A spinning disc study of fouling of cold heat transfer surfaces by gel formation from model food fat solutions

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Abstract

The formation of immobile gels on heat transfer surfaces (‘coring’) caused by cooling fat solutions below their cloud point was studied using a novel spinning disc apparatus (SDA). The SDA features a cooled, removable heat transfer surface with well defined heat and mass transfer characteristics. Measurements of heat flux were combined with computational fluid dynamics simulations to yield reliable estimates of the surface temperature and shear stress. Fouling studies were performed with model solutions of 5 wt% tripalmitin in a paraffin oil operating in the ‘cold start’ mode, wherein the experiment starts with the surface colder than the steady state, simulating one mode of operating a standard ‘cold finger’ experiment. Local heat flux measurements allowed the thermal fouling resistance to be monitored: deposit mass coverage and composition were also measured. The cold surface promotes the rapid formation of an initial gel layer, followed by a period of linear fouling, and finally falling rate fouling behaviour. The linear fouling rate was relatively insensitive to temperature and shear rate, while the fouling rate in the falling rate regime was found to depend on the temperature driving force for crystallisation kinetics. The solids fraction within the deposit layer increased over the duration of a 12 h fouling test, indicating rapid ageing. The rheological properties of the deposits were highly sensitive to solids fraction.

Keywords: crystallisation, fats, fouling, freezing, gel
1. Introduction

The accumulation of unwanted solids on process surfaces is a persistent problem. These materials usually have low thermal conductivity, creating significant resistance to heat transfer. This fouling deposition can also cause partial or complete blockage of piping and process equipment, reducing flow rate together with increasing pressure drop. The formation of deposits from liquid fats on cold surfaces can affect both heat exchangers and distribution lines in the food sector. This ‘coring’ of distribution lines can impair product quality by contamination and by harbouring micro-organisms, as well as reducing the performance of trace heating or cooling designed to maintain the fat in a particular state.

Coring is an example of crystallisation fouling, which Epstein (1983) defined as deposition resulting from solubility differences, such that solids are generated from solution at the heat transfer surface. The fouling layer can form via heterogeneous growth on the surface, as in water scaling, or by crystallisation of the higher melting point components in the solution at the cold interface to yield a viscous, immobile gel which can harden to give a semi-solid deposit over time. The formation of a viscous gel layer is studied here using model solutions of a high melting point fat, tripalmitin (PPP), in a non-solidifying paraffin solvent.

Food fats are multi-component mixtures and coring shares many features with wax deposition in subsea crude oil pipelines (Elphingstone et al., 1999; Ribeiro et al., 1997). The major components are triglycerides, with smaller quantities of diglycerides, and these crystallise when the temperature falls below their cloud point, \( T_c \). The cloud point is defined as the temperature at which the solid particles are first detected, and is lower than the equilibrium melting temperature for a given composition, as cooling-driven crystallisation requires the solution to be supersaturated. Nucleation is a kinetic process, however, and a crystal seed or foreign surface can promote heterogeneous nucleation and growth in this region (Mullin, 1993). The general scenario for crystallisation fouling of food fats is that the solution must firstly reach supersaturation, followed by nucleation and orientation of crystallites, ending up with aggregation and growth of the crystals (Hartel, 2001; Walstra et al., 2001). Whereas a number of models have been reported for deposition on oil pipelines from waxy crudes (Akbarzadeh and Zougari, 2008; Bidmus and Mehrotra, 2004; Singh et al., 2000) freezing fouling in food fats has attracted relatively little attention:
Fernandez-Torres et al. (2001) presented a fouling regime map of binary fat/solvent solutions based on simple thermodynamic and heat transfer relationships, and developed a heat transfer based model for coring in pipes carrying a laminar flow of fat. The controlling step in deposit growth is largely unknown, and key mechanisms, whether normal growth (Jackson and Chalmers, 1956), two-dimensional nucleation (Hillig, 1966) or screw dislocation (Hillig and Turnbull, 1956), remain to be identified.

Freezing fouling may occur at rest, such as in a reservoir, or under conditions of high shear rate and large temperature gradients, as in a scraped surface heat exchanger. For this reason, several experimental studies have investigated the effect of thermo-mechanical history on fouling behaviour. The temperature difference between the bulk liquid and the cold wall is generally regarded as the thermal driving force for deposition. Bidmus and Mehrotra (2004) reported an increase in wax deposition from oils with an increase in the temperature difference across the deposit layer. Increasing the flow rate may cause deposit removal because of the higher shear force. Other studies (Singh et al., 2000) have shown that under conditions of constant temperature difference between the bulk crude oil and deposit/oil interface, the deposit thickness decreased as the Reynolds number was increased under both laminar and turbulent flow conditions. Most tests employ constant overall temperature difference: as deposit accumulates, the solution-deposit interface temperature rises and the driving force for deposition decreases. Fouling resistance-time ($R_f - t$) plots usually exhibit falling rate or asymptotic fouling behaviour (Epstein, 1983) and can often be fitted to the Kern-Seaton (1959) model, even though this model attributes the observed retardation to increased ‘removal’ as the deposit grows rather than temperature effects. Deposit removal by mechanisms such as spalling can occur, as reported by Fitzgerald et al. (2004).

The most commonly used experimental configurations for studying freezing fouling in waxy crude oils are the flow cell loop (Fitzgerald et al., 2004; Parthasarathi and Mehrotra, 2005) and the coldfinger (Jennings and Weispfennig, 2005). Recently, Nigo et al. (2009) reported the design and operation of a novel spinning disc apparatus (SDA) featuring a cooled, rotating, vertical cylinder where deposition occurs on the cold, removable base (the sides are insulated). Food fat fouling behaviour is investigated here using a modified version of the SDA reported by Nigo et al. The SDA was fitted with a heat flux sensor which allows the change in local thermal fouling resistance, $R_f$, to be monitored. Model solutions of 5 wt% PPP in a paraffin oil, similar to those used by Fitzgerald et al. (2004) and Nigo et al. (2009),
were studied and the effects of disc rotation speed, \( \omega_d \), time, and temperature driving force on fouling behaviour were investigated. The thermal performance of the SDA was characterised more fully using computational fluid dynamics (CFD) simulations, so that surface temperature and surface shear stress could be predicted and manipulated readily. The deposit gels formed were analysed to track changes in composition resulting from ageing as reported for wax deposits by Singh et al. (2001b).
2. Materials and methods

2.1. Spinning disc apparatus (SDA)

Figure 1(a) shows a schematic of the deposit generation section in the SDA. A detailed description of the apparatus is given in Nigo et al. (2009). The apparatus consists of a cylindrical can partially immersed in a reservoir of test solution. Rotation is provided by a stepper motor. A recirculating water/glycol bath is used to regulate the temperature of the can, \( T_{cw} \), whilst the temperature of the bulk solution, \( T_b \), is maintained by a water jacket around the reservoir fed by a second, heating, recirculator. The can sides are insulated so that heat transfer occurs primarily through the base. A magnetic stirrer located at the base of the reservoir provides agitation and maintains temperature uniformity in the bulk solution. The temperature of the bulk solution, coolant and water jacket are monitored by T-type thermocouples connected to a datalogger.

Figure 1(b) shows the arrangement of the can base, which was used for heat transfer studies and fouling tests. A micro-foil heat flux sensor (Rhoptio, UK, type 27160) is located between a brass block and a removable 316 stainless-steel disc. The sensor was connected to a battery powered solid-state data collection unit located on the top of the can, so that data could be collected without the use of slip rings etc. and downloaded after an experiment. The thermal resistance of the combined plates and heat flux sensor, \( R_w \), approximately \( 8.2 \times 10^{-4} \text{ m}^2 \text{K W}^{-1} \), is shown later to be small compared to the other terms.

2.2. Heat transfer tests

The surface temperature, \( T_s \), is a key parameter in the fouling mechanism so a series of heat transfer tests were performed to characterize the thermal performance of the system. The same water/glycol mixture was used on the coolant side, in order to maintain coolant flow similarity. Liquid paraffin (the solvent in the fouling tests), 30 wt% and 78 wt% water/glycerol mixtures were used as the reservoir (bulk) liquid. These differed primarily in viscosity, so that a range of hydrodynamic conditions could be studied over the range of rotational speeds accessible with the stepper motor. The bulk liquid hydrodynamics were characterised by the Reynolds number, \( Re_r \), defined as
\[ Re = \frac{r_d^2 \omega_d \rho_b}{\mu_b} \]  

where, \( r_d \), \( \omega_d \), \( \mu_b \) and \( \rho_b \) are the disc radius, the angular velocity of the can, the dynamic viscosity and the density of the bulk solution, respectively. The viscosity was evaluated at the estimated surface temperature, \( T_s \). The experimental conditions and bulk liquid properties are summarized in Table 1.

Heat transfer tests were performed with a charge of 2 L of liquid in the reservoir, as used in fouling tests. The can was initially isolated from the reservoir and coolant circulated to bring it to the required temperature. The data loggers were then activated, the can lowered into the bulk liquid and rotation started. The data reported are the values obtained after steady state was reached.

2.3. Model fat solutions

Test solutions were prepared by dissolving tripalmitin (PPP, >85% purity; density, 984 kg m\(^{-3}\), Sigma Chemicals, UK) in liquid paraffin at 60 °C to give solutions with composition 2.5, 5, 10, 20 and 50 wt%. All fouling tests reported here were performed with 5 wt% solutions. Solution composition was determined by gas chromatography (GC) on samples diluted in n-heptane (Hewlett Packard Agilent 6890 GC equipped with a dimethylpolysiloxane column, 10 m×0.32 mm×0.1 μm, and a flame ionisation detector using helium as the carrier gas). The GC was calibrated and programmed using the BS EN 14105:2003 method.

2.4. Cloud and melting points measurement

The cloud point of PPP solutions was measured under quasi-static conditions, where the only liquid motion was that provided by a magnetic stirrer. The cloud point, \( T_c \), is the onset of spontaneous precipitation and was measured using a turbidity sensor (ODEON meter, NEOTEK-ONLYE, France). The turbidity probe was immersed in the sample held in a 50 mL jacketed beaker (4 cm in diameter and 6 cm in height) and stirred by a Teflon coated magnetic bar at low rotational speed. The sample temperature was measured using a T-type thermocouple and was initially held at 60 °C for 10 min to eliminate any thermal history. The jacket liquid temperature was manipulated to lower the sample temperature at (i) 0.5 and (ii) 1.5 K min\(^{-1}\). Turbidity was recorded at 10 s intervals and \( T_c \) taken as the temperature at which a significant and continuous rise in nephelometric turbidity units (NTU) was observed.
The melting point and melting enthalpy of samples was determined using a Perkin-Elmer Pyris 1 differential scanning calorimeter (DSC) fitted with a refrigerated cooling system. Scans were recorded from 0 to 80 °C at heating and cooling rates of 10 K min\(^{-1}\).

2.5. Rheological tests

A short rheological study was performed to investigate the effect of shear on the crystallisation of PPP from model solutions undergoing cooling. In the SDA the deposit gels are formed at a cold surface subject to shear. Samples of 5 wt% PPP solution were tested using serrated parallel plates on a Bohlin CV120 controlled stress rheometer (25 mm diameter, 0.1 mm gap). Temperature control was provided by a Peltier device located beneath the lower plate. The plates were preheated to 55 °C before loading the sample and the sample was initially held at this temperature, at rest, for 10 min. The sample was then subjected to steady shear at a fixed shear stress of 2 Pa while subject to a temperature ramp of steady decrease from 55 to 5 °C at 5 K min\(^{-1}\), followed by 10 min at 5 °C. The shear stress was set at 2 Pa in order to avoid shearing any developing microstructure severely while maintaining sufficient measurement accuracy. This shear stress is also representative of the shear stress exerted by the solution on the disc and deposit surfaces in fouling experiments.

2.6. Numerical simulation

Computational fluid dynamics simulation of the flow and temperature fields in the bulk liquid in the SDA were used to generate estimates of film heat transfer coefficient from the bulk liquid and the can base plate and the surface shear stress. These simulations can be performed with reasonable accuracy as the device is operated in the laminar flow regime and yield estimates of surface temperature and shear stress. The commercial finite element method software COMSOL Multiphysics (version 3.5, Chemical Engineering Module) was used to solve the continuity, Navier-Stokes and the steady state energy equations for a Newtonian liquid. The flow was modelled as being axisymmetric, incompressible, and steady state. Physical properties, such as density, thermal conductivity and specific heat capacity, were assumed not to change significantly with temperature and were assumed uniform throughout. This study extends the numerical simulation of Nigo et al. (2009), using similar boundary conditions but using modified fouling cell plate and quantitative settings to include a heat flux sensor and different base plate geometry. Moreover, the effect of thermal resistances from the coolant flow and combined plates were taken
into consideration to yield more accurate simulation results. The converged solution took approximately 15 min on a desktop PC with a 3.16 GHz dual core processor and 3.33 GB RAM.

2.7. Fat deposition studies

Deposition experiments reported here were performed in a ‘cold start’ mode, wherein the test surface and reservoir were initially brought to their respective operating temperatures by circulation of the heat transfer medium before the can was immersed in the bulk liquid and rotation started. This resulted in an initial transient where the test surface temperature increased from one close to that of the coolant liquid to an intermediate value. The transient was studied separately using PPP-free paraffin liquid and data reported later show that the transient lasted ~ 60 s. This cold start mode resulted in the formation of an initial gel layer but avoided transients associated with mixing if the PPP charge were added to the solution to start the fouling test. An alternative mode, of warm start, is under development and will be reported in due course.

The rotation speeds, \( \omega_d \), employed were 2.2, 3.5, 5.4 and 7.3 rad s\(^{-1}\), corresponding to \( \text{Re}_r \) of 26, 41, 64 and 86, respectively. The cloud point of the 5 wt% solution was 29.8 °C (at a cooling rate of 0.5 K min\(^{-1}\)) and coolant temperatures, \( T_{cw} \), were selected to give a range of precipitation driving forces, namely 9.8 °C (\( T_c - 20 \)), 19.8 °C (\( T_c - 10 \)), 24.8 °C (\( T_c - 5 \)) and 29.8 °C (\( T_c \)). Fouling tests normally lasted 12 h, but a series of interrupted tests were performed at one condition, for 1, 3, 6, 9 and 12 h, in order to establish reproducibility and to allow stages in deposit growth to be evaluated. The final deposit was removed from the surface using a plastic spatula, weighed and stored for further analysis.

2.8. Deposit analysis

The deposits consisted of gels of PPP crystals in a liquid matrix and contained PPP in the form of solids and entrained solution. The total PPP content, \( w_T \), was determined by dissolving samples in a measured excess of \( n \)-heptane and measuring PPP content by GC as described above.

The solids content of the deposit, \( w_X \), was determined by filtration. About 3 g of deposit was loaded on to a 0.2 μm polytetrafluoroethylene membrane filter paper (Cole-Parmer, USA) and vacuum applied. The filtered solids were washed with hexane (analytical reagent grade, Fisher Scientific, UK) at ambient temperature to remove any entrained paraffin followed by acetone to remove traces of hexane, then
allowed to dry at room temperature for an hour before weighing. The rheology of the deposits was studied using a Bohlin CVO 120 controlled-stress rheometer operating in oscillatory testing mode as reported by Nigo et al. (2009) with the temperature of the bottom (stationary) plate set at 20 °C. Tests were performed using roughened plates (top 25 mm diameter, base 50 mm diameter, gap 1 mm). The oscillating frequency was set at 10 rad s\(^{-1}\) (1.59 Hz) and the sample subjected to shear stress from 2.4 Pa to 29.3 kPa in an ascending ramp.

2.9. Statistical analysis

Analysis of variance (ANOVA) was applied and, when the effect of the factors was significant \((p < 0.05)\), the Fisher least significant difference (test was used at 95% confidence level.)
3. Results and discussion

3.1. Crystallisation under quiescent and shear conditions

Figure 2 summarises the melting and cloud point results for the model solutions of PPP in liquid paraffin and can serve as a simple crystallisation map. Both \( T_m \) and \( T_c \) decrease with decreasing PPP concentration, as expected. The melting point of the PPP melt (i.e. no paraffin added), at 65.4 °C, compares favourably with the value of 66 °C reported by Sato and Kuroda (1987). The enthalpy of melting, \( \Delta H_{fp} \) of the PPP melt was 108 kJ mol\(^{-1}\), which is lower than that obtained by Kellens et al. (1990), which was attributed to the differences in purity (165 kJ mol\(^{-1}\), PPP ~ 99 % pure). The purer the sample, higher the enthalpy value as there are more PPP crystals per unit mass.

\[ \ln X_{ppp} = \frac{\Delta H_{fp}}{R} \left[ \frac{1}{T_m^p} - \frac{1}{T_m} \right] \]  

(2)

where \( X_{ppp} \) is the mole fraction of PPP in solution, \( R \) is the universal gas constant, and \( T_m^p \) is the melting point of the pure component. The solutions were prepared with set mass fractions (as plotted in Figure 2) and the PPP mol fraction was estimated using relative molecular masses for PPP of 807.35 g mol\(^{-1}\) and 324 g mol\(^{-1}\) for paraffin \( (C_nH_{n+2}, \text{where } n = 23, \text{from GC analysis}) \). The \( \Delta H_{fp} \) value obtained from the data fit, of 211 kJ mol\(^{-1}\), differs noticeably from the DSC analysis, which can be attributed to impurities in the material and solvent-solute interactions (Davey and Garside, 2000).

The cloud points show a noticeable influence of cooling rate, with lower values observed for slower cooling rates, as discussed by Saiban and Brown (1997). The width of the metastable zone lay between 10 – 21 K of supercooling over the concentration range studied. The simple crystallisation map deduced from the study can serve as a guide to predict the likelihood of crystal formation at a given temperature and concentration. For the cold start operating mode employed in the experiments reported here, immersion of the cold test section in the warm solution at
the start of the experiment results in a high initial cooling rate: Figure 2 indicates that this is likely to promote homogeneous nucleation and gel formation, as observed in the experiments.

An example of crystallisation in the presence of shear is shown in Figure 3, where a 5 wt% PPP solution sample was subjected to a steady shear stress of 2 Pa as it was cooled from 55 to 5 °C at 5 K min\(^{-1}\). The \(T_m\) value for this solution under quiescent conditions was measured at 49.1 °C and the cloud point (at 0.5 K min\(^{-1}\)) was 29.8 °C. The viscosity was calculated from the measured shear rate and is presented as the relative viscosity, \(i.e\). made dimensionless by division by the viscosity of the paraffin solvent measured under similar conditions. The solution exhibits an initial region, labelled A, in which the relative viscosity is close to unity, indicating the absence of particles.

After about 7.5 minutes the relative viscosity increases markedly (region B), corresponding to the appearance of crystals and formation of a suspension: the AB transition marks the ‘mechanical’ cloud point. The sample temperature, at \(\sim\) 17 °C, is 12 K lower than the quiescent \(T_c\) value. This is partly due to the turbidity meter detecting the onset of crystallisation, \(i.e\). a low number concentration of particles, whereas the rheometer is sensitive to the formation of a network. The relative viscosity increases by four decades, indicating the formation of a gel, which increases in stiffness across region C as the solution continues to cool. The change in mass of crystals in this region is small (compare the solubilities in Figure 2), indicating that the change in viscosity is related to gel behaviour. A high viscosity plateau is reached at \(\sim\) 10 °C, where the peaks are indicative of wall slip and a network in the bulk phase. These characteristics were confirmed by tests in a Linkam shear cell, a controlled shear device fitted with observation windows. Similar transitions in fats have been reported previously (De Graef \textit{et al.}, 2009; Tarabukina \textit{et al.}, 2009; Walstra \textit{et al.}, 2001).

Figure 3 confirms that as the fat solution cools, once it starts to nucleate, it can quickly form a highly viscous gel. The BC transition could be interpreted as a gelation temperature. Comparing change in relative viscosity in region C with the solids content estimated from the equilibrium locus in Figure 2 (data not reported) showed the relative viscosity is not a simple function of the final solids content of gel, indicating that the crystallisation under shear condition is under kinetic control. Similar behaviour was observed with other PPP concentrations. Whilst this test does not replicate the conditions on the SDA exactly, it does confirm that cold start
experiments will form gels rapidly, and that the first appearance of crystals is sensitive to shear and cooling conditions.

3.2. Heat transfer

Spinning disc devices are employed in mass transfer studies as the mass flux (and film mass transfer coefficient) does not vary with radial position across the surface. Sparrow and Gregg (1959) analysed the heat transfer behaviour of spinning discs operating in the laminar regime and showed that the heat flux, $q$, film heat transfer coefficient, $h_b$, and surface temperature, $T_s$, are similarly uniform. It should be noted that the laminar regime extends to Reynolds numbers of $\sim 100,000$. Measurements of the heat flux yield the overall heat transfer coefficient, $U$, defined as

$$ q = U (T_b - T_{cw}) = h_b (T_b - T_s) $$  \hspace{1cm} (3)

Writing $U$ in terms of resistances in series yields

$$ \frac{1}{U} = \frac{1}{h_b} + R_{cw} + R_w = \frac{1}{h_b} + R_{else} $$  \hspace{1cm} (4)

where $R_{cw}$ and $R_w$ refer to the thermal resistance of the coolant flow and plates, respectively, and are lumped together as $R_{else}$. The $h_b$ term is expected to follow a dependency of the form (Sparrow and Gregg, 1959);

$$ Nu \equiv \frac{h_b r_d}{k_b} = a Re_0^{0.5} Pr^{1/3} $$  \hspace{1cm} (5)

where $Nu$ is the Nusselt number based on the disc radius, $k_b$ the thermal conductivity of the bulk liquid, $Pr$ its Prandtl number and the parameter $a$ is a constant. Equation (5) suggests that the data obtained for different liquids should follow similar behaviour when plotted as $Nu/Pr^{1/3}$ against $Re_0$. The film heat transfer coefficient is not directly accessible, however, and some estimation is required.

An alternative Nusselt number, $Nu^*$, can be based on $U$ and its relationship to $Nu$ given by Equation (4):

$$ Nu^* \equiv \frac{U r_d}{k_b} = \frac{Nu}{1 + R_{else} h_b} $$  \hspace{1cm} (6)

Inspection shows that at low $Re_0$, when $h_b$ is small and is expected to influence heat transfer strongly, $Nu^* \sim Nu$ and the dependency on $Re_0$ is expected to mirror Equation (5). At larger values of $Re_0$, Equation (6) indicates that $Nu^*$ is likely to give a poorer estimate of $h_b$. This is confirmed by Figure 4, where the heat transfer dependency on $Re_0$ (Equation (5)) starts to differ from the trend in the data at higher $Re_0$. $Nu^*$ differs noticeably from $Nu$ estimated using Equation (5) with $a = 0.62$, indicating that the parameter $a$ required modification, which was achieved by simulation (next section).
The above analysis assumes that $R_w$ does not vary strongly with rotation speed, which is considered reasonable because of turbulence in the coolant flow geometry and the influence of constant, conduction contributions. The thermal resistance of coolant side, $R_{cw}$, in the SDA system becomes more significant as $Re_r$ is increased and this is estimated by difference, as outlined below.

### 3.3. CFD simulations

The CFD simulations yield predictions of velocity and temperature distributions in the bulk liquid in the heat transfer experiments. The coolant flow is not modelled: its contribution is expressed as a uniform film heat transfer coefficient. Figure 5 shows the stream functions (contour lines) and the temperature profiles (coloured background) obtained for one of the sets of conditions employed in deposition tests. Two vortices are evident in the bulk liquid: an upper one driven by the rotation of the disc, and a lower one induced by the magnetic stirrer acting in the opposite direction.

Simulations were run for a set of cases covering the experimental conditions reported in Figure 4 and values of $h_b$ extracted from the simulations. $R_{else}$ is then estimated from $(1/U - 1/h_b)$ and $R_{cw}$, the thermal resistance on the coolant side, is then evaluated from $(R_{cw} - R_w)$. The static resistance associated with the base plates, $R_w$, is constant over the range of temperatures considered here. A subset of the results related to deposition testing are presented in Figure 6. The film resistance $1/h_b$ decreases with increasing $Re_r$, as predicted by the correlation (Equation (5)), and the simulation values show good agreement with the correlation: fitting the data to a power law trend yielded a power law index of 0.48, in good agreement with the value of 0.50 in Equation (5). $R_{cw}$ also decreases with rotation speed, and is consistently smaller than $1/h_b$, indicating that the dominant resistance to heat transfer lies on the bulk (fat) side.

These results allow $T_s$ to be estimated by combining Equations (3) and (4), viz.

\[
T_s(t) = T_b - \frac{q}{h_b} = T_b - \frac{T_b - T_{cw}}{R_{cw} + R_w + R_f(t) + \frac{1}{h_b}} \frac{1}{h_b},
\]

where $R_f$ is the thermal resistance of any fouling layer. Assuming that the other resistances do not vary with temperature as fouling develops allows $R_f$ to be evaluated from

\[
R_f(t) = \frac{1}{U(t)} - \frac{1}{U_0},
\]
where $U_0$ is the initial, clean, overall heat transfer coefficient. Similarly, the temperature at the base plate/deposit interface, $T_w (= T_s$ when $R_f = 0)$ can be evaluated from

$$T_w(t) = T_{cw} + q(R_{cw} + R_w)$$  \hspace{1cm} (9)

Figure 7(a) shows the initial, clean, steady state $T_s$ at over the range of $T_{cw}$ and $\omega_d$ of interest. The plots show that $T_s$ is more sensitive to $T_{cw}$. A key assumption in these estimates is that the release of enthalpy of crystallisation during deposition is negligible: this is considered further in Section 3.5.

The average shear stress imposed on the disc surface can also be calculated from the simulated velocity field. The shear stress is directly proportional to radial position, $r$, and Figure 7(b) shows the average wall shear stress calculated at $r$ from 0 to 0.035 m. The CFD calculations indicate that the average shear stress is of the order of 0.3 – 2.3 Pa for the initial clean surface. This compares favourably with the range of surface shear stresses reported by Fitzgerald et al. (2004) using a rectangular duct flow cell, of 0.5 – 4.0 Pa. Larger shear stresses could be obtained in the SDA using a different motor. Since surface temperature and shear stress are key parameters in freezing fouling, Figure 7 illustrates the range of surface conditions that can be investigated in deposition studies.

3.4. Fouling Profiles

The reproducibility of the fouling test was confirmed by conducting a series of experiments under identical conditions for various times. The $R_f - t$ plots in Figure 8 show good agreement, with a maximum difference of 3%. Subsequent tests were repeated only when the results were inconsistent with observed trends. These interrupted tests also allowed the evolution of deposit properties to be investigated.

The $R_f - t$ plots obtained in the ‘cold start’ experiments discussed here all showed similar behaviour, exhibiting three stages:

(i) an initial step in $R_f$,

(ii) a period of linear increase in $R_f$, followed by

(iii) a falling rate period, where $R_f$ approaches an asymptote.

The initial sharp increase in $R_f$ derives from the cold start mode. On immersion of the test cell, the warm solution is contacted with the cold surface (below $T_c$), inducing gel formation. At the same time, the heat flow through the cell is larger than at steady
state owing to the transient in heat conduction, resulting in a temporally large heat flux and associated $U$ value. The definition of $R_t$ in Equation (8) means that there is a transient, with negative $R_t$ values, as shown in Figure 9. The Figure shows that a thermal steady state was reached within 2 min, by which time a gel layer had been established. Figure 9 also shows the transient observed under identical conditions with paraffin alone, in which case no gel is formed and the steady state $R_t$ value is zero. Negative values of $R_t$ associated with roughness enhancing convective heat transfer have been reported by other workers, e.g. Crittenden and Khater (1987); Bansal and Müller-Steinhagen (1993), but this transient mechanism is rarely reported, mainly because the experiments start with heat transfer in steady state. The initial step in $R_t$ was reproducible (see Figure 8(a)), and decreased as the coolant temperature increased (Section 3.8).

In the subsequent stages, as deposit accumulates, the deposit-solution interface temperature, $T_s$, increases and the wall temperature, $T_w$, decreases. The calculated values of $T_s$ and $T_w$ in Figure 8(b) show a rapid increase in $T_s$ which slows down following the linear fouling stage and approaches an asymptote, which is close to $T_m$. $T_s$ is not measured directly, so there is some uncertainty in the estimate. Tests under other conditions showed similar behaviour. Figure 8(b) shows a moderate decrease in wall temperature over time, which would promote further crystallisation of the gel. It is noteworthy that the end of the linear fouling regime, at a temperature denoted $T_s = T^*$ (estimated by eye as the end of the linear region), extends beyond the point where $T_s = T_c$ (here, $T_c = 29.8 \, ^\circ\text{C}$ at 0.5 K min$^{-1}$, reached after ~ 50 min). The controlling mechanism does not, therefore, appear to be crystallisation at the deposit interface, with $T_s = T_c$, as reported by Singh et al. (2000).

The $R_t - m$ data from all experiments reported in this study are plotted Figure 10. When deposition is uniform, the relationship between $R_t$ and the mass of deposit per unit disc surface, $m$, is:

$$R_t = \frac{m}{\rho_f k_f}$$  \hspace{1cm} (10)

where $\rho_f$ is the deposit density. The deposit thermal conductivity, $k_f$, was taken to be 0.15 W m$^{-1}$ K$^{-1}$, as the thermal conductivity of solid PPP is conveniently close to that of paraffin. The Figure shows a linear relationship ($R^2 = 0.97$), in good agreement with Equation (10). The gradient, of 0.011 m$^4$ K W$^{-1}$ kg$^{-1}$, yields an estimate of $\rho_f k_f = 91$ W kg m$^{-2}$ K$^{-1}$, which is lower than the value expected for the paraffin, of 130.5 W kg m$^{-4}$ K$^{-1}$ ($\rho_f k_f$ (Fitzgerald et al., 2004) for the PPP solids will be greater again. The difference may be due to the difficulty in collecting all the deposit (affecting $m$), and
any systematic errors in evaluating $R_f$. The linear relationship does, moreover, indicate that the age of the deposit does not affect heat transfer. It is worth noting that the linear relationship of the $R_f - m$ data obtained by the interrupted tests (Figure 8) indicates that the heat flux measurements provided a reliable monitor of deposit growth.

The deposit composition data in Figure 11 exhibit a noticeable change over the course of the 12 h fouling test. The solids fraction, labelled $w_X$, increased almost linearly to about 15 wt% over the first 6 h, coinciding with the linear increase in $R_f$. The subsequent change in $w_X$ was less marked, reaching 20 wt% after 12 h. The composition of the liquid phase within the deposit, $w_S$, is calculated from the measured values of $w_T$ (GC) and $w_X$ (filtration) via:

$$w_S = \frac{w_T - w_X}{1 - w_X}$$  \hspace{1cm} (11)

The data in Figure 11 show $w_S$ being similar to that of the bulk liquid, and increasing over time to ~ 10 wt%. The increase in $w_S$ is less marked than that observed in $w_X$, and the enrichment in solution composition (increase in $w_S$) after 6 h was not statistically significant. These data indicate that the deposit undergoes ageing, mainly through increasing solids content. Further crystallisation of PPP from the solution trapped in the deposit as the local temperature decreases (see Figure 8(b)) would tend to reduce the concentration of PPP in solution. Higher resolution measurements are required to confirm whether the solution is being enriched, which would require diffusion into the deposit from the bulk solution.

One explanation for the increase in solids content is shear-induced hardening. The latter is related to the shear stress imposed on the surface, which is controlled by $\omega_d$, and liquid properties (principally $T_s$). As fouling continues, $T_s$ increases, the viscosity decreases and the solids composition required to form a gel strong enough to resist deformation by the surface shear stress will increase. These competing aspects of temperature have been reported for wax formation in crude oils by Bansal and Müller-Steinhagen (1993). Shear hardening would not require $w_S$ to increase. However, in these tests, the change in $T_s$ also reduces the shear stress imposed by the bulk liquid. Decoupling these factors and establishing the driving force for ageing is the subject of ongoing work, including imaging of the deposits.

3.5. Fouling mechanisms

$R_f - t$ profiles featuring falling rate and asymptotic behaviour are often fitted to the
Kern-Seaton model based on competing deposition and removal processes, viz.

\[ R_f = R_f^* \left[ 1 - \exp\left( -t/t_F \right) \right] \]  

(12)

where \( R_f^* \) is the asymptotic fouling resistance and \( t_F \) is a characteristic time, which can be related to initial fouling rate via \( [dR_f/dt]_{t=0} = R_f^*/t_F \). This approach has been applied to wax formation from oils by Nazar et al. (2005), Akbarzadeh and Zougari (2008) and Ramirez-Jaramilho et al. (2010) but is not appropriate here as (a) the initial linear regime is not consistent with the model; (b) fitting the \( R_f - t \) profiles to Equation (12) gave systematic deviations in fitting; and (c) the asymptotic level, \( R_f^* \), is related to \( T_m \), rather than a removal step. It is noteworthy that no removal was observed in these tests. Fitzgerald et al. (2004) also reported initial, linear fouling behaviour followed by falling rate behaviour but this was interrupted by spalling, which was not observed in these tests. Future work will consider higher rotation rates to see if this can induce spalling in these deposits.

The estimates of \( R_f \), surface and wall temperatures reported here assume that the enthalpy released on forming the deposit, i.e. from PPP crystallisation, is small compared to the total rate of heat transfer. The heat flux sensor measures the rate of heat transfer to the coolant, so if the rate of latent heat evolution were significant, the rate of convective heat transfer would be less than the measured value: Equation (3) indicates that a lower \( q \) would result in higher \( T_s \). The contribution from latent and sensible heat effects on heat transfer was therefore considered by comparing the enthalpy associated with forming the deposit with the total amount of thermal energy transferred through the heat flux sensor over the course of a 12 h fouling test.

The enthalpy of crystallisation associated with the crystals in the deposit, \( H_X \), was estimated from

\[ H_X = mw_X \Delta H_{fp} \]  

(13)

where \( m \) and \( w_X \) are the measured deposit mass coverage and solids fraction, respectively, taken from Figures 10 and 11. The latent heat of crystallisation was taken as 211 kJ mol\(^{-1}\) (Section 3.1), which is a worst case estimate as the true value is likely to be smaller, nearer 100 kJ mol\(^{-1}\). The sensible heat associated with the deposit was estimated from

\[ H_S = mC_p \left[ T_{b} - \frac{1}{2}(T_s + T_w) \right] \]  

(14)

This estimate assumes that the deposit has uniform heat capacity, \( C_p \), taken as that of the paraffin, and a linear temperature distribution across the deposit. Figure 12 shows
that $H_S$ was initially the dominant contribution, with $H_X$ increasing in importance over time, but the combined contribution ($H_O = H_X + H_S$) was consistently small, at $1 - 2\%$ of the total amount of thermal energy transferred. Neglecting latent heat effects in estimating $R_f$ therefore appears reasonable.

3.6. Growth rates

The above calculation indicates that heat transfer was not determining the rate of deposit formation. Further insight into the mechanisms involved can be obtained by comparing the observed deposit growth rate with one estimated from the maximum rate of mass transfer of PPP to the surface. The 5 wt% solutions can be considered dilute (the mole fraction of PPP in the bulk solution is ~ 0.02): the mass flux, $N_{PPP}$, from the bulk solution with PPP concentration, $x_b$, towards the surface where growth occurs (with local concentration, $x_s$) can be estimated from

$$N_{PPP} = k_m(x_b - x_s)$$  \hspace{1cm} (15)

where $k_m$ is the film mass transfer coefficient, and $x_i$ is a mass concentration (in kg m$^{-3}$). The maximum flux was estimated by setting the concentration at the surface, $x_s$, to be that in equilibrium with solid PPP at the surface temperature, $T_s$ (i.e. rapid crystallisation). The rate of deposition, $N_d$, can then be estimated from the solids content of the layer, $w_X$, via

$$N_d = \frac{k_m}{w_X}(x_b - x_s)$$  \hspace{1cm} (16)

Gregory and Riddiford (1956) reported the following expression for $k_m$ for a disc rotating in an infinite Newtonian fluid;

$$k_m = 0.62 \frac{D}{r_d} Re_{r}^{1/2} Sc^{1/3}$$  \hspace{1cm} (17)

where $Sc$ is the Schmidt number, expressed as;

$$Sc = \frac{\mu_b}{\rho_b D}$$  \hspace{1cm} (18)

Here $D$ is diffusivity of solute in solvent, which was estimated from the Wilke and Chang correlation (1955) to lie in the range of $4 - 17 \times 10^{-11}$ m$^2$ s$^{-1}$ for the PPP/paraffin solutions employed here. Calculation of $N_d$ requires evaluation of the transport parameters and $x_s$, and hence the surface temperature: $T_s$ was calculated at selected times using Equation (7) and the experimental $R_f$ value.
The actual deposition rate was estimated from the thickness growth rate, $G$, by $N_d = \rho G$. The thickness growth rate, $G$, was calculated from the $R_f$ measurements by assuming the deposit thickness, $\delta$, to be uniform and $k_f$ known, via:

$$G = \frac{d\delta}{dt} = k_f \frac{dR_f}{dt}$$  \hspace{1cm} (19)

The gradient $dR_f/dt$ (and also the surface temperature, $T_s$) were evaluated by interpolating the $R_f$ and $T_s$ data sets and extracting values at particular points.

Figure 13 shows the rates calculated for the 12 h fouling experiment in Figure 8. Over the first 300 min, where linear fouling was observed, the deposition rate calculated from the $R_f$ data was considerably smaller than the maximum PPP mass flux, indicating that fouling was not controlled by mass transfer in this regime. The mechanism remains to be determined, as the subsequent studies of temperature and velocity show the fouling rate to be relatively insensitive to these parameters.

In the falling rate regime the actual deposition rate is close to $N_d$ estimated by this simple mass transfer model. The main reason for the variation in $N_d$ over time in Figure 13 is the change in $(x_b - x_s)$ with $T_s$, and in the falling rate regime this could be attributed to mass transfer or crystallisation effects.

The transition from linear to falling rate fouling behaviour indicates a change in growth mechanism and this section considers the latter phase. No particulates were present in the bulk solution, and inspection of Figures 8(a) and (b) suggest that the growth rate in this phase is related to the surface temperature and the temperature driving force $\Delta T$, the latter being defined as the difference between $T_m$ and the temperature of crystallisation, here $T_s$. $T_s$ (and thence $\Delta T$) in the falling rate stage was estimated by interpolating the $R_f$ and $T_s$ data sets as described above and extracting values at particular points. Figure 14 shows the results obtained for the runs in Figure 8. A linear trend is evident, which is the form expected for normal growth (Borisov et al., 1968):

$$G = K\Delta T$$  \hspace{1cm} (20)

The fitted line does not pass through the origin, which is attributed to the systematic uncertainties involved in calculating $T_s$ from the experimental data.
The effect of velocity was studied at four rotational speeds, corresponding to average shear stresses on the disc surface of 0.26, 0.52, 1.03 and 1.66 Pa. Figure 15 presents the $R_f - t$ profiles obtained from experiments performed at $T_b = 60$ °C and $T_{cw} = 9.8$ °C for 12 h. Under these conditions the test section surface temperature increases with $\omega_d$, since $h_b$ is more sensitive to $\omega_d$ than $R_{cw}$. The three stages described previously are evident in each case. The initial increase in $R_f$ decreased as $\omega_d$ increased. This may occur because the incipient gel is relatively weak, and was less able to resist the shear forces imposed on it at higher $\omega_d$, or because the new, steady state temperature was warmer at higher $\omega_d$ (Equation (7)).

The linear fouling rate did not change appreciably with $\omega_d$, varying from $1.21 \times 10^{-6}$ to $1.32 \times 10^{-6}$ m² K J⁻¹. Inspection of the (estimated) $T_s - t$ plots indicated that the end of the linear fouling stage occurred at similar values of $T_s$, around 45.7 °C (i.e. $T_m - 4.6$ K). After this point, the fouling rate exhibited normal growth kinetics with $K$ value of $1 - 2 \times 10^{-5}$ μm s⁻¹ K⁻¹ and $T_s$ approaching $T_m$ as discussed above.

The fouling resistance, $R_f$, reached after 12 h decreased with increasing $\omega_d$. The associated mass of deposit ($m$, Figure 16) was similarly reduced, and the plot of $R_f$ against $m$ in Figure 10 indicates that the deposit properties were similar to those reported above. The deposit composition data in Figure 16 show no effect of $\omega_d$ on $w_S$: the variation in $w_S$ lies within the confidence limits for the hypothesis that the value is the same for all cases. The solids content, $w_X$, however increases noticeably with $\omega_d$, by a factor of $3 - 4 \times$ over the range studied. This result is consistent with the formation of a stiffer deposit to withstand the higher shear stress conditions present at its formation. An alternate explanation, given by Jennings and Weispfennig (2005), is that the thinner gel (smaller $\delta_f$) and higher $h_b$ result in a larger temperature gradient across the deposit, creating a higher diffusive flux for hardening. Again, detailed microstructural distribution data are required to elucidate this behaviour.

### 3.7. Effect of velocity

3.8. Effect of thermal driving force

Figure 17 shows the effect of thermal driving force for tests performed at $\omega_d = 5.4$ rad s⁻¹ at four different coolant temperatures. The initial degree of subcooling ($T_c - T_{cw}$) ranged from 20 K to zero, based on $T_c$ measured at a cooling rate of 0.5 K min⁻¹. The $R_f - t$ profiles again exhibit the three stages mentioned above. The initial step
increase is larger and the final $R_f$ value higher with increased subcooling (lower $T_{cw}$),
as expected. It is noteworthy that an initial gel layer is formed at zero subcooling,
indicating that the cloud point calculated under quasi-static conditions does not map
directly to deposition tendency in these systems. For comparison, the initial cooling
rate in the paraffin in Figure 9 is of the order of 37 K min$^{-1}$, and the true cloud point
would thus be raised significantly by the rapid cooling rate.

The amount of deposit also increased with the degree of subcooling, and Figure
18(a) suggests a strongly linear relationship. This extrapolated locus continues beyond
$T_c$, and suggests that no deposition would occur when $T_{cw} \sim 40$ °C. Tests with $T_{cw} > T_c$
were not performed here. Another factor which may play a role is the presence of the
stainless steel surface, which could promote monolayer formation (and heterogeneous
nucleation and growth) at temperatures above the bulk cloud point (for homogeneous
nucleation).

The linear fouling stage was shorter as $T_{cw}$ increased, which is consistent with less
time elapsing before the transition temperature – again, around 46.1 °C – was reached.
The linear fouling rates were similar to those reported above, ranging from $1.04 \times 10^{-6}$
to $1.21 \times 10^{-6}$ m$^2$ K J$^{-1}$. Falling rate fouling could again be described by Equation (20),
with $K$ values of $1 \sim 2 \times 10^{-5}$ μm s$^{-1}$ K$^{-1}$.

The deposit composition data in Figure 18(b) again show little effect of
subcooling on PPP concentration in the liquid entrained in the deposit. The values are
consistently larger than the bulk concentration, but the uncertainty in the measurement
increases with reduced subscooling. The solids mass fraction data exhibit a twofold
difference over the $T_{cw}$ range studied, increasing with lower initial subcooling (and
most of the growth occurring via normal crystallisation). Visual inspection of the
deposits indicated that for this fixed shear rate, those formed under conditions of
smaller subcooling were firmer. The deposits formed under high subcooling
conditions (and faster initial rate) appear to incorporate more solution, which could
either be due to a growth occurring via a dendritic front or the effect of higher liquid
viscosity at lower temperature. Singh et al. (2001a) also reported higher liquid
contents in wax deposit formation at higher subcoolings.

3.9. Deposit rheology

The strength of the gels formed was investigated by oscillatory rheometry. Shear
stress sweeps were performed at a frequency of 1.59 Hz. The results obtained for
many of the deposits obtained from the fouling tests described above are presented in Figure 19. The rheological parameters extracted are those reported by Nigo et al. (2009): the elastic modulus in the linear elastic region, $G'$, the limit of linearity, $\tau_{EL}$, defined as the end of the linear elastic region, where $G'$ deviated more than 5% from its initial value, and $\tau_C$, the crossover point of elastic and viscous moduli. It is noteworthy that all the $\tau_C$ values are > 2 Pa, the maximum surface shear stress estimated in the SDA tests reported here. The Figure shows that all the parameters increased with solids volume fraction in a strongly non-linear manner, and there is good overlap with the data reported by Nigo et al. (2009) for similar materials.

As a particulate soft solid, the rheological parameters are expected to be strongly determined by the solid volume fraction in the gel, $\phi$. The data exhibit power-law behaviour and could be fitted to the form:

$$ Y = b\phi^n $$

where $Y$ is the rheological parameter and $b$ and $n$ are positive constants. The gels are then consistent with the fractal model (Tang and Marangoni, 2007), where $b$ and $n$ are constants related to structural characteristics of the fat crystal network. In that model, the structure of a colloidal network is considered as a collection of fractal flocs, the increase of $G'$, $\tau_{EL}$ and $\tau_C$ with $\phi$ can be explained by the increase in the number of interfloc links with solid fat content (Marangoni and Rogers, 2003; Shih et al., 1990; Venkatesan et al., 2005). The results show that the gel quickly developed an appreciable strength, so that it would resist the shear imposed by the fluid over the surface. However, it was not possible here to determine the local solids distribution: this will be investigated using freezing techniques.
4. Conclusions

The SDA allows one to investigate the heat and mass transfer characteristics of gel formation from model food fat solutions on cold surfaces under well controlled flow conditions. Experimental investigations of the SDA heat transfer performance were combined with CFD simulation studies to generate maps of the temperature and shear stress conditions on the SDA test surface. The results confirmed that the thermal resistance across the disc was small and heat transfer was controlled by the bulk fluid, however, the internal thermal resistances became significant at higher Re.

A series of laboratory experiments were carried out in the SDA with 5 wt% PPP in a non-crystallising solvent to investigate the effects of varying deposition time, shear and thermal driving force on the crystallisation fouling behaviour of food fats. $R_f - t$ profiles for these ‘cold start’ experiments exhibited three stages: transient, linear rate and falling rate stage. $R_f - t$ profiles could be generated reliably and reproducibly, indicating that the heat flux measurements provide a reliable monitor of deposit growth. In all cases, the $H_X$ contribution in $H_T$ is noticeably small, the estimated $T_s$ increased quickly initially and then levelled off to approach the experimentally measured $T_m$.

The amount of gel formed initially could be related to the subcooling of the cold disc. The linear fouling rate was relatively insensitive to temperature and velocity effects and the rate controlling step is yet to be identified. In the falling rate stage, the observed dependence of $G(T)$ on $\Delta T$ was linear, regardless of the operating conditions, indicating that a normal growth stage controls fouling at these temperatures.

For given values of $\omega_d$ and $T_{cw}$, $m$ and $w_X$ increased steadily with $t$, indicating that deposit ageing occurred in the gels over the time scale of the experiments. Ageing did not change the deposit properties in terms of $\rho_f k_f$. The strength of the gel formed, whether described by $G'$, $\tau_{EL}$ or $\tau_C$, increased with solids content and could be related to a fractal model of deposit microstructure.

At the shear and temperature conditions employed in this study, deposition is driven by formation of gels on the heat transfer surface. The composition and structure of the gels reflected the conditions present at their formation. Increasing the amount of shear generally decreased $m$ and increased $w_X$, while decreasing the coolant temperature (and increasing the degree of subcooling) increased $m$ and decreased $w_X$. Spalling was not observed under the moderate shear stress conditions studied here.
Nomenclature

Roman

\(a\) constant in Equation 5

\(b\) constant in Equation 21

\(C_p\) deposit heat capacity, J kg\(^{-1}\) K\(^{-1}\)

\(D\) diffusivity of solute in solvent, m\(^2\) s\(^{-1}\)

\(G\) growth rate of deposit thickness, \(\mu m\) s\(^{-1}\)

\(G^*\) elastic modulus in linear region, Pa

\(h_b\) film heat transfer coefficient in bulk fluid, W m\(^{-2}\) K\(^{-1}\)

\(H_O\) combined contribution of \(H_X\) and \(H_T\), J or kJ

\(H_S\) sensible heat associated with the deposit, J or kJ

\(H_T\) total heat energy transferred during fouling run, J or kJ

\(H_X\) latent heat of PPP crystallisation, J or kJ

\(\Delta H_{fp}\) melting enthalpy of PPP, J mol\(^{-1}\)

\(k_b\) liquid thermal conductivity, W m\(^{-1}\) K\(^{-1}\)

\(k_d\) deposit thermal conductivity, W m\(^{-1}\) K\(^{-1}\)

\(k_m\) film mass transfer coefficient of bulk solution, m s\(^{-1}\)

\(K\) constant in Equation 20, \(\mu m\) s\(^{-1}\) K\(^{-1}\)

\(m\) mass of deposit per unit disc surface, kg m\(^{-2}\)

\(n\) constant in Equation 21

\(N_d\) deposition rate, kg m\(^{-2}\) s\(^{-1}\)

\(N_{PPP}\) mass flux of PPP towards surface, kg m\(^{-2}\) s\(^{-1}\)

\(Nu\) Nusselt number, -

\(Nu^*\) modified Nusselt number, -

\(Pr\) Prandtl number, -

\(q\) heat flux, W m\(^{-2}\)

\(r\) radial position, m

\(R_d\) disc radius, m

\(R\) universal gas constant, J kg\(^{-1}\) K\(^{-1}\)

\(R_{cw}\) thermal resistance of coolant side, m\(^2\) K W\(^{-1}\)

\(R_f\) fouling resistance, m\(^2\) K W\(^{-1}\)

\(R_{f^*}\) asymptotic fouling resistance, m\(^2\) K W\(^{-1}\)

\(R_w\) thermal resistance of disc wall, m\(^2\) K W\(^{-1}\)

\(R_{else}\) thermal resistance of coolant flow and plates, m\(^2\) K W\(^{-1}\)

\(Re_f\) Reynolds number of bulk solution, -

\(Sc\) Schmidt number, -

\(t\) deposition time, s, min or hr

\(t^*\) end time of linear fouling rate regime, s
$t_F$ characteristic time, s, min or hr
$T^*$ surface temperature at the end of linear fouling rate region, °C or K
$T_b$ bulk temperature, °C or K
$T_c$ cloud point, °C or K
$T_{cw}$ coolant temperature, °C or K
$T_m$ melting point, °C or K
$T_w^p$ melting point of pure PPP, °C or K
$T_s$ deposit surface temperature, °C or K
$T_w$ temperature at base plate/deposit interface, °C or K
$\Delta T$ difference between $T_m$ and $T_s$, K
$U$ overall heat transfer coefficient, W m$^{-2}$ K$^{-1}$
$U_0$ initial value of $U$, W m$^{-2}$ K$^{-1}$
$w_X$ solids content in deposit, wt%
$w_S$ PPP content in the solution entrained in deposit, wt%
$w_T$ total PPP content in deposit, wt%
$x_b$ PPP concentration in bulk solution, kg m$^{-3}$
$x_s$ PPP concentration at surface, kg m$^{-3}$
$X_{PPP}$ mole fraction of PPP in solution, -
$Y$ rheological parameters $G'$, $\tau_C$ and $\tau_{EL}$

Greek
$\delta_f$ deposit thickness, μm
$\mu_b$ apparent viscosity of bulk fluid, kg m$^{-1}$ s$^{-1}$
$\rho_b$ density of bulk fluid, kg m$^{-3}$
$\rho_f$ deposit density, kg m$^{-3}$
$\tau_C$ crossover point of elastic and viscous modulus, Pa
$\tau_{EL}$ the end point of the linear region, Pa
$\phi$ solids volume fraction of deposit, %
$\omega_d$ can rotational speed, rad s$^{-1}$

Abbreviations
CFD computational fluid dynamics
DSC differential scanning calorimeter
NTU number of turbidity units
PPP tripalmitin
SDA scanning disc apparatus
References


A spinning disc study of fouling of cold heat transfer surfaces by gel formation from model food fat solutions

Huang, J.Y., Chew, Y.M.J. and Wilson, D.I.

Tables and Figures

Table 1. Experimental conditions employed in heat transfer experiments

<table>
<thead>
<tr>
<th></th>
<th>30 wt% Water/Glycerol</th>
<th>78 wt% Water/Glycerol</th>
<th>Paraffin¹</th>
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<td>Coolant temperature, ( T_{cw} ) (°C)</td>
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<td>10.6</td>
<td>10.6</td>
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<tr>
<td>Bulk temperature, ( T_b ) (°C)</td>
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<td>35.1 – 60.7</td>
<td>35.9 – 61.4</td>
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<td>Dynamic viscosity, ( \mu_b ) (Pa s)</td>
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<td>0.01 – 0.03</td>
<td>0.03 – 0.08</td>
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<tr>
<td>Reynolds number, ( Re_i )</td>
<td>303 – 5443</td>
<td>47 – 307</td>
<td>11 – 48</td>
</tr>
</tbody>
</table>

¹Density at 25 °C, 870 kg/m³, and viscosity 0.166 Pa s at 20 °C; BDH Chemicals, UK
Figure captions

Figure 1. (a) Schematic of the operation of the SDA, (b) construction of the fouling cell plate. Dimensions are in millimeters (not to scale).

Figure 2. Effect of composition and cooling rate on model solution $T_m$ and $T_c$. Loci drawn through the $T_m$ data show regression fit to ideal solution, Equation (2). Lines drawn through $T_c$ data sets as a guide to the eye.

Figure 3. Evolution of viscosity of 5 wt% PPP solutions while cooling in the rheometer at 5 K min$^{-1}$. Constant shear stress, 2 Pa. Regions A-D separated by vertical dashed lines discussed in text.

Figure 4. Effect of $\omega_d$ on $Nu^*$ for different test solutions. Symbols: solid – 30 wt% glycerol-water solutions grey – 78 wt% glycerol-water solutions, open – paraffin. $T_b = 50 ^\circ C$; (triangles) – $T_{cw} = 10.6 ^\circ C$, (squares) $T_{cw} = 10.6 ^\circ C$, (diamonds) $T_{cw} = 10.7 ^\circ C$, (circles) $T_{cw} = 10.7 ^\circ C$. Locus shows trend described by Equation (5) with $a = 0.50$ (Sparrow & Gregg, 1959)

Figure 5. Flow patterns and temperature profiles in the SDA for disc rotation speed of 5.4 rad s$^{-1}$, $Re_r = 27$. Black arrows are velocity vectors. Colour indicates temperature, dark red = 60 $^\circ C$ ($T_b$), dark blue = 9.8 $^\circ C$ ($T_{cw}$).

Figure 6. Effect of $Re_r$ on the thermal resistances in the SDA: circle – $1/U$, triangle – $1/h_b$ from simulation, square – $R_{cw}$ and line – $R_w$. Temperatures as in Figure 5.

Figure 7. (a) Temperature and (b) average shear stress on the disc surface at the radial location from 0 to 0.035 m, extracted from CFD simulations for various $T_c$ and $\omega_d$.

Figure 8. (a) $R_f$ and (b) estimated $T_s$ and $T_w$ during fouling of 5 wt% PPP solutions at $T_{cw} = 9.8 ^\circ C$, $\omega_d = 5.4$ rad s$^{-1}$. Vertical dashed line in (a) separates linear rate stage from the falling rate stage. Horizontal dashed lines in (b) show salient temperatures for the 5 wt% solution used.

Figure 9. Initial transients for tests reported in Figure 8 and for solvent alone.

Figure 10. Correlation of $R_f$ and $m$ for fouling runs in Figures 8, 15 and 17.

Figure 11. Evolution of deposit composition for deposits in Figure 8.

Figure 12. Comparison of cumulative $H_T$, $H_O$ (= $H_X + H_S$) and $H_X$ for fouling runs in Figures 8.

Figure 13. Comparison of deposition rates estimated from $R_f$ data and maximum rate of PPP mass transfer for 12 h fouling experiment in Figure 8.

Figure 14. Growth rate of $\delta_f$ in the falling rate stage for fouling runs in Figures 8 showing linear dependence on $\Delta T$. Line shows fit to linear trend.

Figure 15. Effect of $\omega_d$ on fouling behaviour. Conditions: $T_{cw} = 9.8 ^\circ C$, $t = 12$ h. $T^*$ and $t^*$ are
end temperature and time of linear rate stage respectively.

**Figure 16.** Effect of $\omega_d$ on $m$ and deposit composition for the fouling runs for the fouling runs in Figure 15.

**Figure 17.** Effect of $T_{cw}$ on (a) fouling behaviour and (b) $T_s$ profile. Conditions: $\omega_d = 5.4$ rad s$^{-1}$ and $t = 12$ hr.

**Figure 18.** Effect of $T_{cw}$ on deposit (a) mass and (b) composition, for the fouling runs for the fouling runs in Figure 17. Vertical lines in (a) indicate the $T_c$ and $T_m$.

**Figure 19.** Effect of deposit solids volume fraction on rheological parameters measured at 20 °C. Solid data sets are those reported for similar PPP solutions by Nigo et al. (2009). Loci indicate power law trend lines (Equation (21)) fitted to the data sets: $G' = 9 \times 10^7 \psi^{3.7}$; $\tau_{EL} = 2.9 \times 10^5 \psi^{4.8}$ and $\tau_C = 2.4 \times 10^5 \psi^{2.8}$. 
Figure 1. (a) Schematic of the operation of the SDA, (b) construction of the fouling cell plate. Dimensions are in millimeters (not to scale).
Figure 2. Effect of composition and cooling rate on model solution $T_m$ and $T_c$. Loci drawn through the $T_m$ data show regression fit to ideal solution, Equation (2). Lines drawn through $T_c$ data sets as a guide to the eye.
Figure 3. Evolution of viscosity of 5 wt% PPP solutions while cooling in the rheometer at 5 K min\(^{-1}\). Constant shear stress, 2 Pa. Regions A-D separated by vertical dashed lines discussed in text.
Figure 4. Effect of $\omega_d$ on $Nu^*$ for different test solutions. Symbols: solid – 30 wt% glycerol-water solutions grey – 78 wt% glycerol-water solutions, open – paraffin. $T_b = 50 \degree C$; (triangles) $T_{cw} = 10.6 \degree C$, (squares) $T_{cw} = 10.6 \degree C$, (diamonds) $T_{cw} = 10.7 \degree C$, (circles) $T_{cw} = 10.7 \degree C$. Locus shows trend described by Equation (5) with $a = 0.50$ (Sparrow & Gregg, 1959).
Figure 5. Flow patterns and temperature profiles in the SDA for disc rotation speed of 5.4 rad s$^{-1}$, $Re_r = 27$. Black arrows are velocity vectors. Colour indicates temperature, dark red = 60 °C ($T_b$), dark blue = 9.8 °C ($T_{cw}$).
Figure 6. Effect of \( Re_r \) on the thermal resistances in the SDA: circle – \( 1/U \), triangle – \( 1/h_b \) from simulation, square – \( R_{cw} \) and line – \( R_w \). Temperatures as in Figure 5.
Figure 7. (a) Temperature and (b) average shear stress on the disc surface at the radial location from 0 to 0.035 m, extracted from CFD simulations for various $T_{cw}$ and $\omega_d$. 
Figure 8. (a) $R_t$ and (b) estimated $T_s$ and $T_w$ during fouling of 5 wt% PPP solutions at $T_{cw} = 9.8 \, ^\circ\text{C}$, $\omega_d = 5.4 \, \text{rad s}^{-1}$. Vertical dashed line in (a) separates linear rate stage from the falling rate stage. Horizontal dashed lines in (b) show salient temperatures for the 5 wt% solution used.
Figure 9. Initial transients for tests reported in Figure 8 and for solvent alone.
Figure 10. Correlation of $R_f$ and $m$ for fouling runs in Figures 8, 15 and 17.
Figure 11. Evolution of deposit composition for deposits in Figure 8.
Figure 12. Comparison of cumulative $H_T$, $H_O (= H_X + H_S)$ and $H_X$ for fouling runs in Figures 8.
Figure 13. Comparison of deposition rates estimated from $R_f$ data and maximum rate of PPP mass transfer for 12 h fouling experiment in Figure 8.
Figure 14. Growth rate of $\delta_t$ in the falling rate stage for fouling runs in Figures 8 showing linear dependence on $\Delta T$. Line shows fit to linear trend.
Figure 15. Effect of $\omega_d$ on fouling behaviour. Conditions: $T_{cw} = 9.8 \, ^\circ C$, $t = 12 \, h$.

$T^*$ and $t^*$ are end temperature and time of linear rate stage respectively.
Figure 16. Effect of $\omega_d$ on $m$ and deposit composition for the fouling runs for the fouling runs in Figure 15.
Figure 17. Effect of $T_{cw}$ on (a) fouling behaviour and (b) $T_s$ profile. Conditions: $\omega_d = 5.4 \text{ rad s}^{-1}$ and $t = 12 \text{ hr}$. 

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Figure 18. Effect of $T_{cw}$ on deposit (a) mass and (b) composition, for the fouling runs for the fouling runs in Figure 17. Vertical lines in (a) indicate the $T_c$ and $T_m$. 
Figure 19. Effect of deposit solids volume fraction on rheological parameters measured at 20 °C. Solid data sets are those reported for similar PPP solutions by Nigo et al. (2009). Loci indicate power law trend lines (Equation (21)) fitted to the data sets: $G' = 9 \times 10^7 \phi^{3.7}$, $\tau_{EL} = 2.9 \times 10^5 \phi^{4.8}$ and $\tau_C = 2.4 \times 10^5 \phi^{2.8}$. 