PECULIAR CRYSTALLISATION BEHAVIOUR OF PALM OIL DURING DRY FRACTIONATION

CREMER, G1; DANTHINE, S1; BLECKER, C1 and GIBON, V2*

ABSTRACT

Multi-step dry fractionation of palm oil generates fractions with specific physicochemical properties suitable for many food formulations. The present study addresses the pilot scale production of palm olein iodine value 56 starting from palm oil. The aim was to investigate the influence of the tri-saturated triacylglycerol content (StStSt) in palm oil on its crystallisation behaviour. Four StStSt contents were investigated: 7.0%, 7.6%, 8.2% and 9.8%, and the palm oil crystallisation was examined during a period of 240 min. Oil temperatures were recorded, and crystallisation kinetics were monitored by p-NMR; crystal morphology and polymorphic forms were characterised by optical microscopy and powder X-ray diffraction. One of the compositions (StStSt: 7.6%) was crystallising faster. All the crystallised matrices were filtered at 25°C with a membrane press filter; the olein yields, iodine values, cloud points and triacylglycerol compositions were analysed. The olein from the matrix crystallising faster showed a yield, an iodine value and a cloud point higher than expected. A more detailed HPLC analysis of this olein indicated unusual enrichment in OPP and depletion in OPO, which led to the conclusion of an unusual crystallisation behaviour at this specific composition.

Keywords: dry fractionation, palm oil, palm olein, peculiar crystallisation behaviour, tri-saturated triacylglycerols.

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INTRODUCTION

Edible oils are made up of a wide variety of triacylglycerols (TAGs) with diverse fatty acid (FA) compositions, leading to complex physicochemical behaviour. TAGs can be classified according to their saturation: The tri-saturated (StStSt), the di-saturated (StU), the di-unsaturated (StU) and the tri-unsaturated (UUU). In palm oil (PO), tripalmitin (PPP), oleo-dipalmitin (P2O), palmitodiolein (PO2) and trilinolenin (OOO) are the most abundant of each class. PO crystallisation behaviour has important implications in the manufacture of many edible fatty products like margarine, spreads or shortenings.

Crystallisation operates in two steps: Nucleation followed by crystal growth. The driving force for nucleation is determined by the degree of supersaturation of the liquid. PO crystallisation has been extensively reviewed by Omar et al. (2015). The authors described the effect of chemical composition, crystallisation conditions and the presence of additives. They also focussed on crystallisation kinetics, thermal properties, polymorphism, and recrystallisation aspects. Braipson-Danthine and Gibon (2007) reported a relationship between melting properties, polymorphism and TAG composition of PO and fractions. De Oliveira et al. (2014), Saberi et al. (2011) and Siew and Ng (1999) studied the effect of diacylglycerol (DAG) content on PO nucleation and crystal growth. Basso et al. (2010) and Verstringe et al. (2013; 2014) characterised the effects of PPP and monoglycerides (MAG) on crystal formation. Fredrick et al. (2008) studied the influence of MAG on the crystallisation behaviour of PO. Saadi et al. (2012) reported that 1,2-dipalmitoyl-3-oleoyl glycerol/1,3-dipalmitoyl-2-oleoyl glycerol (PPO/POP) were the major components of the primary nucleus developed during the crystallisation of PO and palm stearin (PS) blend systems. The influence of sorbitan esters and soya lecithin in PO blends were studied by Kwamura (1980) and Miskandar et al. (2006; 2007). The effects of polyglycerol behenic acid esters on PO crystallisation were described by Sakamoto et al. (2003) and of polyglycerol ester additives by Saw et al. (2017), using focused beam reflectance (FBRM)
and differential scanning calorimetry (DSC). The influence of high-intensity ultrasound (HIU) on PO crystallisation was investigated by Fangfang et al. (2013) and Yubin et al. (2015). Hubbes et al. (2018) studied the crystallisation of PO-PS blends to obtain fats with higher PPP contents and PPP/OOO ratios; they showed that the global rate constant of the enriched PO increased with the addition of PPP. PO is a good candidate for dry fractionation, which was extensively reviewed by Deffense (1985), Gibon and Tirtiaux (2002), Gibon (2006; 2012), Gibon et al. (2009; 2020), Kellens et al. (2007), Mei Huey et al. (2015), Tong et al. (2021) and Zaliha et al. (2004).

Multi-step dry fractionation generates a large number of solid and liquid fractions with specific physico-chemical properties; it shows three steps, and three different routes that can be followed (Figure 1); the solid route (for PPP enrichment), the hard palm mid fraction (HPMF) route (for P₂O enrichment) and the liquid route (for PO₂/OOO enrichments). In dry fractionation, crystallisation is followed by crystal separation typically using membrane press filters. Chong et al. (2014a; 2014b; 2015) described technology for higher olein yield using a special cooling programme and additional steps mid-way. Defense (2009) proposed an ultrasound seeding technique for improved crystallisation. Kuriyama et al. (2011) showed that the addition of polyglycerol ester additives enhances the olein yield and alters the fractions characteristics. Tong et al. (2021) reviewed the fractionation conditions that affect the yield and quality of the oil produced and published updates on the influence of seeding agents (DAG, hard fats, etc.) used in fractionation.

In industrial practice, a quantity of StStSt is classically added to palm olein, to ensure an efficient and reproducible crystal initiation for superolein production (Calliauw et al., 2007a; 2007b; 2010). On the contrary, the naturally existing StStSt content of PO permits crystal initiation. In this work, the influence of the StStSt content on PO crystallisation is explored. PO was blended with PS and matrices containing variable contents of StStSt were considered. Crystallisation kinetics were established, operation yields were quantified and the quality of the olein fractions was analysed. The aim of this paper is to highlight and explain a particular crystallisation behaviour observed unexpectedly while studying the effect of StStSt on the crystallisation of PO during dry fractionation.

**MATERIALS AND METHODS**

**Materials**

PO was supplied by Fuji Oil Europe (Ghent, Belgium); refined PS was provided by Desmet Ballestra Group (Zaventem, Belgium). Increasing amounts of PS were added to PO in order to reach StStSt TAGs contents of 7.0%, 7.6%, 8.2% and 9.7% in matrices named M1, M2, M3 and M4 respectively (= internal seeding). To this end, PS was first placed in an oven at 80°C to ensure a complete melting before being mixed with PO. The appropriate quantity was weighed into a preheated glass beaker and then added as liquid in the melted PO in the crystalliser.

**Crystallisation**

Crystallisation was performed in a 30 kg capacity pilot-scale crystalliser having an exchange surface of approximately 15 m²/T; all the tests were carried out with a total oil quantity of 20 kg. The crystalliser was a cylindrical stainless-steel double-jacketed...
tank connected to an external programmable water bath for controlling the temperature. The agitation was conducted by a double-bladed rotor. The oil temperature was measured by a Pt100 probe which was dipped into the oil. Cooling programmes controlling water temperature, oil temperature, and Δ oil/water temperature were applied using the Citect SCADA RunTime application (Citect, Australia). The cooling programme (13 parameters) was first optimised for each system to ensure minimal oil temperature increase during the crystal growth. The oil melting temperature was 75°C, the time allocated for crystal initiation and growth (main crystallisation) was 240 min, and the final temperature was set at 25°C (Figure 2). The crystallisation tests were conducted in triplicate.

Vacuum and Membrane Press Filtrations

The separation of the crystals from the liquid was done by vacuum filtrations (Büchner) and by membrane press filtrations (lab press filter). Vacuum filtrations were performed at regular intervals (every 30 min) during the main crystallisation and press filtrations were only done at the final cooling temperature (25°C).

Vacuum filtrations. Approximately 100 mL of oil were sampled from the crystalliser and immediately filtered over a Büchner (55 mm diameter) covered with filter cloth (500 L min⁻¹dm⁻² at 196 Pa) and connected to a vacuum by means of an MZ 2C NT vacuum pump (Vacuubrand, Germany). The filtrations were stopped based on visual observation of cake dryness.

Membrane press filtrations. Press filtrations were performed with a single chamber lab-scale membrane press filter (Choquenet S.A.S., France) covered with the same filter cloth as for vacuum filtrations. The diameter of the chamber was 5.6 cm and the width was 20 mm. The filter was filled with a volume of approximately 200 mL of oil sampled in the crystalliser (at 25°C). Air pressure, which gradually increased by 0.5 bar min⁻¹ to 3 bar, was used for filling; it was then held at this value for 15 min. Afterwards, additional pressure was applied to squeeze the cake by gradually increasing the air pressure by 1 bar every 2 min to 6 bar. The squeezing pressure was held at this value for 10 min. The olein was collected in vessels and the stearin cake was recovered after opening the filter. The operation yield was derived from weighing the recovered fractions.

Monitoring of the Crystallisation by Pulsed Nuclear Magnetic Resonance (pNMR)

The kinetics of crystallisation was established by measuring the solid fat content (SFC) of the crystallising oil as a function of the time using p-NMR (Minispec-mq20, Germany). Daily calibrations were performed with three standards containing 0.0%, 31.1% and 74.8% of solids, respectively. NMR tubes were filled with crystal slurry directly from the crystalliser using a glass tube with a tight-fitting plunger (Danthine et al., 2003 and Danthine, 2012) and the SFC was immediately measured.

Microstructure by Optical Microscopy

For each sampling, crystal microstructure was observed by optical microscopy (OM) using a microscope coupled to a camera transmitting live images to a computer. The crystals were observed using an Eclipse E400 microscope (Nikon, Japan) equipped with a DS-Fi2 camera (Nikon, Japan).

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**Figure 2. Schematic cooling curve with fast cooling, crystal initiation, crystal growth and final cooling steps.**
The pictures were processed using NIS-elements 4.30 software (Nikon, Japan). A magnification of 20x was selected to analyse the pictures. One droplet of crystal slurry was randomly sampled from the NMR tube with a glass Pasteur pipette; it was then put on a microscope slide. A cover slip was carefully placed and centred over the drop of the sample to ensure homogeneity of the sample thickness. For each sampling, images of crystal microstructures were taken in triplicate.

**Polymorphism by Powder X-ray Diffraction (XRD)**

The polymorphic forms of the crystals were determined by XRD using a Bruker D8-Advance Diffractometer and a Lynx-eye detector (Bruker, Germany) (λ Cu= 1.5406, 40kV, 30mA). Calibration was performed using corundum, silver behenate, and tristearin standards. The X-ray diffraction patterns were analysed using DIFFRAC.EVA V4.2.2 software (Bruker, Germany). Crystals were placed on a sample holder directly after the vacuum filtration and immediately analysed. Small-angle X-ray diffraction patterns (long spacings) were investigated from 2θ = 1 to 10° with 452 steps and at 0.5 s per step. The wide-angle X-ray diffraction patterns (short spacings) were investigated from 2θ = 15 to 27° with 603 steps and at 0.5 s per step. The total duration of a run was approximately 7 min.

**Composition by Reversed-Phase High-performance Liquid Chromatography (RP-HPLC)**

The DAG content and the TAG composition were analysed by RP-HPLC based on the official AOCS Ce 5b-89 method (AOCS, 2017), which does not distinguish between positional isomers. Analyses were performed with a Waters e2695 HPLC system (Waters, USA), equipped with two stainless steel Nova-Pak C18 columns (4 μm, 3.9 × 150 mm). The mobile phase was an isocratic solvent (mixture of acetone and acetonitrile (62.5/37.5 v/v)) with a flow rate of 1.2 mL min⁻¹. The injection volume was 20 μL. The samples were dissolved in methanol/chloroform (1/1 v/v) and a differential refractometer was used for the detection. The assignment was confirmed by comparing retention times with those of some standards. Peaks were integrated with Empower Pro “Apex Track” algorithm (Waters, USA); peak areas below 4000 area counts (equivalent to approximately 0.04% of the total peak area) were not considered. The mean of two independent measurements is reported.

**Iodine Value (IV) by Titration**

The IV was determined by titration based on the official method AOCS Cd 1d-92 (Wijs method) (AOCS, 2017). The mean of two measurements is reported.

**Cloud Point (CP) by Differential Scanning Calorimetry (DSC)**

DSC analyses were carried out using a Q-2000 DSC (TA Instruments, USA) coupled with a refrigerated cooling system (TA Instruments, USA), using aluminium hermetic pans. Calibration was made with indium (melting point 156.6°C; ΔH = 28.7 J g⁻¹) and eicosane (melting point 36.8°C; ΔH = 247.4 J g⁻¹) standards. Nitrogen was used as a purge gas to prevent condensation in the cells. Samples (8-10 mg) were hermetically sealed in aluminium pans; an empty pan was used as a reference.

The crystallisation onset temperatures of the four matrices (M1, M2, M3 and M4) were determined as the intersection of the baseline with the absolute highest tangent of the first crystallisation exotherm. The following time-temperature program was applied: (1) heating to 80°C at 15°C min⁻¹ and holding for 10 min to erase the thermal memory, (2) cooling at -0.5°C min⁻¹ to -60°C. This cooling rate of 0.5°C min⁻¹ was used to mimic the slow cooling conditions used during fractionation.

The CP of the different olein obtained after filtration was determined as the intersection of the baseline with the absolute highest tangent of the first crystallisation exotherm, according to the following time-temperature program: (1) heating to 80°C at 15°C min⁻¹ and holding for 10 min at 80°C to erase the thermal memory, (2) cooling to 30°C at -25°C min⁻¹, holding this temperature for 2 min and finally cooling down to -25°C at a rate of -3°C min⁻¹. All DSC measurements were done in triplicate.

**Positional Isomerism by Silver Ion High-performance Liquid Chromatography - Mass spectroscopy (Ag-HPLC-MS)**

The positional isomerism of some TAGs from selected olein was determined by Ag-HPLC-MS, as described by Santoro et al. (2018). This separation technique uses columns where silver ions are bound to the stationary phase and rely on the ability of the π electrons of unsaturated fatty acids to react with silver ions to form polar complexes. The separation system was an HPLC Ultimate 3000 (Thermo Fisher Scientific, Italy) interfaced through an APCI ionisation source to a linear ion trap coupled to a high-resolution mass analyser (LTQ-Orbitrap Thermo Fisher Scientific, Italy). The ionisation source was heated at 450°C and used in positive ions mode. The instrument was equipped with a silver-modified cation exchange column (Luna SCX, Phenomenex, 150 × 2.0 mm, 5 μm, 100 Å). A gradient separation made up of two isocratic steps at different mobile phase compositions (n-heptane:ethylacetate 93:7 and 90:10) was used. The flow rate was 0.300 mL min⁻¹, and the injection volume was 10.0 μL. The mean of two measurements is reported.
Statistical Analyses

Data were statistically analysed using Minitab 19 software (Minitab LLC, USA). All assays were conducted in triplicate unless otherwise specified. Means and standard deviations were calculated and differences between means were determined with a significance level of \( \alpha = 0.05 \).

RESULTS AND DISCUSSION

Compositional properties by HPLC, IVs by titration and crystallisation onsets by DSC of the 4 matrices are listed in Table 1. The major \( S_tS_tS_t \) TAG was PPP, followed by \( P_tS_t \) (S: stearic acid); the \( S_tS_tS_t \) content increased gradually from 7.0% in M1 to 9.8% in M4. On the contrary, the DAG content was slightly reduced from 9.3% to 8.9% due to the lower DAG content in the stearin. The IV decreased from 52.1 to 49.9. The crystallisation onset shifted towards higher temperatures; this shift could only be attributed to the nucleation effect of higher amounts of \( S_tS_tS_t \) in the matrices.

The crystallisation behaviour of PO was investigated during pilot scale dry fractionation. A water-cooling program was applied, and the oil temperature response was recorded. The crystallised oil was sampled at regular intervals and the SFC was immediately measured by p-NMR; this gave an insight into the rate of crystallisation in the different matrices. At the same time, a microstructural analysis was performed by optical microscopy. Vacuum filtrations were carried out to recover the olein, the composition of which was determined by HPLC, and the polymorphism of the crystals by powder X-ray diffraction. Figure 2 represents the cooling curves, with water and oil temperatures plotted as a function of time. After fast cooling, the oil was brought into supercooling conditions for crystal initiation and the crystal growth was examined for 240 min. This period was called “main crystallisation”. \( T_c \) was the oil temperature at the beginning of the main crystallisation step. Due to crystallisation exothermicity, some heat was released during the crystal growth. When this heat released by crystallisation exceeded the heat of dissipation capacity of the crystalliser, a rise in oil temperature was observed. \( T_{\max} \) and \( T_{\min} \) corresponded to the minimum and maximum oil temperatures observed during the crystal growth respectively. After 240 min of main crystallisation, the oil was cooled down to 25°C (\( T_f \)) before proceeding to membrane press filtration.

Figure 3a shows the oil temperature rise (\( T_{\max} - T_{\min} \)) during the main crystallisation for the four matrices. This temperature rise was particularly important for M3 and M4 having the highest \( S_tS_tS_t \) contents. A sharp increase was observed between M2 and M3 and the highest value was detected for M4 (+1.8°C). Figure 3b illustrates the minimum oil temperature achieved during the main crystallisation before the rise (\( T_{\min} \)); this temperature increased with the \( S_tS_tS_t \) content, sharply from M1 to M2, and then gradually from M2 to M4. The \( \Delta T_{\min} \) between M1 and M4 was ~5°C. The time before the oil temperature rise was also determined (Figure 3c); it was significantly shorter for M2, M3 and M4 compared to M1 (\( p \)-values <0.05), and M2 had the highest reduction (40 min). In all cases, the crystallisation exothermicity caused a rise in oil temperature during the crystal growth. This rise was more important and occurred at higher temperatures in the matrices with higher \( S_tS_tS_t \) content. It led to partial crystal melting, affecting the crystal slurry quality, especially for M3 and M4. The time before the temperature rise was shorter for higher \( S_tS_tS_t \) contents but surprisingly, M2 showed the shortest time tending to indicate a faster crystal initiation.

Figure 4a presents the SFC of the crystallising oils determined by p-NMR as a function of time during the main crystallisation where M3 and

| Table 1. Tri-Saturated Triacylglycerol and Diacylglycerol Contents, Iodine Values (Wijs) and Crystallisation Temperature Onsets for M1, M2, M3 and M4 |
|-----------------|--------|--------|--------|--------|
|                  | M1     | M2     | M3     | M4     |
| **TAG (% HPLC)**|        |        |        |        |
| MP               | 0.5    | 0.5    | 0.5    | 0.5    |
| PPP              | 5.1 (+/- 0.1) | 5.6 (+/- 0.0) | 6.1 (+/- 0.0) | 7.5 (+/- 0.1) |
| P_tS_t           | 1.2 (+/- 0.1) | 1.2 (+/- 0.1) | 1.3 (+/- 0.1) | 1.5 (+/- 0.1) |
| P     S           | 0.1    | 0.2    | 0.2    | 0.2    |
| SSS             | 0.1    | 0.1    | 0.1    | 0.1    |
| Total S_tS_t     | 7.0 (+/- 0.1) | 7.6 (+/- 0.0) | 8.2 (+/- 0.0) | 9.8 (+/- 0.1) |
| **DAG (% HPLC)** |        |        |        |        |
| 9.3 (+/- 0.1)    | 9.2 (+/- 0.1) | 9.1 (+/- 0.0) | 9.8 (+/- 0.1) |
| **IV (Wijs)**    | 52.1 (+/- 0.2) | 51.6 (+/- 0.2) | 51.2 (+/- 0.2) | 49.9 (+/- 0.2) |
| **Crystallisation temperature onset (°C by DSC)** | 22.3 (+/- 0.1) | 25.1 (+/- 0.1) | 26.0 (+/- 0.1) | 27.7 (+/- 0.1) |

Note: DAG - diacylglycerol; TAG - triacylglycerol; M - myristic acid; P - palmitic acid; S - stearic acid; St - saturated fatty acids.
M4 crystallised faster than M1. However, their crystallisation kinetics slowed down starting at 180 min. Interestingly, M2 presented the highest SFC from 90 min, reaching a plateau after 150 min. The final SFC after 240 min was similar (~ 9%) for all the matrices. The crystallising oil was filtered under vacuum every 30 min, and the composition of the collected olein was evaluated by RP-HPLC. The residual StStSt content in the olein is shown in Figure 4b. This content was higher in M3 and M4 oleins, in relation to the higher content in the initial matrices. However, the residual content in M2 olein was lower than in M1 olein after 90 min.

Polymorphism of the crystals was evaluated every 30 min during the main crystallisation. M1 and M2 showed β' polymorphism during the main crystallisation time, with WAXD peaks at 4.3 Å, 4.2 Å, 4.0 Å and 3.8 Å (Figure 5). In both cases, a double-chain length packing was observed (β'-2L). M3 started to crystallise in β'-2L as well but a β form was observed after 210 min, with WAXD peaks at 3.9 Å and 4.6 Å. The β form also corresponded to
double chain length packing. The appearance of this β-2 form was accompanied by a slight decrease in the intensity of the β'-2L peaks. The same was seen for M4 with the β-2 peaks appearing already after 180 min. This β form most probably resulted from re-crystallisation of partially melted β'-2L crystals.

Crystal morphology was observed by optical microscopy, and the differences observed are illustrated in Figure 6a. After 90 min of crystallisation, bigger crystals were formed in M3 and M4, which are the matrices with higher StStSt content (= 150-250 μm, compared to 50-150 μm) for M1 and M2). M1 showed aggregates with cauliflower-like morphology, growing from 90 to 240 min. After 240 min, M2 presented a particular morphology with aggregates made of small round-shaped structures and well-defined smaller urchin-like shaped crystals (= 25-50 μm) growing on the external surface (see arrows in Figure 6b). Very small crystals were also visible in the bulk. M3 and especially M4 crystals were more dislocated in consequence of the partial crystal melting as previously mentioned. Crystal size of M3 did not change over time (150-250 μm). However, a less dense layer was observed on crystal surface, due to re-crystallisation of the liquid phase in β-2L arising from partial melting of β'-2L crystals. A much more disaggregated structure was observed for M4, but still with this less dense layer on the crystal surface (β crystals on the surface of the β' ones).

Membrane press filtrations were carried out at 25°C (final cooling temperature). The olein yields were calculated based on the collected stearin and olein fractions. Results are presented in Figure 7a. The increase in StStSt content is associated with a decrease in olein yield; a Δyield of -12% is observed between M1 and M4. M3 (73.0 ± 1.6%) and M4 (64.2 ± 0.9%) matrices have significantly lower olein yield compared to M1 (76.1 ± 0.5%) (p-value = 0.0034, and 0.0001 respectively). The same is expected for M2 but this is not the case; the olein yield is even slightly higher in M2, highlighting again the unexpected behaviour of this matrix. The IVs by Wijs and cloud point by DSC of the oleins are presented in Figures 7b and 7c. Oleins from M3 and M4 had higher IV which correlated to lower yield. The IV of M2 olein was the highest and was significantly higher compared to M1 olein (p-value = 0.0017). In general, oleins with a higher IV are characterised by a lower CP. This is not the case for M2 olein, which has the highest CP while having a higher IV.

The olein compositions were analysed by RP-HPLC; the StStSt, St2U, StU2, UUU and DAG contents are presented in Table 2. The StStSt and DAG contents decreased from M1 to M4, whereas the unsaturated (UUU) and di-unsaturated (StU2) contents increased. However, the St2U content was weakly modified. Both DAG contents and TAG compositions were in line with normal variations of the IV and the cloud point.

M2 olein has an unusual behaviour. It combines a yield higher than expected with a higher IV and a higher CP. The RP-HPLC profiles did not show any peculiar composition associated with M2 olein. However, this technique does not allow the differentiation of positional isomers. In order to
TABLE 2. TRI-SATURATED, DI-SATURATED, DI-UNSATURATED AND TRI-UNSATURATED TAG CONTENTS IN THE OLEIN FROM M1, M2, M3 AND M4 AFTER PRESS FILTRATION AT 25°C

<table>
<thead>
<tr>
<th></th>
<th>Olein from M1</th>
<th>Olein from M2</th>
<th>Olein from M3</th>
<th>Olein from M4</th>
</tr>
</thead>
<tbody>
<tr>
<td>StStSt</td>
<td>1.2 (+/- 0.0)</td>
<td>1.1 (+/- 0.1)</td>
<td>1.0 (+/- 0.0)</td>
<td>0.8 (+/- 0.0)</td>
</tr>
<tr>
<td>St2U</td>
<td>44.3 (+/- 0.0)</td>
<td>44.7 (+/- 0.1)</td>
<td>44.8 (+/- 0.1)</td>
<td>44.8 (+/- 0.1)</td>
</tr>
<tr>
<td>StU2</td>
<td>38.0 (+/- 0.3)</td>
<td>38.4 (+/- 0.1)</td>
<td>38.6 (+/- 0.0)</td>
<td>38.9 (+/- 0.1)</td>
</tr>
<tr>
<td>UUU</td>
<td>5.9 (+/- 0.1)</td>
<td>6.0 (+/- 0.0)</td>
<td>6.0 (+/- 0.1)</td>
<td>6.1 (+/- 0.0)</td>
</tr>
<tr>
<td>DAG</td>
<td>10.6 (+/- 0.1)</td>
<td>9.7 (+/- 0.1)</td>
<td>9.5 (+/- 0.0)</td>
<td>9.4 (+/- 0.0)</td>
</tr>
</tbody>
</table>

Note: DAG - diacylglycerol; TAG - triacylglycerol; St - saturated fatty acids; U - unsaturated fatty acids.
identify the reasons for the peculiar behaviour of M2 olein, oleins from M1, M2, M3 and M4 were analysed by Ag-HPLC. This technique is the most reliable for the regiosomeric separation of TAG. P2O and PO2, the most abundant TAGs, were highlighted and the corresponding 1,3-dipalmitoyl-2-oleoyl glycerol/1-oleoyl-2,3-dipalmitoyl glycerol (POP/OPP) and 1,2-oleoyl-3-palmitoyl glycerol/1,3-dioleoyl-2-dipalmitoyl glycerol (POO/OPO) contents were calculated as reported in Figures 8a and 8b. The relative percentage of OPP was higher and the one of OPO lower in the olein from M2 compared to the other matrices. These differences are highly significant (p-values = 0.0009 and 0.0002 for OPP and OPO respectively), thus confirming our hypothesis that the atypical behaviour of M2 olein arises from differences in positional isomers.

CONCLUSION

This study showed that the StStSt content in palm oil has a significant impact on crystal initiation and crystal growth. Control of the oil temperature increase was more difficult for high StStSt contents (matrices M3 and M4), inducing a partial melting of the initially formed β'-2L crystals. The resulting liquid is then further recrystallised in β-2L, leading
to a mix of β’ and β polymorphic forms. Although crystallising both in β’ form, M1 and M2 had different crystal morphology. The time before temperature rise was the shortest for M2; M2 also crystallised faster than the other matrices and its StStSt content was more efficiently removed in the olein during vacuum filtrations. The highest IV and the highest cloud point as well as a higher yield than expected were observed for M2 olein after press filtration at 25°C, which is atypical behaviour. RP-HPLC could not explain this unusual behaviour; however, Ag-HPLC showed significantly more POP and less OPO in the M2 olein. A particular PPP/P2O/PO2 co-crystallisation behaviour at M2 composition explains the atypical behaviour of the M2 matrix during crystallisation to produce olein IV 56.

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