative indication of the area occupied by each fatty acid chain. Statically crystallized results are within the same order of magnitude, differing by less than 0.002 nm² between treatments, indicating that all tested systems have extremely similar sub-cell units. Interestingly, the pilot scale sample seems to deviate from the others, as the area per chain increased, suggesting the formation of a less dense triclinic packing.

3.2.2 SAXS. In the SAXS region (between 0.0351 and 4.5055 nm^{-1}), it is possible to identify the formation of lamellar phases (Fig. 6B).⁴⁰ All samples could be classified as having the $2L(\beta)$ configuration. Pure samples (SSS) displayed peaks at slightly higher q-values than their commercial counterparts (FHRO), suggesting slightly smaller lamellar sizes (Table 2). This result suggests that the presence of additional triglycerides led to less compact lamellae (4.48 nm vs. 4.47 nm). In addition, the shear crystallized sample exhibited the longest longitudinal spacing, suggesting a less compact lamellar domain after crystallization in the SSHE. This result seems to deviate from the findings of Mazzanti et al.,⁴¹ which suggest that an increase in the shear rate led to smaller d-spacings and higher lamellar thickness. Differences here are probably related to the differences in the experimental setup, as Mazzanti's research reports domain sizes in situ, during crystallization. On the contrary, in this article, one month of storage was allowed before measurement, allowing for domain relaxation (as confirmed by the low levels of crystalline microstrain).

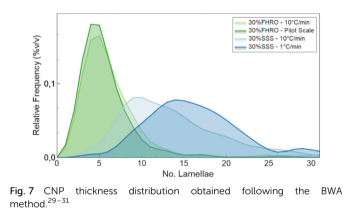
Additionally, the slow-cooled stearic acid sample was characterized by a thinner peak (1 $^{\circ}$ C min⁻¹) and higher intensity than the fast-cooled one (10 $^{\circ}$ C min⁻¹). This result is indicative of larger crystallite sizes as FWHM is inversely proportional to the crystallite magnitude (eqn (2)). Employing the Scherrer equation, the average crystallite size for all triglyceride samples was calculated (Table 2). Notably, the slow-cooled 30% SSS sample exhibited the largest crystallite thickness at 151.1 nm, followed by the fast-cooled 30% SSS sample at 92.8 nm. The FHRO samples followed in close proximity, measuring 39.02 nm when statically cooled at 10 $^{\circ}$ C min⁻¹ and 36.36 nm when crystallized in a SSHE. The corresponding CNP thickness distributions are available in Fig. 7. In the case of the FHRO, some compositional broadening can be expected, as the recorded profile can be the summation of various closely distributed *d*-spacings.

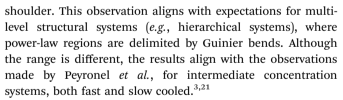
A Williamson–Hall plot was employed to discern the crystalline microstrain, revealing significantly lower levels of strain when compared to the previous literature on fat systems (our results always fall below 1×10^{-3}) (Table 2).⁴² Thus, both the Scherrer equation and the W–H plot yielded similar crystallite size results. The average crystallite size for all 30% fat dispersions seemed to follow the sequence 30% SSS – 1 °C min⁻¹ > 30% SSS – 10 °C min⁻¹ > 30% FHRO – 10 °C min⁻¹ > 30% FHRO – pilot scale.

3.2.3 USAXS. The USAXS data covered a *q*-range between 0.0018 and 0.2453 nm^{-1} . In Fig. 8, the USAXS results of the same four samples are presented. All curves are characterized by at least two discernible slopes interconnected by a Guinier

Sample Cooling rate		30%	SSS		30% FHRO											
		10 °C min ⁻¹			1 °C min ⁻¹			<u>10 °C min⁻¹</u> Pil			Pilot	ot scale				
Miller index	$q (nm^{-})$	-1)	d (nm ⁻¹)	$\stackrel{\rm FWHM}{(nm^{-1})}$	<i>q</i> (nm ⁻	-1)	d (nm ⁻¹)	$\stackrel{FWHM}{(nm^{-1})}$	q (nm	ı ^{−1})	d (nm ⁻¹)	FWH (nm ⁻		$q \pmod{(nm^{-1})}$	d (nm ⁻¹)	$\stackrel{FWHM}{(nm^{-1})}$
001 002 003	1.40 2.81 4.22		4.48 2.23 1.49	0.06 0.08 0.06	1.39 2.81 4.22		4.47 2.23 1.49	0.04 0.04 0.04	1.39 2.78 4.21	8	4.52 2.26 1.49	0.14 0.18 0.15		1.40 2.69 4.22	4.49 2.34 1.49	0.16 0.25 0.15
Domain size (nm)	4.47				4.47				4.48	3				4.51		
Identified stacking	$2L(\beta)$				$2L(\beta)$				2L(/	3)				$2L(\beta)$		
CNP thickness (nm) – Scherrer	92.8				151.1	-			39.0	02				36.36		
CNP thickness (nm) – BWA	59.8				71.7				28.7	8				25.9		
CNP thickness (nm) – WH plot	86.3				152.5	;			36.7	5				28.93		
Microstrain (%) – WH plot	4.90	imes 10	$^{-4}$		5.60	× 10	$)^{-4}$		1.00) × 10) ⁻³			2.30×1	0^{-3}	

Table 2 Small angle X-ray scattering results for statically crystallized 30% dilutions of SSS in OOO and 30% dilutions of FHRO in HOSO. Results include estimated peak positions and FWHM, identified lamellar stacking, domain sizes (Scherrer, BWA and WH) and microstrain estimations





In the high *q*-region, all scattering curves decayed in intensity with a power-law close to q^{-4} . Local slopes indicate characteristics of the surface roughness/compactness of a specific structural level.²¹ A slope of 4 suggests that the CNPs formed have well-defined interfaces and are compact objects with sharp interfaces.²¹ In the context of fat systems, previous interpretations are linked to surface roughness, where a slope of 4 is interpreted as a smooth surface.⁴³

As the *q*-value decreases ($q < 0.05 \text{ nm}^{-1}$), a notable feature in the curves emerges in the form of a distinct Guinier bend. These knees exhibit variability across the different samples, providing a clear indication of differences in conformation within the examined systems. This result suggests that the mesoscale is modified both by TAG purity and processing conditions. The same data are presented in the inset of Fig. 8

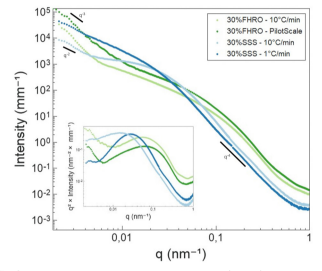


Fig. 8 Ultra small angle X-ray scattering results (*I* vs. *q*). Results were merged at low *q* values with the SAXS data, leading to some gaps in the data. The inset depicts a Kratky representation $(q^2 l vs. q)$ of the same data.

through a Kratky representation $(q^2I(q) vs. q)$, which allows a better visualization of the bend. In fact, in this representation, the Guinier inflection appears as a peak, which can assist in interpreting the CNP conformation. The location of the peak provides information on the size of the cross-section. From Fig. 8, we can infer that the CNPs of the two FHRO samples have smaller cross-sections, followed by the 30% SSS – 10 °C min⁻¹ and ultimately the 30% SSS – 1 °C min⁻¹.

Fig. 8 also reveals that, for the FHRO samples and the fast cooled 30% SSS sample, a secondary slope is visible in the lowest *q*-range. This additional slope suggests a secondary aggregation level and the difference between slopes in this region suggests changes in the aggregation behaviour between samples. This range will be interpreted through fractal dimensionality. For the slow cooled 30% SSS sample, this feature is

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not fully resolved in the available *q*-range, suggesting that the crystallites aggregate at higher dimensions than 3.5 μ m (the detection limit of USAXS). This result aligns with the findings presented in the microscopy section, as some of the CNPs are found to be out of the 3.5 μ m range.

As highlighted in the Introduction section, exploring the USAXS range in triglycerides is particularly intriguing due to its capacity to disclose essential details about CNPs and CNP aggregates. The scattering profile in this context exhibits numerous degrees of freedom (CNP shape, CNP size, CNP aggregation, fractality, polydispersity, *etc.*). However, a substantial amount of structural information is lost upon transforming from the real space (*d*) into the reciprocal space (*q*). As the USAXS range does not present sharp characteristic feats (*e.g.*, peaks), it is necessary to utilize models to retrieve information from this area.

Table 3 presents the results for the unified fit and Guinier-Prod models. SasView $5.0.4^{44}$ was used as a fitting software. The corresponding fits are available in Fig. 9. Further information on the models can be found in Appendix 8 (ESI[†]). In the case of the 30% FHRO – 10 °C min⁻¹, two different unified fits were performed, one assuming 2 levels of aggregation and one assuming 3 levels.

From the unified fit model, we can recognize two parameters associated with the CNP size, namely $R_{g_{CNP}}$ and R_{CNP} . R_{CNP} is calculated based on $R_{g_{CNP}}$, following the assumption of monodisperse spherical scatterers (eqn (A7), ESI†). Table 3 confirms that the smaller CNPs are obtained for the 30% FHRO – pilot scale, while the biggest CNPs are reported for the slow cooled 30% SSS dispersion. Interestingly, whenever 2 levels were used for the fit, the 30% SSS crystallized at 10 $^{\circ}$ C min⁻¹ exhibits smaller CNPs than the fast-cooled 30% FHRO. These results are not in line with the order suggested in the microscopy findings, as well as the initial Kratky plot interpretation. Conversely, whenever 3 levels are assumed, the order of magnitude is respected, as the fast cooled 30% FHRO sample has smaller CNPs than its pure counterpart.

Regarding fractal dimensionality, the factor P was utilized, following the interpretation suggested by Pink et al.¹⁸ Peyronel often reports a fixed slope of 4 (P_{CNP}), which is interpreted as a smooth-surfaced scatterer.³ As seen in Table 3, P_{CNP} , associated with the surface of the CNPs, was close to 4 for most samples (suggesting sharp interfaces and smooth surfaces). These findings align with those of Peyronel for similar systems,⁴³ as well as the neat surfaces visualized through a cryo-SEM (see Section 3.1 – Microscopy results). However, the result obtained for the 30% FHRO - 10 °C min⁻¹ whenever two levels were assumed did not seem to follow the norm, as the fit yielded a P_{CNP} value of 2.26. This slope would be traditionally interpreted as mass fractal, indicative of density fluctuations²¹ and a loose conformation. This result is not in accordance with the cryo-SEM images (Fig. 3) nor portrays the expected surface configuration for highly homogeneous FHRO CNPs. In contrast, whenever 3 levels were utilized for the fit, the surface roughness of the CNPs seems to have an appropriate magnitude of 3.4, indicative of a surface fractal.

At this point, a critical discussion concerning the selection of levels in the unified fit is imperative. In fact, based on the microscopy results, there are no indications of the existence of an intermediate aggregation level at a nanometre scale (as suggested by the R_{INT} of 154 nm in the 3 level fit), and therefore,

Table 3 Unified fit and Guinier–Porod model results. The reported q-ranges are those used to initialize the fit. Once the fit is performed in the whole range, the fit ranges between the q-max of the CNPs to the q-min of the CNP aggregates

Sample				30% SSS		30% FHR	0	
Cooling rate			Levels	10 °C min ⁻¹	1 °C min ⁻¹	10 °C mir	Pilot scale	
Unified fit	Scattering unit	Parameter		2	2	2 3		2
	CNP	$R_{ m g_{CNP}}$	nm	77.06	134.17	100.16	42.24	33.31
		R _{CNP}	nm	99.48	173.21	129.31	54.53	43
		$P_{\rm CNP}$	_	4.2	3.83	2.26	3.44	3.9
		$D_{\mathrm{f}_{\mathrm{CNP}}}$	_	4.2	2.17	2.26	2.56	2.1
		Fractal		Diffuse	Surface	Mass	Surface	Surface
		<i>q</i> -min		0.0054	0.007	0.02	0.05	0.01
		q-max		0.2	0.2	0.2	0.4	0.2
	Intermediate level	$R_{\rm g_{\rm INT}}$	nm	_	_	_	119.97	_
		R _{INT}	nm	_	_	_	154.88	_
		P _{INT}	_	_	_	_	1.11	_
		$D_{f_{INT}}$	_	_	_	_	154.88	_
		Fractal	_	_	_	_	Mass	_
		<i>q</i> -min	_	_	_	—	0.008	_
		q-max	—	—	—	—	0.025	_
	CNP aggregates	P_{Agg}	_	4.12	2.19	3.9	3.91	2.85
		$D_{f_{Agg}}$	_	>4	2.19	2.1	2.09	2.85
		Fractal		Diffuse	Mass	Surface	Surface	Mass
		<i>q</i> -min		0.0021	0.0021	0.0021	0.0021	0.0028
		q-max		0.004	0.006	0.004	0.004	0.004
Guinier–Porod	CNP	S		0.62	1.62	1.44		1.6

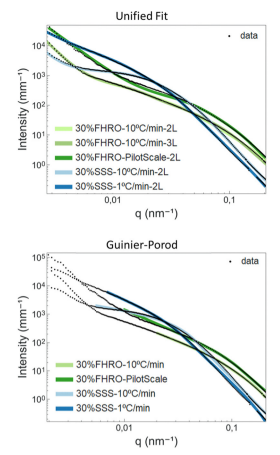


Fig. 9 Unified fit and Guinier–Porod curve fitting results. In black, the fitted data and in coloured lines, the resulting fits. For the unified fit model, the number of levels are specified in the legend (2L = 2 levels, 3L = 3 levels).

phenomenologically speaking, a two level model should be the most appropriate. However, whenever the transition between two levels covers a broad *q*-range, the model requires the addition of the "artificial" level in order to deliver phenomenologically sound outputs (*e.g.* appropriate R_{CNP} and P_{CNP}).

In addition, a noteworthy aspect of concern in the unified fit relates to the reported practice of adjusting the number of levels in the model based on the fit quality.⁴⁵ Although seemingly pragmatic for enhancing fit performance, this practice raises questions about the physical validity of the added levels. In other words, by adding an additional aggregation level, there is a risk that, in pursuit of achieving an optimal fit, the inclusion of additional aggregation levels might be driven more by the exigencies of the fitting process rather than by a grounded physical justification of the existence of a specific aggregation level. When interpreting scattering data, it is essential to have alternative techniques to support the hypothesis of additional aggregation levels (*e.g.*, TAG woods).

3.3 Proposed model

Based on the previous discussion, it seems imperative to generate a shape-dependent model that can better represent the morphology of CNPs and their aggregation behaviour. Eqn (4) presents the simplest model to describe scattering data.

$$I(q) = K \cdot \langle P(q) \rangle \cdot \langle S(q) \rangle \tag{4}$$

where *K* is the scaling parameter, $\langle P(q) \rangle$ is the orientation- and size-averaged form factor of the particles, related to the shape of scattering particles and $\langle S(q) \rangle$ is the structure factor, which accounts for the interactions between particles.

3.3.1 Scale factor *K*. The scale factor *K* is the part of the model that accounts for the concentration of particles and their scattering density. In the proposed model, the *K* term can be described using eqn (5).

$$K = V_{\rm P} \cdot \phi_{V} \cdot (\Delta \rho)^2 \tag{5}$$

where $V_{\rm P}$ is the mean volume of the scattering particles, ϕ_V is the volume fraction of the CNP particles (which is assumed to be proportional to the SFC), and $\Delta\rho$ is the difference between scattering length density of the CNPs (solid TAGs) and the solvent (mainly oleic TAGs and any dissolved fractions).

The chemical formula and mass density were used to calculate the scattering length density (ρ) of SSS and OOO utilizing the SasView SLD Calculator.⁴⁴ Given that that the main triglyceride in all cases is SSS and that all profiles analysed are in the β form, the density of SSS in the β phase was used for calculating the ρ of the solid TAGs (mass density = 1.03 g cm⁻³).¹⁰ The solvent was assumed to be mostly composed of triolein, and therefore, the scattering length density of triolein was utilized (mass density = 0.915 g cm⁻³).⁴⁶ The resulting ρ was found to be 9.82 × 10⁻⁶ Å⁻² for SSS and 8.67 × 10⁻⁶ Å⁻² for OOO. The model here proposed accounts only for the solid contributions. In fact, as experimentally observed, the scattering from the liquid oil fraction remains always below the scattering of the solid fat (Appendix 9, ESI[†]).

3.3.2 Form factor $\langle P(q) \rangle$. The orientation- and sizeaveraged form factor $\langle P(q) \rangle$ describes the geometrical arrangement of the particles. In other words, it is the part of the model that accounts for the shape of the primary scattering unit (CNPs). To find an adequate form factor it is necessary to have a reasonable assumption of the shape of the primary scattering unit. In the proposed model, we assume that CNPs can be represented as parallelepipeds based on (1) in-house microscopy techniques and (2) the literature. As previously explained in Section 3.1 – Microscopy results, CNPs are visualized as platelet-like structures characterized by length, width and thickness. These findings were also compared with the literature, where CNPs have been consistently approximated to platelet particles with three distinct dimensions (length, width and thickness).^{2,6,8,13,17,33,34}

The orientation- and size-averaged form factor of the particles was calculated following a parallelepiped model^{47,48} and implemented in SasView.⁴⁴ As a result, in the USAXS scattering data, the shoulder appearing between 0.005 and 0.3 nm⁻¹ is attributed to the cross-section of the CNPs (Fig. 10). Differently sized CNPs result then in shoulders placed in different *q*-ranges. As the peak in the Kratky plot is shifted to lower *q*, an increase in the cross-section of CNPs is suggested.

Soft Matter

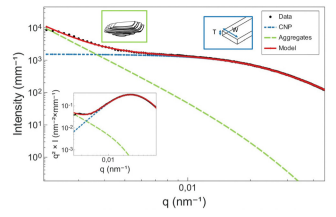


Fig. 10 Graphical representation of the proposed model. At low *q*, the aggregates dominate the scattering, modelled through a structurally limited power law. The Guinier bend is attributed to the cross-section of the CNPS, here modelled through a parallelepiped form factor. The results are also satisfactory in a Kratky representation (inset).

In order to better understand what information in the model is relevant, a simple sensitivity analysis was conducted (see Appendix 10, ESI†). Here, variations in the parallelepiped dimensions, including thickness, width and length, were systematically studied, and the outcome model was plotted. The intention behind this analysis was to identify which factors modify features in the intermediate bend. In principle, only the factors that have an impact on the knee should be taken into account. Notably, variations in thickness and width had a discernible impact on the modelling of the Guinier bend. Conversely, changes in length predominantly affected the low q part of the curve. This region, as mentioned before, is heavily influenced by the scattering of CNP aggregates, and therefore, it is not advised to conclude in terms of length. In this study, we limit the interpretation of the model to the cross-section of the CNPs (thickness and width).

3.3.3 Structure factor $\langle S(q) \rangle$. Finally, the structure factor accounts for the spatial correlation between scattering particles.¹⁹ In fact, the parallelepiped model is only valid over a limited *q*-range, as the Bragg reflections characteristic of the lamellar stacking dominate the scattering at high *q*, while the scattering of CNP aggregates starts to dominate at low *q* values ($q < 0.0055 \text{ nm}^{-1}$). Given that there is no strong correlation peak in the intermediate *q*-region of the USAXS range, $\langle S(q) \rangle$ is assumed to be equal to 1 whenever aggregates are not present (CNPs in solution).

In the low q-region, whenever a protruding slope is visible (attributed to the aggregation of CNP domains), the structure factor can be modelled using a power-law dependence, assuming the formation of fractal aggregates. The structure factor is defined in eqn (6).

$$S(q) = \left[1 + \frac{B}{q^{\mathrm{P}}}\right] \tag{6}$$

3.3.4 Suggested model. The proposed model follows eqn (7). The multiplying factor between the aggregates and the parallelepiped function allows for a structurally limited power law, which essentially truncates the power law at high q. A visual representation of the model is presented in Fig. 10.

$$I(q) = \phi_V \cdot V_{\mathbf{P}} \cdot (\Delta \rho)^2 \cdot \langle P_{\mathbf{P}\mathbf{P}} \rangle \left[1 + \frac{B}{q^{\mathbf{P}}} \right]$$
(7)

Unless polydispersity is included in the model, the model can deviate from the data in high q ranges (Appendix 11, ESI†). In fact, as the polydispersity on the cross-section of the system increases, the smoother the decaying Porod slope at high q. The assumption of polydispersity was considered reasonable, as observed in some cryo-SEM images (Appendix 11, ESI†). These findings are also supported by previous quantification efforts on CNPs, both in this publication (Fig. 4) and the multiple CNP histograms presented in the previous literature.^{2,13,17,33,34} For computational reasons, a fixed polydispersity value was utilized (here, arbitrarily set to 0.3).

It remains relevant to disclose that the model described here was inspired by similar models existing in the literature. Patil et al.¹⁹ used a comparable approximation to extract structural parameters from SAXS data on organic thin films (poly(3hexylthiopene)) for solar cell applications. Their scattering results, similar to those reported here for fats, include three areas, including (1) a Braggs reflection in a high q-region, (2) a broad shoulder in intermediate *q*-ranges and (3) a power law at low q. Similar to triglycerides, the Braggs reflection was linked to lamellar stacking and the power law was attributed to bigger amorphous domains. The intermediate and low-q domains were modelled with a combination of a cylinder form factor and a Debye-Bueche structure-function. There, the shoulder was attributed to the cross-section of the polydisperse cylinders (primary scattering units), whereas the signal from the amorphous network was also parametrized utilizing fractal dimensionality.¹⁹ In addition, a secondary inspirational model was presented by Newbloom et al. In their work, also linked to organic photovoltaics, poly(3hexylthiopene)s,⁴⁷ and poly(3-alkylthophene)s,⁴⁹ scattering results were modelled by combining a parallelepiped model (to model the polymer rectangular cross-section) and a power law to determine the fractal dimension of bigger aggregates. Again,

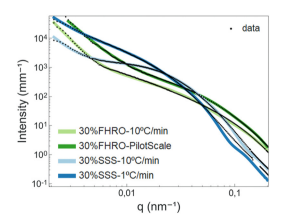


Fig. 11 Proposed model curve fitting results. In black, the fitted data and in coloured lines, the resulting fits.

 Table 4
 Proposed model fit results following eqn (7). Here, P is the power law exponent and D_f is the corresponding fractal dimension

Sample			30% SSS		30% FHRO			
Cooling rate			10 °C min ⁻¹	$1 ^{\circ}\mathrm{C} \mathrm{min}^{-1}$	10 °C min ⁻¹	Pilot scale		
CNP	Thickness	nm	41	57	12	15		
	Width	nm	96	192	68	47		
	Polydispersity	_	0.3	0.3	0.3	0.3		
	Cross section	nm^2	3892	10938	836	718		
	No. lamellae	_	10	13	3	4		
	T/W ratio	—	0.4	0.3	0.2	0.3		
CNP aggregates	Р		3.3	2.72	3.8	2.42		
	D_{f}	_	2.7	2.7	2.2	2.4		
	Fractal	_	Surface	Mass	Surface	Mass		

the scattering profiles were found to be very similar to those of triglyceride systems.

3.3.5 Model results. The resulting fits are available in Fig. 11. The numerical results are presented Table 4. In the upper section of the table, parameters associated with CNPs are presented, while in the lower part, information on the aggregates is provided. Overall, the order of magnitude of the cross-section follows the order: 30% FHRO-pilot scale < 30% FHRO – $10 \,^{\circ}\text{C} \text{min}^{-1} < 30\%$ SSS – $10 \,^{\circ}\text{C} \text{min}^{-1} < 30\%$ SSS – $1 \,^{\circ}\text{C} \text{min}^{-1}$. This outcome aligns with the initial Kratky plot interpretation and is in good agreement with the order reported in the microscopy results (Fig. 4).

In terms of CNP thickness, when comparing the model results to the TEM data (for the 30% SSS – 10 °C min⁻¹) (Fig. 2), we can conclude that the results are in sensible agreement, as TEM microscopy suggest a thickness of 31 nm (equivalent to around 7 lamellae), while the model output was 41 nm (about 10 lamellae). This deviation is considered acceptable, as the overall order of magnitude is well represented by the results. Moreover, the output is in agreement with the order obtained in the SAXS results. Particularly interesting is the alignment between the crystallite thickness distribution obtained *via* de the BWA approximation (see Fig. 7) and the model results. In fact, the peak of each of the distributions is notably close to the obtained thickness, suggesting a good method cross-validation.

At this point, a discussion concerning the limitations of the Scherrer equation is necessary. Whenever the crystallite size exceeds a certain threshold, typically within the range of 100–200 nm, peaks become too narrow, and inevitably, the Scherrer equation is subject to big errors.¹⁷ The sizes reported here for the 30% SSS – 1 °C min⁻¹ are within this range, and therefore, results are within the limits of the method. Moreover, distortions in crystal packing or instrumental broadening can further contribute to inaccuracies in size characterization utilizing the Scherrer equation. The model proposed herein bypasses the constraints of the Scherrer equation, offering an alternative method for quantifying CNP thickness in a different *q*-range.

As seen in Table 4, while the CNP thickness seems to be dominated by the purity of the triglyceride, the width is clearly influenced by the processing conditions. In fact, the slowcooled 30% SSS system exhibited wider platelets than the fast-cooled equivalent. This result confirms the generalized findings of Acevedo *et al.*, suggesting that slow-cooled samples form wider CNPs.^{33,34} In addition, it was also possible to confirm that shear crystallization reduced platelet width significantly,³³ although the same finding could not be corroborated for thickness.

While the general conclusions remain valid, there are some deviations in the absolute width values. In fact, modelled values appear to be smaller than those measured in the microscopy section. In this case, the USAXS results are deemed more reliable, as cryo-SEM results are bound to have diverse sources of errors. In fact, unlike TEM, cryo-SEM is better suited for surface characterization rather than cross-section measurements, and therefore depth artifacts need to be considered when performing the measurements. In addition, cryo-SEM measurements are limited to a subset of platelets, which restricts the analysis to a small subgroup of nanostructured elements ($n \approx 50$). Finally, cryo-SEM results are subject to extensive sample handling, including deoiling, which can have important consequences in sample dissolution and re-crystallization in the solvent.

In the lower section of Table 4, data concerning the aggregation behaviour are presented. The interpretation remains consistent with that provided in the unified fit section. For both rapidly cooled systems, 30% SSS and 30% FHRO, *P* lies between 3 and 4, indicating tight CNP packing and surface fractality. In contrast, slow-cooled 30% SSS and 30% FHRO pilot scale samples exhibited fractal dimensions ranging from 2.4 to 2.7, indicative of mass fractals, suggesting looser conformations.

4. Conclusions

In this study, for the first time, a shape-dependent model to interpret triglyceride USAXS data was proposed. The model, covering two orders of magnitude, describes the CNPs as polydisperse parallelepipeds, while describing the crystallites (CNP aggregates) through fractal dimensionality. The fit yields information on the cross-section of the CNPs, providing an alternative to the Scherrer equation for thickness calculation. In addition, the proposed model bypasses some of the limitations associated with the unified fit model, as it accounts for the morphological features of the primary scattering unit and removes the need to add an artificial level whenever the transition between two levels covers a broad *q*-range. To validate the proposed model, four stearic acid based samples crystallized through different protocols were characterized. The average CNP size for all 30% fat dispersions seemed to follow the sequence (from smaller to bigger): 30% FHRO – pilot scale, 30% FHRO – 10 °C min⁻¹, 30% SSS – 10 °C min⁻¹, and finally 30% SSS – 1 °C min⁻¹. These results were consistent across different methods (X-ray scattering and microscopy) and models (unified fit and proposed models) utilized. We anticipate that the implementation of this model will significantly enhance our capacity to characterize CNPs, thereby advancing our understanding of the mesoscale dynamics dominating triglyceride systems.

Author contributions

Conceptualization: I. A. P., T. R., K. D., F. V. B.; data curation: I. A. P., W. C.; formal analysis: I. A. P., F. V. B.; funding acquisition: I. A. P., K. D., F. V. B.; investigation: I. A. P., F. D. W., T. R., W. C.; methodology: I. A. P., F. D. W., T. R., W. C., I. P., F. V. B.; project administration: K. D., F. V. B.; resources: K. D., F. V. B.; software: I. A. P., F. D. W.; supervision: T. R., K. D., F. V. B.; visualization: I. A. P., I. P.; writing – original draft: I. A. P.; and writing – reviewing and editing: F. D. W., T. R., W. C., I. P., K. D., F. V. B.

Data availability statement

Following the ESRF data policy, the raw XRS data are made openly accessible to the public *via* 10.15151/ESRF-DC-1543611960. Any additional data that substantiate the outcomes of this research are publicly and openly available in Zenodo at 10.5281/zenodo.10693917.

Conflicts of interest

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