

Deposit layer formation during skim milk dead-end filtration with ceramic hollow fiber membranes using magnetic resonance imaging

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A dead-end filtration process of skim milk with ceramic hollow fiber membranes was investigated using *in situ* Magnetic Resonance Imaging. Special emphasis was put on contrast optimization exploiting water relaxation times being sensitive to casein concentration in the formed deposit layer.

Introduction

Membrane filtration is known to be a major process in liquid food processing. The main applications of membranes in this field are in dairy, followed by beverage industry.¹ An important application of membrane filtration processes in the dairy industry is the separation of casein micelles and whey proteins from milk by means of microfiltration.² The resulting protein fractions are in their native states and have different nutritive and techno-functional properties. Hence, they are added to a wide range of food products. The challenge during the filtration is to completely retain casein micelles with a diameter between 50 and 400 nm ($d_{50,3} = 180$ nm) while maximizing the permeation of whey proteins with 2–4 nm ($d_{50,3} = 3$ nm). Although the pores of ordinary microfiltration membranes are by far bigger than whey proteins, the permeation is not 100% but only around 50% so far. This is mainly due to the casein micelles, which are retained and form a deposit layer that acts as a secondary layer on the membrane's surface. The consequence is a reduced permeate flux and protein permeation with increasing filtration time.³

Typically, the filtration flux is used to characterize the filtration performance. However, by using e.g. resistance in series models, only global filtration resistances for pore blocking and cake layer formation phenomena are obtained.^{4,5} These models enable the study of fouling phenomena. Nevertheless, they do not take into account, that the transmembrane pressure (TMP) decreases along the membrane due to friction forces in crossflow filtration processes which lead to an axial dependence of the filtration flux and permeation characteristics.^{2,6} Studies dealing with the length dependence of flux and deposit layer formation show diverging results. Gernedel (1980)⁷ justified the declining flux along the membrane with the axial transportation of already deposited material towards the outlet of the membrane and therefore a thicker deposit layer. Based on the results of a sectioned membrane recent studies assume a thicker deposit layer at the inlet of the membrane due to a higher TMP and thus a lower permeation. However, the drawback so far is that all

studies use indirect and off-line methods to measure the deposit layer. The actual thickness of the fouling layer during the filtration is not known.² Complementary characterisation techniques are needed to gain more information about these spatially varying fouling phenomena.

Nuclear magnetic resonance imaging (MRI) is known to be a powerful technique for non-invasive observation of membrane processes.⁸⁻¹⁰ Recent examples for the application of MRI in membrane filtration research are the investigation of the fouling behaviour of spiral wound modules, the influence of feed spacers on the velocity distribution and biofouling of these modules.¹¹⁻¹⁵ MRI was also applied to measure flow, concentration polarization and cake layer formation in hollow fiber geometries.^{8, 16-19} Typically, colloidal particle suspensions^{16, 20, 21} or oil-water emulsions^{18, 22} were used as model substances, exploiting the relaxation contrast in the images.

In this work the question is answered, whether a protein deposit layer on the membrane's surface can be detected based on different relaxation times of the retained substances and the feed medium. Therefore, the transverse and longitudinal relaxation were measured as a function of casein concentration, which is the main retained component during skim milk filtration. Based on these results, *in situ* MRI during a dead-end filtration was performed in order to detect the casein deposit layer on the membrane's inner surface.

Material and methods

In situ filtration set up

The experiments were performed in a 200 MHz MRI tomograph (Bruker Avance 200 SWB, Bruker BioSpin GmbH, Rheinstetten, Germany). Figure 1 shows the flow diagram of the *in situ* filtration set up.

The ceramic hollow fiber module is mounted into a ¹H-NMR bird cage (10 mm inner diameter) and installed in the tomograph. For the experiments the hollow fiber membrane is fixed in a glass module and connected via hose connections to the feed source and

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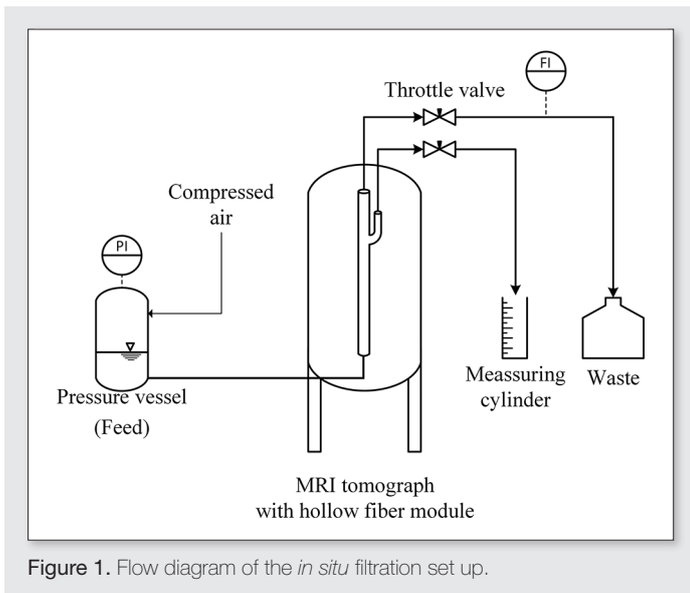


Figure 1. Flow diagram of the *in situ* filtration set up.

the collecting vessels. A pressure vessel is used for feed transport to avoid pulsation due to pumping and consequently to ensure an artefact-free measurement. Filtrations were performed with a constant transmembrane pressure of 100 kPa. Dead-end filtration rather than cross-flow filtration was chosen in order to realize a fast build-up of a deposit layer on the membrane's surface. The permeate was collected in a measuring cylinder volume to determine V_{Perm} over time t . Knowing the active membrane area A_{Mem} , the permeate flux J_p can be calculated according to:

$$J_p = \frac{1}{A_{Mem}} \frac{dV_{Perm}}{dt} \quad (1)$$

Ceramic hollow fiber membranes

Ceramic hollow fiber membranes were used due to their high chemical and thermal stability as well as high specific membrane surface. The ultrafiltration membranes have a nominal pore diameter of 40 nm. They consist of α - Al_2O_3 and have a radially asymmetric structure: A microporous support layer provides the mechanical stability, and a thin active layer on the membrane's feed facing surface determines the separation properties (Figure 1). Due to the localization of the active layer on the membrane's inner surface, the filtration mode is from the inside to the outside (in/out filtration). The membrane had an active filtration length of 445 mm, an inner diameter of 1.74 mm and a clean water flux of 350 l/m²/h (measured at TMP of 100 kPa).

Production of milk samples

To avoid thermally induced interactions between serum proteins and casein micelles raw milk was collected directly from a local farm. The milk was separated with a pilot cream separator type MM 1254 D (GEA Westfalia, Oelde, Germany) at 8,000 g and 50 °C to obtain a skim milk with a fat content below 0.1%. The skim milk was divided into two batches. The first batch was used for the dead-end filtration directly whereas the second fraction was processed further to obtain a whey protein free casein concentrate. The casein micelles were fractionated from the whey proteins by means of microfiltration operated in diafiltration mode. It was coupled with an ultrafiltration membrane (cut off 10 kDa) according to Toro-Sierra

et al.²³ For microfiltration different conditions were used: It was operated in crossflow mode with ISOFLUX[®] membranes with a nominal cut-off of 0.14 μ m and a membrane area of 2.45 m², consisting of seven ceramic modules in form of multichannel elements (1178 mm length, 3.5 mm diameter per channel, 23 channels, TAMI Industries, France). After concentrating the skim milk by a factor of two the same amount of milk and ideal whey (microfiltration permeate) were collected in the receiving tank of the microfiltration and ultrafiltration, respectively. Seven diafiltration steps were carried out at constant filling level by transferring the permeate of the ultrafiltration back to the receiving tank of the microfiltration to keep the natural milk milieu and avoid changes of the casein micelles. In this way, the whey protein concentration in the microfiltration retentate was reduced by 97.2%. The final protein content of the casein concentrate was adjusted to 9.0% by concentration and spray dried (Spray Dryer B-290, Büchi, Switzerland) to a dry matter of 96.2%. Casein suspensions were prepared by dissolving the proper amount of spray dried casein in ultra-pure water.

T_1 and T_2 determination

In order to measure T_1 and T_2 skim milk and the casein samples were filled into 5 mm NMR sample tubes and placed in the tomograph altogether. MSME with varying inversion times was applied to measure the magnetization recovery. During analysis of signal intensities T_1 relaxation times were determined by fitting an exponential growth function to the measured data points:

$$\frac{M(t)}{M_0} = 1 - \exp\left(-\frac{t}{T_1}\right) \quad (2)$$

$M(t)$ is the magnetization at time t , M_0 is the equilibrium magnetization and T_1 the longitudinal relaxation time. Transverse relaxation times T_2 were determined by measuring the samples in a multi echo sequence at different echo times ($t = n \cdot T_E$) at a fixed repetition time. T_2 was obtained using a fit of an exponential decay:

$$\frac{M(t)}{M_0} = \exp\left(-\frac{t}{T_2}\right) \quad (3)$$

Results and discussion

Influence of casein concentration on relaxation properties

Figure 2a) displays the transverse relaxation times of casein suspensions with four different casein concentrations at two different repetition times. An increasing casein concentration leads to a faster signal decay of the transverse magnetization. As expected, the variation of TR had no significant influence on T_2 . Casein not only affects T_2 but also leads to a decrease of T_1 . Casein causes the longitudinal and the transverse magnetization of the suspension to relax faster compared to the relaxation of pure water. These findings are in good agreement with the observations described in Ref. 24, 25. In the context of MRI during filtration processes of milk, this fact allows the exploration of relaxation contrast to detect the deposit layers formation.

In situ MRI for observation of layer formation

During dead-end filtration of skim milk, the retained feed fractions accumulate at the membrane and form a deposit layer. Due to the large mean diameter of casein micelles ($d_{50,3} = 180$ nm), they will be rejected by the hollow fiber membrane with a pore size of 40 nm.

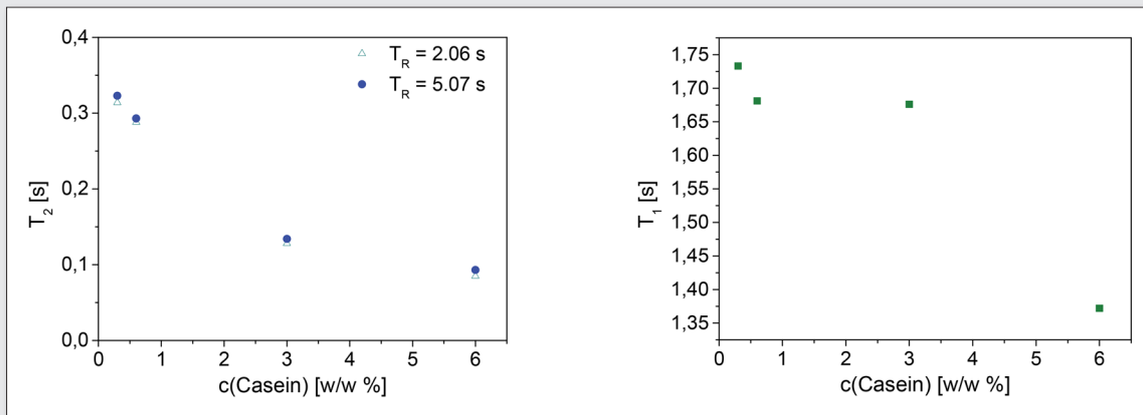


Figure 2. Both relaxation times T_2 and T_1 of a casein suspension depend on the casein concentration. The relaxation times decrease with increasing concentration and are therefore good measures for contrast generation in MRI.

The consequence is an accumulation of casein micelles in the deposit layer and hence a much higher concentration than in the skim milk feed solution. The effect on relaxation time in the region of the deposit layer is therefore even more pronounced as in the relaxation studies (Figure 2). When setting up an MRI experiment knowing that both relaxation times are influenced by the casein concentration, the contrast in the images has to be considered. A reduction of T_1 is equivalent with a signal enhancement, i.e. a positive contrast in T_1 weighted images, whereas a reduction in T_2 leads to a signal reduction, i.e. a negative contrast in echo based MRI sequences. Thus, with a shorter repetition time TR a stronger T_1 weighting can be achieved. Vice versa, an increase of the echo time leads to a more pronounced differentiation along T_2 . An increase, a reduction or even an inversion of the image contrast can be obtained depending on the pulse sequence's timing. Good results were obtained at an average repetition time of $TR = 800$ ms and the minimum echo time (Figure 3) when applying Rapid Acquisition with Relaxation Enhancement (RARE) MRI. The inner circle in the axial image of the unfouled membrane (Figure 3a) represents the feed

channel, followed by the membrane itself. The membrane shows a low intensity and therefore appears blue in the false colour images. This is not only due to the lower proton density in the membrane but also due to the short transverse relaxation times of molecules in the pores of the membrane. The observation of a brighter circle of higher intensity roughly in the middle of the membrane affirms this interpretation. This circle is caused by the water molecules located in the larger pores of the supporting layer (Figure 3f). The pore structure is not uniform over the cross section of the hollow fiber membrane. This is attributed to the production process and is in good agreement with the μ CT image in Figure 3f). Furthermore, a quite thin ring with high intensities can be identified at the inner surface of the ceramic hollow fiber membrane prior to filtration (see Figure 3). This phenomenon could result from magnetic susceptibility differences of the ceramic hollow fiber and the surrounding skim milk²⁶ but also from adsorption of proteins to the membrane surface.

During dead-end filtration of skim milk the deposit layer formation was measured as a function of time. In previous studies, RARE imaging has turned out to be feasible for time resolved and almost

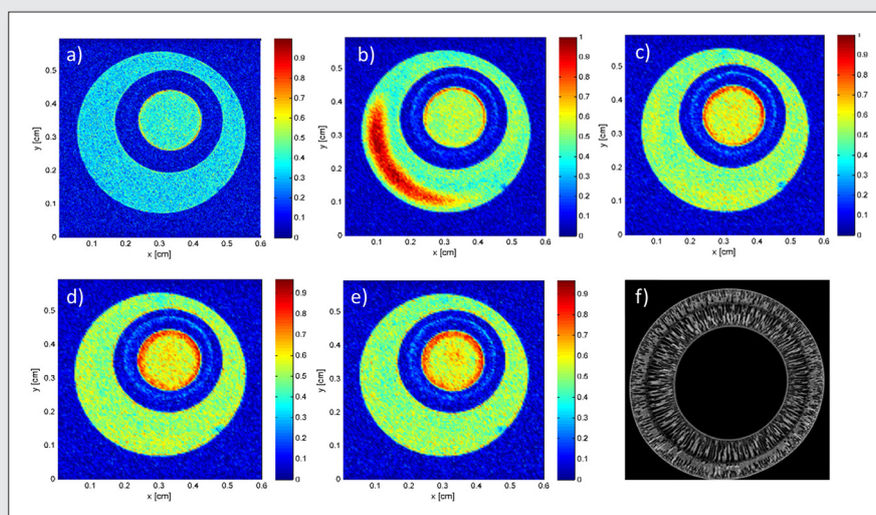


Figure 3. RARE-images of skimmed milk dead-end filtration at 1 bar TMP. a) before start, b) 7 min, c) 27 min, d) 39 min, e) 46 min – Acquisition time: 2:08 min, $T_R=800$ ms, $\tau_E=3.709$ ms, FOV: 6x6 mm, Matrix: 128x128, RARE Factor: 8. Please note, that the images are individually scaled to the maximum intensity of each image, indicated by the corresponding colorbar. f) μ CT image of a ceramic hollow fiber membrane.^{26, 27}

artefact free *in situ* monitoring of filtration. Five axial slices were measured, each having a thickness of 1 mm. The results for the first of the five axial slices at different filtration times can be seen in Figure 3. With increasing filtration time, an area with high intensities is detected on the membrane's surface. This area corresponds to the formed deposit layer with increased concentration of T_1 -shortening substances. Additionally, a region of high intensity in the lower left section of the image is recognizable (Figure 3b). This is likely due to flow artefacts resulting from the initial high flow in the permeate channel at the beginning of the filtration corresponding to the high filtration flux.

Furthermore it can be seen, that the deposit layer is not uniform. In the upper region of the membrane the deposit layer is much more pronounced than in the lower region. Simultaneously, the pore structure seems to be more open in the upper part of the membrane leading to a lower local membrane resistance. This in turn leads to a higher flow and hence more convective transport of casein micelles towards the membrane's surface resulting in a higher deposit layer.

The RARE images were quantitatively analysed as a function of filtration time. The thickness of the fouling layer was determined by a self-written MATLAB script, exploring the symmetry and the uniformity of the layer at the inner side of the hollow fiber membrane. Two ellipses, one describing the inner membrane's surface and one describing the interface of the fouling layer to the filtrate channel are defined interactively. The flexibility of this procedure takes the fabrication errors into account as well as the fact that an automated segmentation is difficult due to the varying image contrasts. Figure 4 displays the development of the deposit layer height as a function of filtration time.

The thickness of the deposit layer increases significantly at the beginning of the filtration process. This is due to the higher flux during in the beginning of dead-end filtration, which leads to a high convective transport of casein micelles to the membrane's surface. After this initial phase, the build-up of the deposit layer leads to an increased filtration resistance and therefore to a decrease of the filtration flux at constant pressure. The consequence is that the

deposit layer increases and the filtration flux slows down. During dead-end filtration experiments it has to be considered, that not only casein micelles are rejected, but also whey proteins are partly embedded in the deposit layer, also leading to an alteration of T_1 and T_2 in this region. Regarding the extremely low values for the filtrate flux, it is worth to note, that on an industrial scale the fractionation of casein micelles and whey proteins is performed in crossflow filtration mode. The higher crossflow velocity leads to shear forces at the membrane's surface. These cause an at least partial removal of the deposit layer and therefore lead to a lower filtration resistance resulting in much higher filtrate fluxes. The removal of the deposit layer is very important, because the permeation of the whey proteins is reduced by the deposit layer, too.² The objective of this article was to show that it is possible to measure the height of the fouling layer *in-situ* during the filtration of skim milk by means of MRI.

Conclusion

An increase of casein concentration results in a decrease of the transverse and longitudinal relaxation times of the aqueous suspension. Therefore it is possible to differentiate areas with high casein concentrations, e.g. in the fouling layer, from the feed solution with relaxation weighted MRI. The fouling layer was monitored time resolved during build up, and the thickness of the fouling layer was analysed as a function of duration of dead-end filtration of skim milk. Relaxation weighted MRI is shown to be a powerful tool to elucidate the underlying mechanism of length dependent deposit layer formation during the fractionation of skim milk and hence optimize filtration processes to avoid fouling. Further investigations will be performed regarding the application of MRI in cross flow filtration and the length dependence as well as the composition of the resulting deposit layer.

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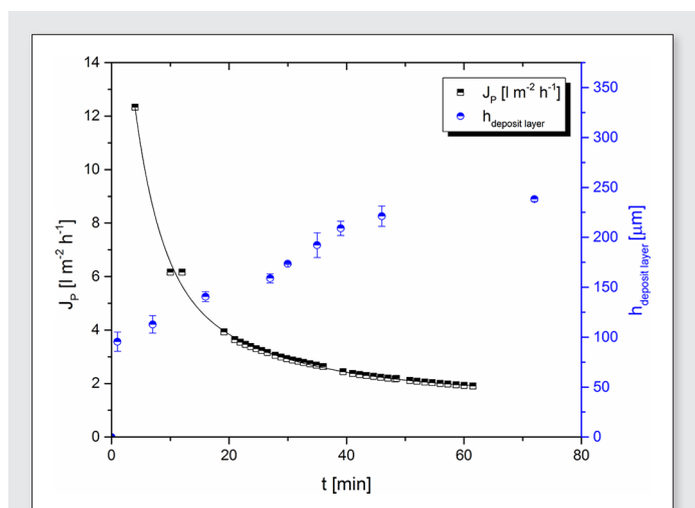


Figure 4. The flux J_p decreases as a function of filtration time, where J_p was measured integrally at the outlet of the module. In the same time, the height of the deposit layer increases as quantified from MRI images as shown in Figure 3.

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