

In-Situ Characterization of Deposits in Ceramic Hollow Fiber Membranes by Compressed Sensing RARE-MRI

S. Schuhmann, N. Schork, K. Beller, and H. Nirschl

Karlsruhe Institute of Technology (KIT), Institute of Mechanical Process Engineering and Mechanics, Karlsruhe 76131, Germany

T. Oerther

Bruker Biospin GmbH, Rheinstetten 76287, Germany

G. Guthausen

Karlsruhe Institute of Technology (KIT), Institute of Mechanical Process Engineering and Mechanics, Karlsruhe 76131, Germany

Karlsruhe Institute of Technology (KIT), Chair of Water Chemistry and Water Technology, Karlsruhe 76131, Germany

Ultrafiltration with ceramic hollow fiber membranes was investigated by compressed sensing rapid acquisition relaxation enhancement (CS-RARE) magnetic resonance imaging (MRI) to characterize filtration mechanisms. Sodium alginate was used as a model substance for extracellular polymeric substances. Dependent on the concentration of divalent ions like Ca^{2+} in an aqueous alginate solution, the characteristics of the filtration change from concentration polarization to a gel layer. The fouling inside the membrane lumen could be measured by MRI with a CS-RARE pulse sequence. Contrast agents have been used to get an appropriate contrast between deposit and feed. The lumen was analyzed quantitatively by exploring the membrane's radial symmetry, and the resulting intensity could be modeled. Thus, different fouling mechanisms could be distinguished. CS-RARE-MRI was proven to be an appropriate in situ tool to quantitatively characterize the deposit formation during in-out filtration processes. The results were underlined by flux interruption experiments and length dependent studies, which make it possible to differentiate between gel layer or cake filtration and concentration polarization filtration processes.

Keywords: fouling, alginate, NMR imaging, ceramic hollow fiber membranes, compressed sensing

Introduction

Ceramic hollow fiber membranes (CHFMs) are used for ultrafiltration because of their resistance against high temperatures and chemicals. They have a wide range of applications and are used for example in food, pharmaceutical, and water industries. The CHFMs in this publication are operated in in out filtration mode, which leads to higher shear rates on the inside of the CHFMs compared to out in filtrations.¹ CHFMs can be used in two operation modes: dead end and cross flow filtration. The operation mode depends on the nature of feed and application. Dead end filtration is applied at low feed concentrations whereas cross flow filtrations are usually used for higher concentrations to realize higher shear forces on the deposits due to the parallel flow.² This article focuses on dead end filtrations.

Sodium alginate is often used as a model substance for extracellular polymeric substances, which makes experiments

comparable and reproducible. The filtration mechanism depends on the feed composition. Literature distinguishes between cake filtration, concentration polarization, and gel layer filtration.³ In the filtration of aqueous solutions of sodium alginate, concentration polarization was observed. The properties of the sodium alginate feed can be changed by adding divalent ions to the feed, for example, in form of CaCl_2 . Sodium alginate will then form a gel. This characteristic of sodium alginate allows to in situ study these two different filtration mechanisms by MRI noninvasively.

MRI is known in medical applications, but also explored in technical and scientific processes and filtration.⁴⁻¹⁷ In this article, MRI is used for the in situ, time resolved observation of the filtration progress in CHFMs lumen. Especially the time evolution and the nature of the deposit is of major interest. The MRI measurement time depends on the pulse sequence, the resolution and the desired signal to noise ratio of the images. Due to the time evolution of the filtration process, comparatively short measurement times are essential. Compressed sensing (CS) allows to reduce the measurement time drastically. Nearly the same image quality can be achieved by

Correspondence concerning this article should be addressed to S. Schuhmann at sebastian.schuhmann@kit.edu.

CS reconstruction,¹⁸ which permits the observation of the filtration's progress with an adequate time resolution.

The approach in the present article is to elucidate the filtration mechanisms in CHFMs dependent on the composition of the feed solution by CS RARE MRI. Two extremes were studied to develop a quantitative description of deposit formation for concentration polarization and gel layer formation with an adequate spatial and time resolution.

Experimental

Feed solutions with contrast agents

An aqueous sodium alginate solution (Alginic acid sodium salt from brown algae "medium viscosity" from Sigma Aldrich) was used as a model substance for extracellular polymeric substances. The concentration of sodium alginate was 200 mg/L. Sodium alginate is known to form gel structures in the presence of divalent ions like Ca^{2+} . In former experiments, loose polarization layers or gel layers could be observed.^{19–21} Ca^{2+} ions were added in form of CaCl_2 (Sigma Aldrich) with a concentration of $c_{\text{Ca}^{2+}} = 2\text{mmol/L}$. Due to the poor contrast between alginate and water a contrast agent (Magnetite Alginate 100 MagAlg100, also named MION)²² was added to the feed solution in concentrations of 1 mM and below. These MIONs show predominately a T_2 contrast, the regions with significant concentration of the MIONs will therefore appear with lower intensities in relaxation weighted MRI. Long term MRI experiments proved that MIONs in the used concentration have no influence on the filtration mechanism. Also the permeate flux did not change by adding MIONs to the feed solution. Additional experiments revealed the following behavior of MIONs which is essential in the interpretation of the MRI data: When the MIONs are dissolved in water, they tend to sediment. When MIONs were dissolved in an alginate solution, they diffuse together with the alginate without sedimentation. Dissolved in an alginate gel, the MIONs don't migrate out of the gel into the surrounding water. Thus, these MIONs can be used to increase the MRI contrast by paramagnetic relaxation enhancement.

Hollow fiber membrane

For the ultrafiltration in wastewater treatment, in out filtration hollow fiber membranes were used as provided by MAN N + HUMMEL GmbH, Ludwigsburg, Germany. The fibers with an inner lumen diameter of about 1.9 mm and an outer fiber diameter of 3.2 mm^{20,23} are made of porous Al_2O_3 ceramics. The hollow fiber membranes have a radially asymmetric structure showing different layers along the radius. The outer and highly porous layer is responsible for the mechanical stability and the form of the hollow fiber whereas the inner layer is responsible for the filtration characteristics. This inner active layer has an average pore diameter of 40 nm and thickness in the order of 20 μm . CHFMs have the ability to withstand high temperatures and are tolerant against aggressive chemical cleaning. A water permeability between 170 $\text{L}/(\text{m}^2 \text{ h bar})$ and 315 $\text{L}/(\text{m}^2 \text{ h bar})$ could be measured on the used ceramic hollow fiber membranes.

In situ filtration experiments

Filtration experiments were performed in a single fiber filtration setup (Figure 1). The filtration module was investigated using a Bruker Avance HDIII SWB 200 MHz spectrometer equipped for MRI.^{19–21} The probe was a 20 mm birdcage of

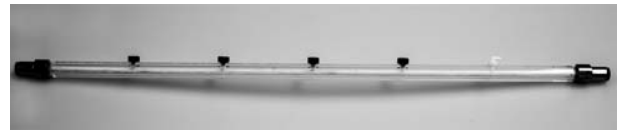


Figure 1. Single fiber filtration module which is mounted in the tomograph. Multiple permeate outlets are available, but only one is used in the experiments.

the MICWB40 series. This allows for an in situ observation of the membrane lumen during the filtration. Due to the module length of 420 mm, the module was observed in a first experiment only in a segment of 25 mm near the permeate outlet. Shifting the module during the filtration in a second experiment allows a limited but meaningful observation of the deposits along its length axis (z in the following) of the hollow fiber membrane. The feed solution was pumped into the module by a pressure vessel applying a transmembrane pressure of 1 bar leading to typical permeate volume flow rates of 1–5 mL/min which corresponds to laminar flow. By closing the valve of the retentate outlet, a dead end filtration can be performed. The permeate volume was measured volumetrically over time.

MRI methods

A fast spin echo method is needed for in situ filtration in CHFMs because of the inherent T_2 contrast which is generated by the contrast agent but also because of large susceptibility differences between liquids and ceramics.¹⁹ A RARE pulse sequence was used to measure the deposit formation in the membrane lumen while exploring the possibilities of CS. A series of 180° pulses after a 90° excitation pulse generates a series of echoes with different phase encodings but also with different transverse relaxation weighting. Multiples lines of k space are measured during a given repetition time T_R . The number of echoes which is collected by one T_R is named RARE factor RF .²⁴ RF has a significant influence on the images, conversely RARE is known to be sensitive to movements, that is, in our case to velocities in the feed lumen and in the permeate. Specifically, the inflow effect should be mentioned. Low RF and large T_R diminish this effect, which makes it hard to reduce the measurement time with commonly used NMR pulse sequence parameters. However, the time resolution of the MRI experiments has to be much smaller than the filtration progress. For example, at the beginning of the filtration, the feed velocities change very fast on the subminute time scale as the volume flow rate does. Short measurement times are therefore mandatory to detect the formation of the deposit layer spatially and time resolved. To reduce the measurement time without reinforcing the inflow effect or reducing the spatial resolution or signal to noise ratio, a CS RARE sampling pattern was used with 50% of the full phase encoding steps (Figures 2a, b).

A Box Müller method was applied to obtain the sampling scheme with a standard deviation $\sigma = 0.33$.²⁵ Fifty percent of the points were mainly sampled in the k space center along the phase encoding direction. The read direction (x direction) was sampled completely. The image reconstruction was performed with a l_1 penalized nonlinear conjugate gradient reconstruction solving the following optimization problem

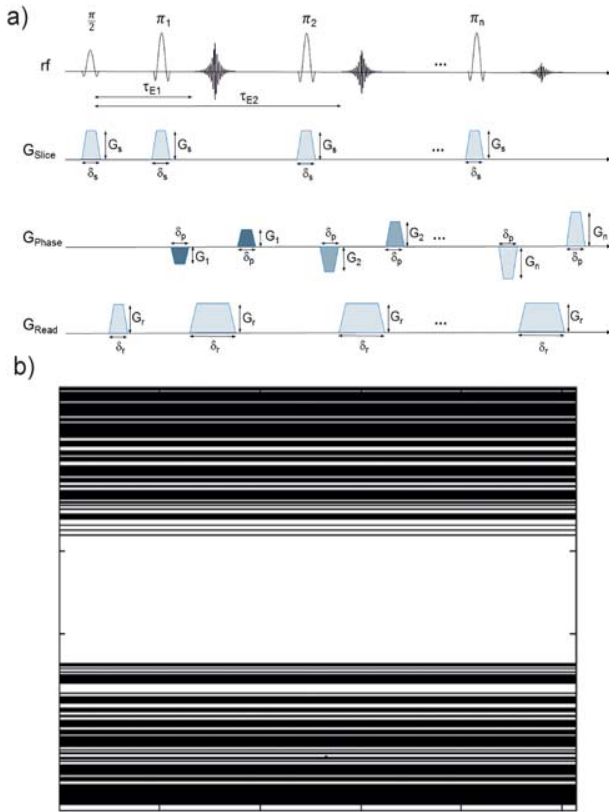


Figure 2. (a) RARE sequence and (b) CS-RARE sampling scheme: Only 50% of the phase encoding steps were measured (y -direction) to reduce the measurement time. All 256 points were measured in read direction (x -direction). The phase encoding steps were obtained by applying a Box-Müller sampling pattern.

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$$\operatorname{argmin}_S \left\{ \|FS - K\|_2 + \lambda_1 \|S\|_1 + \lambda_2 TV(S) \right\} \quad (1)$$

F is the Fourier operator applied to the image S , K describes the measured data in k space, $\lambda_{1,2}$ are the regularization parameters, and TV is the Total Variation. The reconstruction was performed with a modified MATLAB script available in Ref. 26. Due to the CS sampling pattern the measurement time was reduced to 3 min 12 s with pulse sequence parameters summarized in Table 1.

Quantification of the deposit layers

To characterize the fouling layer, the cylindrical geometry of the hollow fiber membrane was explored. Concentric rings

Table 1. MRI Parameters of the CS-RARE Pulse Sequence

MRI Parameter	Value
T_R	3 s
$\tau_{E, \text{effective}}$	5.8 ms
Field of view	8×8 mm
slice thickness	2 mm
no. pixels	256×256
in plane resolution	$31.25 \mu\text{m}$
RARE factor	2
CS undersampling	50%
measurement time	3 min 12s

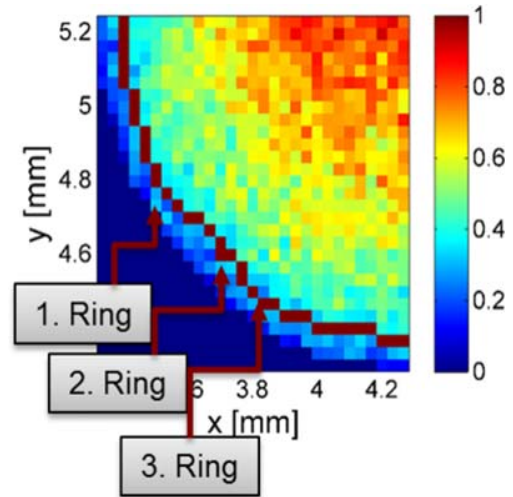


Figure 3. Scheme for determination of the radial intensity distribution: starting from the membrane wall an arbitrarily shaped ring is identified by the intensity jump between wall and feed in the beginning of filtration. The consecutive rings are placed concentrically and self-similar with decreasing radius toward the center of the lumen.

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were inscribed into the lumen from the membrane wall toward the center of the lumen (Figure 3).

The pixel intensities in this ring, which adapts the inner shape of the membrane, are averaged, and the mean intensity is obtained as a function of the radius in the lumen. Conventionally the radius is defined starting in the center of the lumen. Thus, despite of the procedure starting at the membrane's inner wall, the radial MRI intensity distribution is calculated with $I(0)$ being the intensity in the center of the lumen. By visual inspection of the MRI images it was confirmed that the deposit was accumulating uniformly and symmetrically during the filtration processes, that is, no angular dependence of the deposit was observed. The average over all angles was thus found to be a good assumption. In a next step the radial intensity distribution was quantified as a function of time. The intensity gradient from the membrane center to the membrane's inner surface can thus be detected and modelled in terms of filtration theory. During the filtration, the module can additionally be shifted to different z positions to measure the axial dependence of deposition.

Modelling of the MRI intensity distribution

Concentration profiles in the membrane lumen were described in literature.^{3,27} A model was derived under stationary conditions by considering the convective and diffusive mass transport. Under the additional assumption that the dissolved molecules, here alginate and the contrast agent, do not pass the membrane, their concentration c_S as a function of a spatial coordinate can be modelled by an exponentially increasing function

$$\frac{c_S}{c_{s, \text{feed}}} = \exp\left(\frac{J_V r}{D}\right) \quad (2)$$

The quotient of the spatially varying concentration c_S and the feed concentration $c_{s, \text{feed}}$ is an exponential function of the flux $J_V \frac{dV}{A dt}$, that is, the differential permeate volume V per time

interval and filtration area A , the diffusion coefficient D , and the radial coordinate r . The equation results from considerations of mass balance and Fick's law of diffusion.

In the case of in situ MRI experiments, the alginate macro molecules together with the contrast agent accumulate at the membrane surface, which leads to lower MRI intensities, that is, negative image contrast, due to the dominant relaxivity r_2 of the MIONS. As a function of filtration time, the region increases where lower intensities are observed. Assuming that the model in Eq. 2 is reasonable also in the case of the non equilibrium conditions in MRI, that is, nonvanishing permeate flux, an empirical approach is chosen to describe the radial growth of the deposit: An exponential increase of concentration toward the membrane surface is assumed. Reformulating Eq. 2 in terms of MRI relevant quantities in an ongoing filtration, the radially averaged MRI intensity $I(r)$ can be written as

$$\frac{I(r, t)}{I(0)} = 1 - B \exp\left(-\frac{r}{h(t)}\right) \quad (3)$$

$h(t)$ is the time dependent size of the deposit and $I(0)$ the NMR intensity of the feed solution, B is a numerical factor for improving the fit quality. The value of B is between 0 and 1 and describes the amount of deposit. Care was taken to avoid inflow artefacts to guarantee an appropriate reference intensity $I(0)$ in the lumen. The enrichment of alginate, that is, MIONS, toward the membrane wall leads to an intensity reduction, also leading to the modification of the mathematical expression of Eq. 3 when compared with Eq. 2. The center of the coordinate system is located in the center of the hollow fiber with $r = 0$. Thus, the radial symmetry of the CHFM can be exploited by averaging over the angle in the axial images while retaining the radial spatial resolution to derive the deposit height $h(t)$. Please note that h (Eq. 3) is defined in this context as the length at which the intensity decreased by a factor of $1/e$ with respect to the feed value. h does not represent the total height of the deposit although being related to this quantity. The geometric characteristics of the fouling layer is given by h within this description.

With progressing filtration time, the deposit will grow. The analysis of $h(t)$ as a function of filtration time t results in detailed information on an in out ultrafiltration process regarding deposit formation. The simplest approach is an exponential, which is modeled in the present case by the commonly known exponential growth function (4) describing the data well within the experimental error

$$h(t) = h_0 \left(1 - \exp\left(-\frac{t}{t_c}\right) \right) \quad (4)$$

As a result, the time constant t_c , describes the build up of the deposit. t_c , depends on the filtration process.

Model for gel relaxation and equilibration of concentration gradients

A common definition to distinguish gel, that is, fouling layers, from concentration polarization is the characteristics when releasing the filtration pressure and subsequently starting a second filtration. While a gel can relax and swell, a concentration gradient as proposed in the case of pure concentration polarization will equilibrate according to Fick's law when stopping the filtration process. Macroscopically, the different filtration processes can be differentiated accordingly by recording the flux after relaxation and equilibration of the deposit in a second filtration. In the case of concentration

polarization, J_V will return to approximately the value of the native fiber, whereas a continuation with the small values at the end of the first filtration will be observed in the case of cake or dense gel layers.²⁸

The gel compressibility and swelling can be observed by MRI. It was empirically modeled similar to Eq. 3 but applying a Gaussian function revealing the Gaussian width h_G as a function of swelling, that is, equilibration. Again, the time dependence of h_G can be described by an exponentially growing function according to Eq. 4. This approach reveals the characteristic gel relaxation time on a sudden pressure release.

In the case of pure concentration polarization, the time scale is rather short given by Fick's law of diffusive concentration equilibration. It was found that the time needed for pressure release and the subsequent MRI experiment is already too large in the present case to allow for a quantitative modeling of the concentration equilibration. Already after one image acquisition (3 min 12 s) after pressure release, the concentration polarization has almost completely disappeared. The question to be answered in this case therefore concerns the degree of reversibility which was addressed by repeating the filtration.

Results and Discussion

Quantification of deposition and impact of the filtration mechanisms as measured by MRI

CS RARE MRI images allow to detect the membrane lumen in situ during the in out filtration. With progressing filtration time, an increasing deposit layer is observed on the membrane's inner surface. It appears with low intensities caused by the dominantly negative T_2 contrast induced by MagAlg100 as was shown earlier.^{19,20} The CS sampling pattern allows faster measurements with same spatial resolution but lower inflow artifacts as the repetition time T_R can be chosen adequately high to avoid inflow effects without losing the required time resolution for observing the progress of the filtration process. The quantification of the deposit height and the time scale was performed from the time series of MRI experiments as described above.

Filtration of Sodium Alginate with Ca^{2+} : Gel Formation and Compressibility. A filtration using a transmembrane pressure of 1 bar was performed with the described sodium alginate solution while axial CS RARE MRI images were acquired (Figure 4a). A small inflow artefact is visible in the permeate channel, which however does not contribute to the quantification in the feed channel. This is due to the vanishing MION concentration in the permeate leading to longer longitudinal relaxation when compared with the feed solution. The inflow effect could be avoided by longer repetition times, which, however, would lead to inadequately time resolution. The normalized intensities were calculated as a function of the radial distance from the membrane wall (Figure 4b). Equation 3 was fitted to the data and reveal a typical time evolution of the deposit layer characterized by the parameter $h(t)$. The growth of the deposit layer in terms of $h(t)$ was modeled by Eq. 4 (Figure 4c) revealing the growth time constant t_c or its reciprocal value, the growth rate $r_c = 1/(482 \text{ min}) = 0.125/\text{h}$.

After interrupting the filtration, the fouling layer remains sticking to the membrane surface. The fouling layer even increases its size, which is a sign of a swelling gel: When the pressure is released the compressible gel will swell, which leads to a dilation of the fouling layer. This underlines the

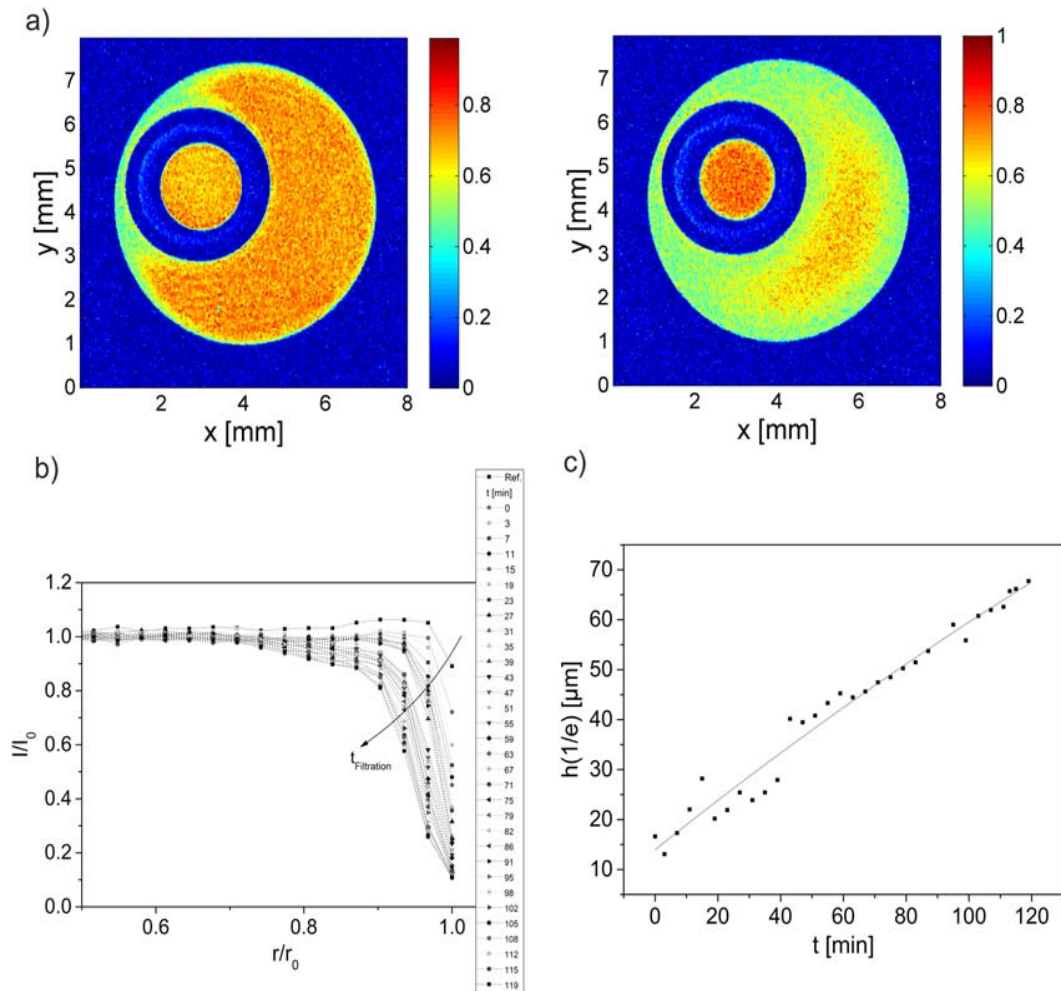


Figure 4. (a) Axial images at start and end of an ultrafiltration of aqueous sodium alginate solution with Ca^{2+} . (b) Radial intensity profiles along the filtration process from which the height h of the deposit layer was calculated. (c) Time evolution of h and modeling by an exponential growth function.

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statement of a gel layer model. The time constant t_c for swelling of the gel layer amounted to 16 min according to Eq. 4. After the interruption, a second filtration was performed with same filtration parameters under in situ observation by MRI. In this second filtration period, the fouling layer continues to build up on the 'old' fouling layer of the first filtration interval which still sticks on the membrane surface. The flux is found to continuously decrease further without any step as observed in earlier experiments.

Filtration of Sodium Alginate Without Ca^{2+} : Concentration Polarization and Equilibration. A loose layer can be observed in the filtration of sodium alginate without Ca^{2+} as described previously.^{19,20} The MRI data were quantitatively analyzed to characterize the filtration process in more detail by applying the mentioned angular average (Figure 5a). The data points slightly oscillate due to MRI data acquisition artefacts. However, the fit of Eq. 3 to the data is numerically sufficiently stable to compensate for this imperfection. As a result, $h(t)$ obviously depends on the filtration time (Figure 5b) and is much larger than in the case of alginate filtration without Ca^{2+} . When fitting Eq. 4 to the data, the time scale t_c is determined, describing the deposition process during filtration. For the filtration without Ca^{2+} , t_c amounts to 14.5 16 min compared to $t_c = 484$ min for the filtration with Ca^{2+} . These findings confirm that the filtration mechanisms differ and depend

on the concentration of the divalent ions. $h(t)$ quantitatively describes the concentration polarization, that is, a concentration gradient from the feed channel in the middle of the lumen toward the membrane's inner surface in the present case of filtration without Ca^{2+} . The smaller values for t_c are a hint for a tendency to build up a polarization concentration. Higher values of t_c will indicate a more gel layer like filtration.

After 90 min, the filtration was stopped, and the pressure was set to zero to observe the relaxation of the deposit in analogy to the experimental design for the filtration with Ca^{2+} . Due to pressure release, the signal intensities in the membrane lumen for the filtration without Ca^{2+} equilibrate. This is another indication for concentration polarization. A different behavior was observed compared to the filtration with Ca^{2+} .

An important question concerns the degree of reversibility of filtration. Apart from the technical and practical issues related to this question, the answer also influences the modeling of the filtration process itself. In the case of a pure concentration polarization the reversibility should be measurable either by repeated filtration cycles in MRI or by time resolved measurements of the classic integral values of flux to be discussed in the next section. After the first filtration and the equilibration of the MRI intensities in the feed channel, a second and third filtration were performed subsequently with MRI measurements at the same position of the fiber. Between

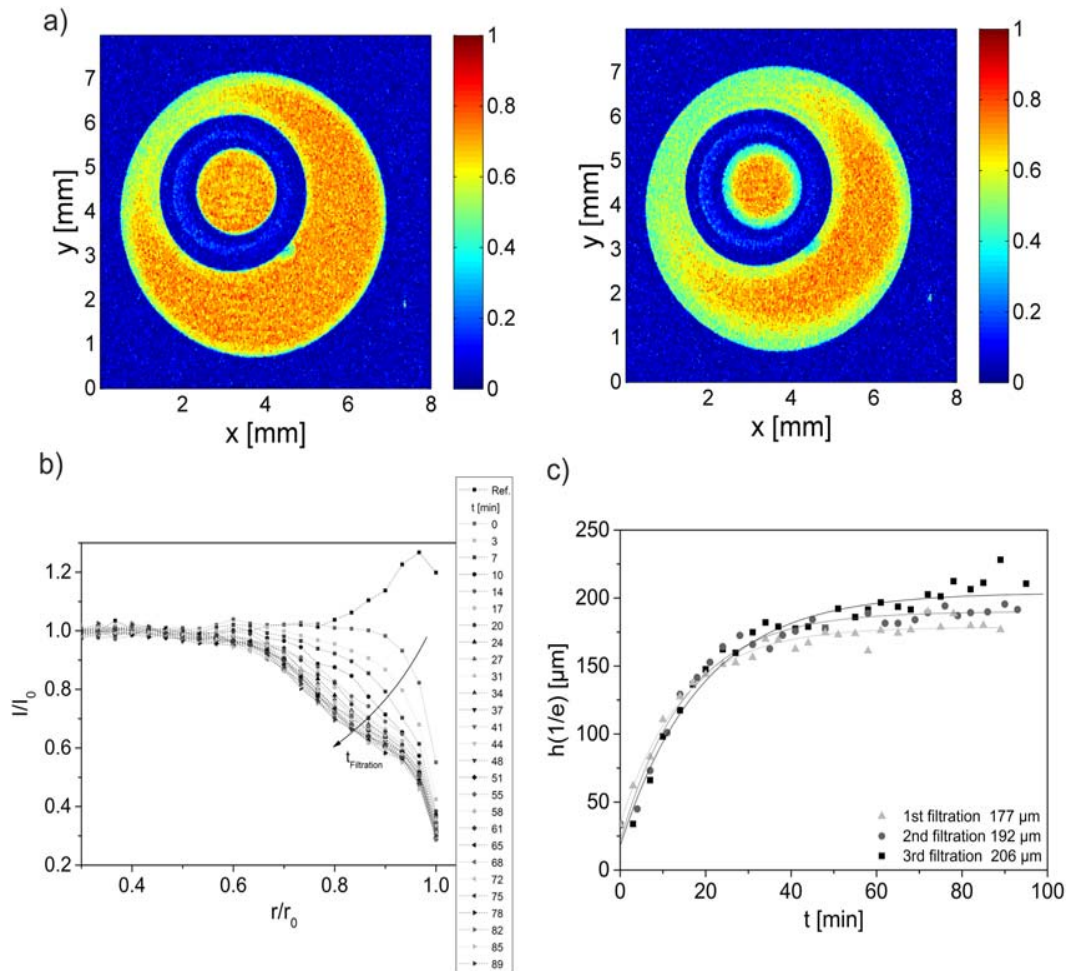


Figure 5. (a) Axial images at start and end of an ultrafiltration of aqueous sodium alginate solution without Ca^{2+} . (b) Radial intensity profiles along the filtration process from which the height h of the deposit layer was calculated. (c) Time evolution of h and modeling by a much faster exponential growth function.

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the first and second filtration the fouling layer diffuses back toward the center of the lumen leading to a larger initial alginate concentration in the second filtration. Whereas the liquid in the lumen was exchanged via cross flow setup between the second and the third filtration. When modeling and fitting in analogy to the previously described experiments, h was determined as a function of filtration time for all three filtration

processes (Figure 5b). The overall behavior is the same, while the time scales slightly differ, but are in the same order of magnitude. The final value $h(t \rightarrow \infty)$ slightly increases with the number of filtration processes as expected. The small ratio $h_1(t \rightarrow \infty) / h_3(t \rightarrow \infty)$ of about 14% indicates, however, that the reversibility is good, but not perfect. To a large extent the filtration of sodium alginate without Ca^{2+} can be regarded as a

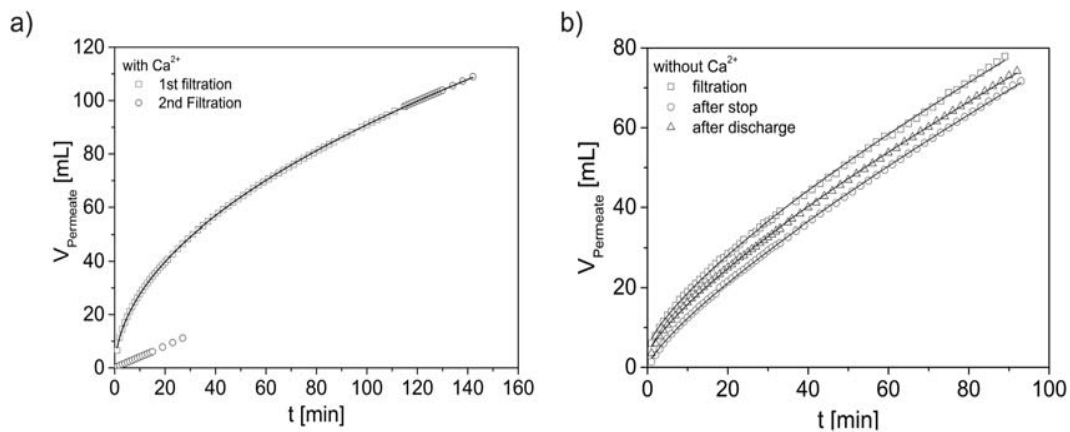


Figure 6. (a) Permeate volume V_{Permeate} as a function of filtration time t for sodium alginate filtration *with* Ca^{2+} . The expected square root dependence of $V_{\text{Permeate}}(t)$ was observed. (b) In the filtration of sodium alginate filtration without Ca^{2+} $V_{\text{Permeate}}(t)$ can be described empirically by $V_{\text{Permeate}}(t) \propto t^{0.7-0.9}$.

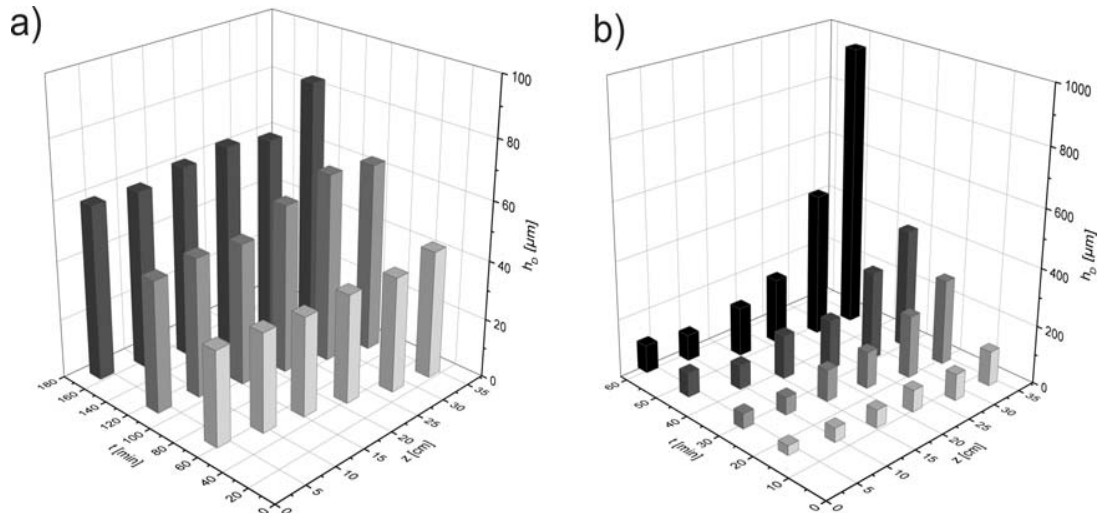


Figure 7. h as a function of filtration time t and z , that is, along the long axis of the cylindrical fiber.

(a) Ultrafiltration of aqueous sodium alginate solution with Ca^{2+} : the deposit layer is more or less equally distributed and grows with time. (b) The concentration polarization layer build-up is a strong function of z and t in the case of filtration without Ca^{2+} .

reversible concentration polarization, whereas the filtration of sodium alginate with Ca^{2+} showed the characteristics of a dense gel layer formation.²⁸ This detailed characterization was possible by the CS RARE MRI, revealing the characteristics of the filtration processes in great detail.

Flux after interruption of ultrafiltration

To underline the MRI data analysis, the permeate flux was integrally observed during the experiments. The ultrafiltration processes with and without Ca^{2+} were interrupted, and the pressure was reduced to 0 bar. After observation of the relaxation of the deposit, both filtrations were started again with the same filtration parameters. During these experiments, the permeate volume was volumetrically measured as a function of filtration times (Figure 6).

The flux is different in both experiments. For the filtration without Ca^{2+} , an almost complete flux recovery is observed. The minimal deviations in the beginning of the three filtration processes show up in the measured volumes and might be attributed to pore blockage or small alginate agglomerates sticking to the membrane surface. The permeate volume for the filtration with Ca^{2+} behaves differently: In the second filtration after the interruption, the flux increases monotonically and uniformly in continuation of the end of the first filtration.

These volumetric and integral results are in good agreement with the MRI measurements, where a compact and irreversible layer could be measured for the filtration with Ca^{2+} (Figure 4a) and a loose, fluffy layer with a larger extent could be observed for the filtration without Ca^{2+} (Figure 5a).

Deposit formation along the fiber axis

As the pressure varies along a ceramic hollow fiber membrane, the deposit is expected to be a function of length, too. The coordinate is denoted as z according to the NMR nomenclature and is 0 at the inlet of the feed into the fiber (Figure 1). The point at $z = 35$ cm is located near the permeate outlet. In additional experiments, ultrafiltrations of sodium alginate with and without Ca^{2+} were investigated as a function of z and of filtration time t . The MRI data were recorded and processed applying the abovementioned procedures. As the filtration mechanisms depend on the Ca^{2+} concentration, the

z dependence is expected to differ for these two experiments (Figures 7a, b).

In the case of gel layer formation (Figure 7a), the height of the deposit is again much smaller compared to concentration polarization (Figure 7b), but the increase as a function of z is still pronounced and measurable. With increasing filtration time, the gel layer thickness increases slightly.

The z dependence of the concentration gradient is much more pronounced in the filtration *without* Ca^{2+} : The deposit increases drastically with increasing t and z , which was also reported by Panglisch in simulations of particle filtration in dead end mode.²⁹ Please note the scale of h . The findings are in agreement with the expectation from classic filtration experiments and is now quantified noninvasively and time resolved during filtration experiments. Concluding, the experiments show the capability of fast MRI to detect deposits in CHFMs and to provide the basis for quantification of typical time and length scales involved in filtration processes. Based on the spatially and time resolved MRI results the mechanisms can be described in detail which could lead to further development of dedicated numerical and analytical models in filtration theory.

Conclusions

In situ filtration MRI experiments were quantitatively analyzed to describe the mechanisms of deposit formation during ultrafiltration of sodium alginate in CHFMs. The CS RARE pulse sequence allowed short measurement times while using relatively large repetition times to avoid inflow artefacts especially in the feed channel in the images. The membrane lumen was mapped in the images by angular averages of the intensity as a function of the radius. The radial intensity profiles were modeled to obtain the characteristic length scales of the deposit h . $h(t)$ could further be modeled with an exponential growth function with the characteristic time constant t_c . This parameter depends strongly on the Ca^{2+} concentration indicating the fouling mechanisms of the two ultrafiltration processes. Low values for t_c are attributed to concentration polarization and large values of t_c indicate a dense and compact, irreversible gel layer formation. This interpretation was underlined by flux interruption experiments, which gave

insight into the reversibility of the deposit structures. As the deposit formation depends on pressure a height dependence is expected in both cases of filtration and is observed in MRI. Using the same modelling of the data, the deposit height was measured as a function of filtration time and fiber height z . The results are in agreement with the expectations from the postulated filtration mechanisms.

Notation

Symbols

A = active filtration area of the membrane, m^2
 B = numerical factor,
 c_s = concentration of alginate/contrast agent, kg/m^3
 D = diffusion coefficient, m^2/s
 h = geometric characteristics of fouling layer, μm
 J_V = permeate flux, $L/(m^2h)$
 r = radial position, mm
 r_2 = transverse relaxivity, $(s\text{ mM})^{-1}$
 T_2 = transverse relaxation time, s
 T_R = repetition time, s
 t = filtration time, min
 t_c = time constant, min
 V = filtrate volume, mL

Greek letters

σ = standard deviation,
 λ_1 = regularization parameter,
 λ_2 = regularization parameter,
 $\tau_{E, effective}$ = effective echo time, ms

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