Bachelor Thesis

Optimization of Polyelectrolyte Multilayer Capsules

Optimierung der Polyelektrolyt Kapseln

by

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Abstract

Polyelectrolyte multilayer capsules based on CaCO₃ and SiO₂ cores coated with the polyelectrolyte systems PAH/PSS and PDADMAC/PSS have been analyzed. A series of measurement has been conducted to study the impact of temperature, rotation speed during precipitation and concentration of the stock solutions on the CaCO₃ core diameter. Also the effect of annealing on the empty capsules has been observed. As a result an impact of the heating on the diameter was observed, but in a few cases swelling was detected although shrinking was expected. Furthermore a strong dependency of the CaCO₃ particle size on the rotation speed during the particle precipitation has been observed. The CaCO₃ microspheres increased in size if the rotation speed was decreased from 1200 rpm to 250 rpm while in general keeping their monodisperse character. This could be used to obtain monodisperse particles with tunable size that can be employed in a wide field of applications like biotechnology, medicine or catalysis.

1 Introduction

In the 1990’s the method of layer-by-layer thin films was developed by Decher et al. [1] on flat surfaces. It is based on the electrostatic attraction of oppositely charged functional groups, that can subsequently be adsorbed onto a charged template. The driving force of this method is charge overcompensation after each layer [2] which allows several layers to be applied.

In particular polyelectrolytes as layer material have several advantages: Because of their ionic nature polyelectrolytes can be adsorbed from aqueous solution in a cheap and environmentally friendly way. Moreover the coating with polyelectrolytes is mostly independent of the template form as long as it has a surface charge. Soon the potential to coat also spherical particles was found out as well as the fact, that after core removal spherical hollow capsules are obtained. These capsules can be loaded with molecules of different molecular weight and the polyelectrolyte walls can be functionalized additionally with nanoparticles[3]. The production of these polyelectrolyte multilayer capsules is of great interest because of their potential as drug delivery systems[11] with controlled releasing properties[12][16][3], as sensor systems [26], as microreactors[13][14][15], as catalytic systems[13][14], etc.

The basic polyelectrolyte multilayer capsule is composed of two components: the sacrificial core and the polyelectrolyte multilayer wall. The coating process for the typical system (PSS/PAH)$_n$ on CaCO$_3$ can be seen in figure 1.

Fig. 1: Fabrication of hollow polyelectrolyte microcapsules via layer-by-layer assembly, taken from reference [3]
1 INTRODUCTION

1.1 Templates

For the template a wide range of materials have been used. If the spherical particle is not the cargo itself, these templates are sacrificial. This is the case most of the time. Generally, polyelectrolyte capsules are permeable to small molecules with a dimension of a few nanometers, but impermeable to bigger molecules. Thus the core can be removed, if the particles fulfill the qualification of being dividable into low molecular weight fragments.

Materials used until now reach for example from organic colloids like polystyrene latex [4] and melamine formaldehyde (MF)[10] to inorganic template materials like CaCO$_3$[9] to SiO$_2$[16]. They differ significantly in their properties like monodispersity, porosity and solubility as well as biocompatibility.

While CaCO$_3$ cores can be dissolved easily and environmentally friendly with aqueous ethylenediaminetetraacetic acid (EDTA) solution, their size range is relatively large and difficult to control. SiO$_2$ cores on the other hand can be produced monodispersely but their solution with HF is complicated because of its low dissociation constant. The originally employed weakly crosslinked MF particles dissolve at low pH, but some MF oligomers can remain[10]. Because of the biological incompatibility these cores are limited in their use and not further analyzed in this work.

A further property that depends on the core is the loading method. In general the methods are categorized in 'pre-loading' and 'post-loading' methods.

If 'pre-loading' is used either the cores are filled by coprecipitation with the molecules before the layer-by-layer assembly or the molecules are adsorbed onto the porous core surface before coating.

In the 'post-loading' strategy hollow capsules are produced that are loaded afterwards by changing the permeability of the capsule walls. This can be achieved for example by changing the pH or the temperature and can be reversible or irreversible.

In the presented work the pre-loading method is used with CaCO$_3$ particles and for SiO$_2$ cores post-loading is used.

CaCO$_3$ Cores

CaCO$_3$ particles show many of the desired properties for templates and have therefore been studied a lot recently. For the composition of CaCO$_3$ different reactions can be used. In this work NaCO$_3$ and CaCl$_2$ have been used so that, following the reaction equation 1, calcium carbonate is produced.
There are several polymorphs in which CaCO$_3$ can crystallize:

**Calcite** is the thermodynamically most stable form of CaCO$_3$ and is widely available in nature. Its crystal structure is trigonal-rhombohedral and consists of alternate CO$_2^-$ and Ca$^{2+}$ layers.

**Aragonite** is thermodynamically less stable but often found in nature. Alternating CO$_2^-$ and Ca$^{2+}$ layers are arranged in an orthorhombic crystal structure.

**Vaterite** is a thermodynamical metastable form rarely found in nature. It has an hexagonal crystal structure of alternating CO$_2^-$ and Ca$^{2+}$ layers.

For the usage as sacrificial cores for polyelectrolyte multilayer capsules spherical charged particles with a size of 2 $\mu$m-8 $\mu$m are needed. For this purpose the polymorph vaterite is applicable, since the small vaterite crystals ($\approx$ 50 nm) aggregate to spherical particles of the desired size range[17] and surface potential[19].

**Amorphous CaCO$_3$,** that does not have a long range order of the atom position, also forms particles in the desired size range[21]. These particles are in general spherical and obtain a vaterite coating after 30 min[18], so that they could be used as template for polyelectrolyte adsorption. In this work no structure analysis has been done, so it could not be distinguished whether clusters of vaterite or amorphous CaCO$_3$ was the final product in the reaction. Nonetheless mainly vaterite particles are expected, because Yan et al. reported, that the crystallization of CaCO$_3$ particles depends on the stirring speed during precipitation. By X-ray diffraction they found out, that the spherical particles consisted mainly of vaterite while the hexahedral crystals were mainly composed of calcite[20].

In figure 2 A-C SEM pictures of the resulting particles are shown. In order to coat these particles it is important to know the polarity of their charge. Volodkin et al.[5] found out, that the surface charge of CaCO$_3$ particles depends on the pH of the surrounding fluid. For pH 7 and 8 the surface charge is positive and for pH 9 and 10 the zeta-potential changes to negative. The CaCO$_3$ particles produced this way are reported to be porous and to have a rather big surface area[7]. Because of this they are particularly suitable for pre-loading via coprecipitation.

Because neither the vaterite nor the amorphous CaCO$_3$ is stable, it is necessary to either dry the spheres for storage or to coat them in order to prevent the formation of the more stable calcite or aragonite.

After the assembly of the desired number of polyelectrolyte layers the CaCO$_3$ core can be removed from the microsphere using EDTA as the chelating agent under mild conditions. This procedure has been reported to not affect the encapsulated material[9].
Fig. 2: SEM images of CaCO$_3$ particles. A, in Overview; B, broken particle; C, recrystallization of a particle; D, calcium carbonate microcrystals from Aldrich, taken from reference [5]

Fig. 3: Change of $\zeta$-potential in mV of CaCO$_3$ microparticles as a function of pH, taken from reference [5]
1 INTRODUCTION

SiO$_2$ cores

For the preparation of hollow microcapsules based on SiO$_2$ as template, a commercial product was used (4.78+/-0.19 µm).

In order to dissolve the cores after coating 0.3 M hydrofluoric acid is used following the reaction 2.

$$SiO_2 + 4HF \rightarrow SiF_4 + 2H_2O$$ (2)

This procedure is more hazardous because of the acid and therefore needs to be handled with special care. However the monodisperse and non aggregated SiO$_2$ microspheres are apt for polyelectrolyte capsule preparation.

1.2 Polyelectrolyte capsule walls

The capsule walls are build by polycation–polyanion complexes that are formed through electrostatic interaction. In order to understand this mechanism it is necessary to have a closer look at polyelectrolytes.

Polyelectrolytes are charged polymers in which each monomer has at least one functional group that can be charged. They are divided into 'strong' and 'weak' polyelectrolytes describing their charge dependency on the pH. Strong polyelectrolytes have a charge that is mainly independent of pH while for weak polyelectrolytes the charge can be modified by changing the pH. An attribute with that they can be categorized is their acid dissociation constant $K_a$ that is the equilibrium constant for the dissociation:

$$HA \rightleftharpoons A^- + H^+$$ (3)

The dissociation constant can then be written as the quotient of the $A^-$, $H^+$ and HA equilibrium concentrations:

$$K_a = \frac{[A^-][H^+]}{[HA]}$$ (4)

In order to reduce the range of this constant the negative decadic logarithm is often used:

$$pK_a = -\log_{10}K_a$$ (5)
Polyelectrolytes with a small $\text{pK}_a$ value are considered strong. In contrast they have the characteristics of weak polyelectrolytes, if the $\text{pK}_a$ value is between $\approx 2$ and $\approx 10$.

The amount of charged functional groups has also an effect on the structure of the polyelectrolyte: The more functional groups are charged the more the polyelectrolyte is stretched because of electrostatic repulsion. Therefore a less charged weak polyelectrolyte will have a rather coiled polymer conformation.

The system PSS/PAH

Poly(allylamine) (PAH) and Poly(styrenesulfonate) (PSS) are polyelectrolytes whose properties in multilayer build up are well studied. PSS is a strong polyanion with a $\text{pK}_a$ value of about 2 while PAH has a $\text{pK}_a$ value of about 8.5 and therefore is a weak polycation. In order to have a system in which a similar amount of both polyelectrolytes is adsorbed, nearly equally charged polymers are necessary. Because of that PAH with a pH of 6.5 is typically used for the coating. It has been shown, that capsules build of $(\text{PSS/PAH})_n$ multilayers change their size if heated. This effect will be explained further later.

The system PSS/PDADMAC

Poly(diallyldimethylammoniumchlorid) (PDADMAC) is like PSS a strong polyelectrolyte and has an cationic character. Earlier studies have shown, that hollow capsules with an even polyelectrolyte number and PSS as the outmost layer shrink upon temperature increase. The wall thickness increases during this process and consequently the permeability for smaller molecules should decrease\cite{25}.

The structure of PSS, PAH and PDADMAC is displayed in figure 4.

1.3 Heat sensitivity

The procedure of post-loading is especially interesting for macromolecules with particularly high or low molecular weight.

For the system PDADMAC/PSS it has been observed, that a swelling of the capsule resulted in a decrease of wall thickness while the wall thickness increased during capsule shrinking\cite{23}.
This suggests, that the permeability increases with swelling and decreases during shrinking. In order to use this effect for amplification of the loading possibility, the thermal behavior of polyelectrolyte capsules has been under examination.

Two systems, that are sensitive to the temperature have been described above:

The systems PAH/PSS and PDADMAC/PSS.

The change in size seems to be dependent on the system used, the number of layers[24] and the charge of the last layer. The results for even layer number given in the literature are summarized in the table below.

<table>
<thead>
<tr>
<th>Polyelectrolyte</th>
<th>Last layer</th>
<th>Thermal effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSS/PDADMAC</td>
<td>PDADMAC</td>
<td>swelling (reversible)[23]</td>
</tr>
<tr>
<td>PDADMAC/PSS</td>
<td>PSS</td>
<td>shrinking (irreversible)[25]</td>
</tr>
<tr>
<td>PSS/PAH</td>
<td>PAH</td>
<td>shrinking (irreversible)[27][31]</td>
</tr>
<tr>
<td>PAH/PSS</td>
<td>PSS</td>
<td>shrinking (irreversible)</td>
</tr>
</tbody>
</table>

Table 1: Thermal effect on polyelectrolyte multilayer capsules

Additionally it has been shown, that corresponding with the (PDADMAC/PSS) capsule shrinking the wall thickness increased and the permeability for low molecular weight macromolecules is decreased[28]. These properties allow to post–load hollow capsules with (PDADMAC/PSS) walls via heat shrinking and to encapsulate molecules with a low molecular weight (10 kDa)[29]. The swelling of capsules with PDADMAC as the last layer was explained with an electrostatic repulsion of the excess positive charge[23], while the shrinking in the case of PSS as last layer
was explained with a polymer rearrangement to reduce the surface tension [25].
The procedure applied in this work was mainly according to W. Song et al. [30].

1.4 Goal

As explained above the fabrication of polyelectrolyte multilayer capsules based on 
CaCO$_3$ cores is of great interest. For this production it is necessary to be able 
to influence the particle size and to get as high reproducibility as possible. With 
capsules based on size controlled and monodisperse particles, the opening of with 
gold nanoparticles functionalised capsules can be analyzed better and the usage 
as sensor system can be improved. Because the precipitation process of CaCO$_3$ is 
influenced by several aspects, the aim of this work is to optimize the influenceability 
of size.

Once the problem of the size range is solved, finding a way to load the capsules 
with particles of low molecular weight is desirable. In order to accomplish this, it 
would be necessary to reduce the permeability of capsules. The idea is to archive 
the permeability reduction by using heat sensitive polyelectrolytes that show an 
irreversible size reduction at high temperatures. This 'heat shrinking' could be 
used to post-load capsules with low molecular weight molecules.
2 Analysis

For the analysis of the cores confocal laser scanning microscopy (LSM) was the main tool. The capsules were also analyzed by the LSM and the loading success was observed by fluorescence microscopy. In all experiments the macroscopical precipitation process has been observed as well as the dissolution process of the cores.

2.1 Confocal laser scanning microscopy

The basic idea of confocal laser microscopy is to focus on one point in the probe at a time and to ignore the rest of the reflected light. This way the sample can be scanned point by point and the resulting picture is a 2D cut of the area of interest. Thus a much higher resolution than with conventional microscopes can be achieved. By combining several 2D layers, a 3D picture of the sample can be produced and recorded.

A scheme how this idea is technically realized is shown in figure 5.

![Diagram of confocal laser scanning microscope](image)

Fig. 5: Schematic structure of a confocal laser scanning microscope, taken from reference [8]
The Laser beam is first expanded and directed to a dichroic mirror, that reflects only the part of the light with the selected wavelength. This light is focused by the objective and hits the sample. Fluorescent samples absorb the light and emit a light with a usually longer wavelength and lower energy. For non fluorescent probes the transmission channel is used in which the wavelength does not change. The emitted light with an apt wavelength passes the dichroic mirror again. In front of the detector a pinhole blocks the light from outside the focus point and an emission filter again selects the light with the desired wavelength range. Afterwards the remaining light reaches the detector and can be recorded. Because only one point in the focus level is recorded by this procedure, it is necessary to scan the whole probe to get a full picture.

2.2 Data interpretation

Size study

For the CaCO$_3$ size analysis a photo was taken with the LSM using the Zeiss software ZEN 2007. According to the magnification a scale bar of 5 $\mu$m was added. This picture was then loaded in the public domain program ImageJ in which the scale was set apt to the scale bar. Afterwards the picture was made binary and the aggregated templates were erased. If necessary the colors were inverted, so that the cores were represented by black circles. As soon as only nearly spherical and non aggregated templates were left, the program ImageJ could calculate the black areas. This data could then be transferred into diameter information via the following equation.

$$d = 2 \cdot \sqrt{\frac{A}{\pi}}$$  \hspace{1cm} (6)

In this equation $d$ gives the diameter in $\mu$m and $A$ is the area in $\mu$m$^2$. In the following discussion the word size is used as a synonym for the particle diameter. A source of error in this analysis is, that the CaCO$_3$ particles are not perfectly spherical. Therefore the diameter calculation using equation 6 is not exact. This error can be reduced by statistical analysis of many microparticles as has been done for these experiments.

Another problem is that in LSM pictures the choice of focus has an impact on the apparent size of the particle. As is shown in figure 6, the impact of this error is proportional to width of the size range: Particles smaller than the average will
seem to be even smaller, because the scanning layer will be above their middle section and bigger particles will also appear smaller because of the lower scanning layer. Therefore pictures with a low variation in particle size will be more exact. This effect could be reduced by analyzing one microsphere at a time with the most suitable focus level. On the other hand this is a very time consuming procedure which has not been done for this analysis.

![Fig. 6: The difficulty of focus for spheres with different sizes](image)

**Heat shrinking**

The heat shrinking procedure consists of different steps, which are explained further in the section methods. In between the various steps the samples were looked at under the microscope. This way pictures were produced from before the coating, after the coating, after the core dissolution, after the incubation with the fluorescein isothiocyanate–Dextran (Dextran-FICT) and after the heat exposure. Additionally parts of the probes were dried after core dissolution in order to test, if the core was removed completely. The size distribution of SiO$_2$ based capsules was analyzed in the same way as for the CaCO$_3$ cores in order to evaluate the shrinking process. Furthermore the ability to load these capsules with fluorescent labeled 10 kDa Dextran was studied with a confocal fluorescence microscope as described above.
3 Experimental

Materials

**milliQ water**: deionized water (Millipore R ≥ 18.2 MΩ) was used for all solutions and washing steps

**PAH** (Poly(allylamin hydrochloride), Sigma, no. 283223, MW ≈ 56 000 g/mol): weak polycation

**PDADMAC** (Poly(diallyldimethylammoniumchloride), Sigma, no. 409022 MW ≈ 350 000 g/mol): strong polycation

**PSS** (Poly(sodium-4-styron sulfonate), Aldrich, no. 243051, MW ≈ 70 000 g/mol): strong polyanion

**EDTA** (Ethylenediaminetetraacetic acid, Sigma, no. E5134): chelating agent for Ca\(^{2+}\) ions

**HF** (Hydrofluoric acid, 0.3M): dissolvent for silicon dioxide

**SiO\(_2\) spherical particles** (diameter 4.78 ±0.19 µm) template for capsules

**Dextran** Dextran (Mw 2000 kDa) and Dextran–FITC (fluorescein isothiocyanate–Dextran, Mw 10kDa)

**Na\(_2\)CO\(_3\)**

**CaCl\(_2\)**

Equipment

**Centrifuge** Thermo Science, Varifuge 3.0RS for the CaCO\(_3\) core dissolution and microfuge ALC, Centrifugette 4214 for the washing processes

**Ultrasound sonicator** Bandelin, Sonorex super RK103H

**Magnetic stirrer**

**Vortex generator** Sartorius, Certomat MO II

For analyzing the experiments always a Zeiss microscope with Argon Laser and a 100x Oil Immersion objective was used.

3.1 Methods

3.1.1 CaCO\(_3\) core synthesis

For the CaCO\(_3\) precipitation 0.3 M solutions of each CaCl\(_2\) and Na\(_2\)CO\(_3\) were prepared. As substituent for any cargo in the pre–loading method a solution of
Dextran 2000 kDa with the concentration $5\frac{mg}{ml}$ was prepared and mixed with the required amount $CaCl_2$ solution prior to precipitation with $Na_2CO_3$. If not written otherwise the experiments were conducted at room temperature (20 °C-24.5 °C) in a lab environment (atmospheric pressure: 983 hPa-1020 hPa; humidity: 22 %-41.5 %). For each of the different test series one parameter was modulated as will be described afterwards.

In the beginning of the synthesis 770 µl of aqueous Dextran solution was mixed with 615 µl of $CaCO_3$ solution while the rotation speed of the magnetic stirrer was adjusted to 1000 rpm. Afterwards equally 615 µl of the $Na_2CO_3$ was added and left under invariant stirring conditions for 30s. After this precipitation time the glass vial was removed from the magnetic stirrer and the resulting particle suspension was divided into two 1.5 ml tubes. The tubes were left to rest for 3 min at room temperature and afterwards three washing steps with water as well as one with acetone were applied. In each washing step 1 ml of water or acetone were first added to the precipitant. In order to wash the particles and to reduce the aggregation, they were held onto the lab vortex, so that they were mixed with the fluid. For sedimentation of the particles in order to remove the supernatant again, the tubes were centrifuged at 6000 rpm in a centrifuge with a radius of about 5 cm which results in a maximal relative centrifugal force

$$G = \frac{a_z}{g} = \frac{\omega^2 r}{g} = 2012 \quad (7)$$

with g the earths gravitational acceleration. The relative centrifugal force is also called g-force.

After the centrifugation the fluid on top of the precipitant was removed and new water was added to perform the next washing step.

After the washing with acetone the tubes were punctuated with a needle and stored under vacuum for drying.

Three different parameters were analyzed: The impact on the particle size of the temperature, the rotation speed during precipitation as well as the concentration of the stock solutions and Dextran. For each parameter five modulations were chosen and two samples were prepared. The analyzed temperatures were 5 °C, 10 °C, 15 °C, 20 °C and 25 °C. For the different volume concentrations 250 µl, 500 µl, 650 µl, 770 µl and 1000 µl of the aqueous Dextran was used in the synthesis which results in volume concentrations of $0.17 \frac{l}{l}$, $0.29 \frac{l}{l}$, $0.35 \frac{l}{l}$, $0.39 \frac{l}{l}$ and $0.45 \frac{l}{l}$. As rotation speeds during the precipitation 250 rpm ($G=0.35$), 500 rpm ($G=1.40$), 750 rpm ($G=3.14$), 1000 rpm ($G=5.59$) and 1200 rpm ($G=8.05$) were used. For the calculated g-forces a mean rotation radius of 0.5 cm was assumed. Each of these experiments was performed five times and, because of the fluctuations in size, on different days.
3.1.2 Heat shrinking

This experiment was carried out at room temperature if not written otherwise. For the preparation of the heat shrinking experiments the spherical SiO$_2$ particles needed to be coated and the cores needed to be removed afterwards, because the shrinking effect only takes place on hollow capsules.

100 µl of SiO$_2$ particle suspension were washed 5 times with water in order to obtain clean particles for the following coating steps. As coating materials two different systems were used: (PSS/PDADMAC)$_4$ and (PSS/PAH)$_4$.

The solutions of PSS ($2 \text{ mg} \text{ ml}^{-1}$ in 0.5 M NaCl), PAH ($2 \text{ mg} \text{ ml}^{-1}$ in 0.5 M NaCl) and PDADMAC ($2 \text{ mg} \text{ ml}^{-1}$ in 0.5 M NaCl) were prepared with a pH of 6.5. For the adjustment of the pH HCl and NaOH were added.

Because the surface charge of the SiO$_2$ is negative, it is necessary to start the coating process with the polycation: PAH or PDADMAC. This means, that the last layer is PSS if an even number of layers is applied.

For each layer the particles of one synthesis were suspended in the polyelectrolyte solution (1 ml each time) and left in an sonication bath for two minutes to avoid aggregation. Afterwards they were shaken further ten minutes. Following to this polyelectrolyte exposure the microspheres were sedimented with a centrifuge and the excess of polyelectrolyte was removed. Three washing steps with water followed as described above, before the polyanion was added and the process was repeated.

Following this protocol 4 bilayers of polyelectrolyte were applied.

The cores of the coated particles were then dissolved by using 1 ml of the acid HF in a concentration of 0.3 M. This suspension was agitated and centrifuged several times for 40 min. Afterwards the acid was removed and, in the case that a white sediment remained indicating incomplete core dissolution, new acid was added and the process was repeated until no sediment was visible anymore. Two washing steps with water followed to withdraw the remaining acid.

After this preparation the heat shrinking process could be started. A solution of 10 kDa Dextran fluorescent labeled with FITC in a concentration of 50 µM was added and left one or two hours for incubation at room temperature. During this time the probe was wrapped in aluminum foil in order to protect it from light. In the meanwhile a water bath was heated to 68 °C in which the tubes were placed subsequently for the heating. They were left for 30 min in thermal treatment for the shrinking process protected from light. Afterwards the suspensions were allowed to cool down to room temperature for 5 min and washed twice to remove the non encapsulated FITC-Dextran.

These probes were analyzed with a fluorescence microscope and stored at about 5 °C.
4 Results

4.1 Impact of rotation speed, temperature and concentration on the particle size

In the following diagrams the mean values of each measurement are plotted versus the studied parameter. As error bars the standard deviations of the mean value are used for the measurements in order to make the experiments with different amounts of data comparable: The number of particles measured is for example much smaller if the particles are bigger since the same amount of CaCO$_3$ is available and the area of the microscope picture remains the same. Because of this problem and in order to get a more significant result by reducing the error at least four pictures of each sample were analyzed.

In the following paragraph the analysis of one rotation speed experiment is presented as an example.

After the synthesis the samples were studied under the microscope and pictures were taken from different locations in the sample. In figure 7 a) an original image from the transmission channel of the microscope is shown exemplary. A scale bar was added and the photos were saved in .jpg format. Afterwards the pictures were processed with ImageJ as described above. The corresponding resulting picture is displayed in figure 7 b).

![Fig. 7: a) Photo of the particles; b) After processing with ImageJ](image)

ImageJ gives a table as output in which the measured black areas are listed. The scale has been set according to the scale bar in the original image and only those black spots with an area from 1 $\mu m^2$ to 100 $\mu m^2$ were chosen. The resulting tables from all pictures of two equally prepared samples were then combined and the
diameter of the microspheres was calculated according to equation 6. The mean value as well as the standard deviation of the mean value were determined and histograms were generated with the program Origin 6.0. An example is presented in figure 8.

Fig. 8: Histogram of experiment 4, 1000 rpm

Histograms of the other experiments can be found in the supplementary information.

Rotation speed

The Diagram in figure 9 summarizes the data of all experiments in which the rotation speed was variated. The different colors represent the different experiments done and the lines connect the five data points of one experimental series. As ordinate in all three summary graphs the size range 0 µm–7 µm was chosen, in order to make the size changing effect better comparable.

As can be derived from the standard deviation of the mean value the measurements of the different experiments are not identical and in some cases not even comparable, although the same parameter was changed.
4 RESULTS

Fig. 9: Variation of the particle size as a function of the rotation speed during synthesis

Fig. 10: Variation of the particle size as a function of the effective g-force during synthesis
This can not be explained clearly. It was assumed that another parameter, like for example slight variations in pH of the solutions, has an impact on the precipitation and resulting particle size. This was not further analyzed.

Despite this unexplained variations a clear dependency of the particle size on the rotation speed could be found. By assuming the not understood parameter as constant the gradient of the curve for the experiments was very similar: The mean value of the size became smaller with increasing rotation speed. In the analysis of the rotation speed, especially significant seems to be the effect on the size in the area between 500 rpm and 750 rpm, where the biggest decrease of particle size could be observed resulting in the steepest slope in the figure 9. The reason for the rise in size at the second experiment with the stirring speed of 1200 rpm is the fact, that in one of the two samples the microspheres had a significantly bigger diameter, which influenced the mean value of this experiment. The histogram, that belongs to this experiment is displayed in figure 11. During the rest of the experiments no large difference in size between the two samples was observed. An explanation for this might be, that in the case of the unusual sample there were seed crystals already present in the glass vial or in the pipette tips. If this sample is neglected, the result of this experiment is more similar to the others.

If a mean rotation radius of 0.5 cm for the precipitation is assumed, the resulting g-force that affects the particles can be calculated. In this calculation the radius has an linear impact, so that slight changes in radius do not effect the form of the curve progression. Furthermore the variation of the radius inside the glass vial is the same for all experiments, so that the impact should not change the results. The resulting diagram is show in figure 10. In this figure the particles become smaller with higher g-force as well, because G is proportional to the square of the rotation speed. Nonetheless the curve progression is different: For small g-forces the particle size increases without showing an upper bound in this region.

In the figures 9 and 10 it can be seen, that the mean values of the microspheres size variate in between 6.5 µm and 2.9 µm, if the rotation speed is modulated in the range of 250 rpm—1200 rpm. Although this is a big difference in size, the maximum difference with equal parameters is 1.6 µm which is rather narrow. This means, that a wide range of particle diameters can be reached using this procedure of variating the speed, which is useful for a wide range of applications.
Fig. 11: Histogram of the particle size in the second experimental serie with a rotation speed of 1200 rpm

Temperature

Fig. 12: Variation of the particle size as a function of the temperature
In figure 12 the dependency of particle size on the temperature was plotted for all of the five experiments. In comparison with the rotation speed experiment the effect of the temperature seems to be much less predictable, because there was a bigger variance in the resulting gradients for the different experiments conducted. In three of the five experiments a tendency towards smaller particles with higher temperatures could be seen and for the experiment 3 the size is oscillating a little bit, but the general tendency is a decrease in size for warmer solutions. However for the fifth experiment the diameter was the smallest at 15 °C and increased towards the more extreme temperatures. Also the biggest size difference with equal parameters was 2.2 µm which is big compared to the other experiments. The mean size range was between 2.7 µm and 5.7 µm and hence smaller than in the rotation speed experiment.

If not only the mean values but also the size distribution is studied it stands out, that for the 5 °C measurement of the third experiment the particles in one sample were again significantly bigger than in the other sample. This results in a much smaller mean value. The diagram that is the evidence for that is shown in figure 13.

In the following paragraph some possible reasons for the smaller reproducibility are explained. One of the problems could be, that the temperature was not stable...
during the experiment. The solution temperature in the beginning of the synthesis was documented as the experiment temperature. Afterwards the vials were not protected from the room temperature, so that the reaction mixtures became warmer during the precipitation step. Another reason for the observed difference could be, that the initial temperature differed slightly in the compared measurements because the synthesis was not performed immediately after temperature measurement. Also an effect could have been the fact, that the volume of the tempered solutions was not always exactly the same because different amounts of solution were used of aqueous Dextran (770 \(\mu\)l) than of CaCl\(_2\) and Na\(_2\)CO\(_3\) (615 \(\mu\)l) for each synthesis. Therefore the temperature was not exactly the same for all of the solutions.

For the synthesis of CaCO\(_3\) particles at very low temperatures around 5 °C a further observation was made: If the temperature was particularly low the CaCO\(_3\) was not sedimented during centrifugation but remained in the solution as a cloudy suspension. Only after several washing steps and with the gradual warming of the suspension, the precipitant was formed. In those samples the size distribution was especially wide: The cloudy solution was stable even after the 30 s resting time for Experiment 1 (0 °C and 6.5 °C) and 2 (5 °C).

Nonetheless it could be shown, that in general the size range was larger when the temperature was lower. At lower temperatures bigger microspheres were produced although some very small particles remained in the suspension as well.

In order to produce bigger particles the effect of lower temperature could be used. The comparatively big size range can be decreased after synthesis by using sedimentation of the bigger particles via centrifugation.
Concentration

In the volume concentration experiment the size variation was the smallest: In all the experiments the sizes ranged between 2.9 \( \mu m \) and 3.9 \( \mu m \) with a size distribution of 0.9 \( \mu m \) at a constant parameter. This variation in size is even smaller than the difference between equal parameters for the temperature experiment, which makes this effect irrelevant in comparison to the unknown size effect responsible for the variation between the experiments. Especially the measurements in experiment one and five are nearly constant in diameter. In all experiments except the third one, there was a very small rise in size between the 250 \( \mu l \) data and the 1000 \( \mu l \) measurement. This could mean, that with a higher Dextran concentration during the synthesis also more Dextran is enclosed in the microspheres. Besides this effect the size range is the narrowest at a volume concentration around 0.39 \( \frac{l}{l} \), the concentration used in the traditional routine. For this concentration 770 \( \mu l \) of aqueous Dextran was used.

Fig. 14: Variation of the particle size as a function of the Dextran aq. volume concentration during precipitation
Other than that the volume concentration seems to have no impact on the size distribution.

4.2 Post–loading of polyelectrolyte capsules prepared on SiO\textsubscript{2} templates via heat shrinking

In the following images the (PAH/PSS)\textsubscript{4} and (PDADMAC/PSS)\textsubscript{4} capsules prepared on SiO\textsubscript{2} templates are shown at the different production steps as well as after the loading with FITC-Dextran.

PAH/PSS

In figure 15 a) the coated microspheres are shown. They are rarely aggregated, monodisperse and present in abundant number. But during the washing steps for the core dissolution the particle number decreased drastically and the particle aggregation increased. Also the layering appeared rather fuzzy under the microscope – the samples were not as spherical any more.

The challenge in this step is the complete removal of the template material. But this seems to happen much slower than expected because of the multilayer coating. Furthermore there was a high loss of capsules because of the washing steps during and after dissolution.

After the core seemed to be removed successfully, the heat shrinking procedure was carried out as described in the section methods. The figure 16 shows some capsules after the heat shrinking in an image divided into fluorescence channel, transmission channel and the corresponding merged image. It can be seen, that the resulting effect was not the same for all of the particles: Two out of the three
capsules in the image were fluorescently labeled while a third one showed no fluorescence. By analyzing the size of the fluorescent and non fluorescent capsules it was observed, that the non fluorescent capsules were even bigger in diameter than before the thermal treatment and that the fluorescent capsules were shrunk between 5% and 16%. In the samples with capsules coated with PAH/PSS about 91% of the analyzed capsules were fluorescent.

The fluorescent capsules had mainly a fluorescent ring structure under the microscope. This suggests, that the FITC-Dextran was caught in between the polyelectrolyte layers and either did not enter the cavity at all or could not be reliably kept inside the capsule.

**PDADMAC/PSS**

The System (PDADMAC/PSS) showed in the first two steps no difference to the system (PAH/PSS): Because of the core removal the capsules became fewer, more aggregated and less spherical.
Fig. 17: a) The PDADMAC/PSS coated SiO$_2$ particles; b) After core dissolution

Fig. 18: PDADMAC/PSS capsules after the heat shrinking and loading
After the heat shrinking procedure not all of the particles were labeled with FITC-Dextran: several particles were either not fluorescent or only a little bit on the outmost edge. In this experiment much less PDADMAC/PSS capsules remained and only 14% of the remaining capsules were clearly fluorescent. Some of the rest had a thin fluorescent boundary that could not always be distinguished. Moreover the shrinking process was not as reliable in this experiment as for the PAH/PSS capsules: some capsules were swelling instead of shrinking during the heat treatment. This is not compatible with the expectations according to the literature. Capsules that were fluorescent showed a much higher fluoresceine concentration in the polyelectrolyte layers than in the capsule cavity.

**Analysis of the problem in the heat shrinking procedure**

In order to find out, whether the Dextran enters the cavity during the incubation, pictures were taken during the incubation time with Dextran-FITC. In both, the PAH/PSS system and the PDADMAC/PSS system, the Dextran could not be found inside the cavity, even if the incubation time was elongated to two hours. Two representative images can be found in figure 19. In the pictures the green fluorescent Dextran–FITC can be seen with two black circles that are the polyelectrolyte capsules. This is evidence, that the Dextran–FITC with a molecular weight of 10 kDa is not able to diffuse in the capsules cavity, since there is no green fluorescent inside. This is unexpected because the polyelectrolyte walls should not be too tight for
Fig. 20: a) PAH/PSS capsules with free fluorescent; b) PDADMAC/PSS capsules with free fluorescent

particles of this low molecular weight to enter. In the literature even particles with higher molecular weight of 10 kDa and more have been reported to be enclosed following this procedure. Maybe the reason for this problem is, that a small amount of the core is left so that a repulsive potential remains. In order to find out, if the capsules are impermeable for even smaller particles, free fluorescent has been tried to encapsulate. The pictures 20 show the results after one hour. Again for most capsules no encapsulation could be seen: The capsules appear as black circles in the fluorescence channel, which means that the intensity of the fluorescence is much lower inside the capsules.

This confirms the interpretation, that there is a further problem with the produced capsules like remains of the SiO$_2$ core, that induce a repulsive potential. For a successful encapsulation this needs to be analyzed further.
5 Discussion and Outlook

The research on this interface of physics, chemistry, material science and biology is much less based on theoretical equations than other fields of physics. This results sometimes in a different approach to problems. Often the principles are first analyzed empirically before the theoretical model follows. In the two areas, that were tried to optimize in this work, the first topic of calcium carbonate precipitation has been subject to research for a rather long time and some models to describe it exist. Despite this, the reproducibility is still limited because the conditions in the lab change continuously: The climate changes, the composition of the surrounding air, the experimental tools were not sterile and other experiments have been conducted alongside. These reasons as well as the fact that the clustering of the vaterite particles and of particles with amorphous CaCO$_3$ is a statistical effect could have an impact on the measurements. This clustering is not yet calculable. This might explain why the results measured are not always very predictable.

The second field of study, the post loading of capsules, on the other hand is a new topic researched only for a few years until now. Some theories exist about why the shrinking takes place for one of the systems, but for the next one it does not apply any more. Because of this and for the reason of its applicability it would be interesting to improve this loading method to make it simple and reproducible. Although an ideal optimization of the synthesis and loading process was not achieved, some helpful tendencies and better defined problems were achieved:

5.1 Rotation speed

Although a prediction of the particle size by adjusting the known parameters is not possible yet, at least the size can be influenced by changing the rotation speed after the size is measured for one synthesis. This pilot synthesis is necessary, to find out the impact of the unknown parameter.

Because the rotation speed is such a nice tunable parameter and relatively reproducible this technique could thus be standardized. It would therefore be interesting to analyze the rotation speed in smaller intervals with the goal to find the mathematical dependency. After a pilot synthesis has been done this could be used to calculate the rotation speed needed to produce the desired size under otherwise constant parameters. This would also help to understand the crystallization process much better.
5.2 Temperature

The effect of a temperature change was not as predictable as for the rotation speed. For low temperatures around 10 °C in general bigger particles than with higher temperatures of 25 °C occurred, but in some cases there was also a rise in size towards the synthesis with warmer solutions. What can be said is, that precipitation at low temperature leads to a wider range of size and thus would make a further isolation of a certain particle size necessary for gaining monodisperse templates. At temperatures close to 0 °C this effect was especially significant. This became obvious during the synthesis, because of the quite stable cloudy suspension, which makes this procedure less predictable and less suitable for standardized production.

5.3 Concentration

The volume concentration of the solutions during the synthesis seemed to have a negligible impact on the particle size. This might mean, that a bigger amount of the cargo can be loaded without increasing the diameter in a way resulting into too big capsules.

At this point it would be interesting to study on the one hand, how much of Dextran was internalized in the particles. This could be quantified by labeling the Dextran for example with a fluorescent dye and to measure and compare the fluorescence levels after synthesis.

On the other hand it would be interesting to find out how much importance the molecular weight and charge of the incorporated cargo has. The Dextran used had a molecular weight of 2000 kDa but it can be expected, that smaller molecules would result in smaller particles. This could be an interesting study for further research.

5.4 Heat shrinking

In this study it was possible to produce a few non aggregated and spherical polyelectrolyte capsules of the systems PAH/PSS and PDADMAC/PSS. These capsules could then be labeled, but not filled with Dextran–FITC and treated with heat. Other than it has been reported in the literature, the shrinking effect of the PDADMAC/PSS capsules was not reliable since some of the capsules swelled instead. Also the main amount of Dextran–FITC was not inside the capsule cavity, from which it could be released in a controlled way, but in the capsule walls.
A reason for these problems might be a non constant temperature in the waterbath, a too short annealing time or a too short incubation time of one hour. These parameters could further be varied to archive a loading of the capsule cavity instead of walls. Furthermore other molecular weights could be used to find the cutoff in which the Dextran can not enter the capsules any more and in which the Dextran can not be kept inside at all. Moreover it would be useful to understand quantitatively how much has been loaded into the cavity by this method under the conditions given.

Because even for free dye and after a long incubation time the fluorophors did not permeate the walls of all capsules, it would be interesting to increase the dissolution time and HF amount to make sure, that all of the core is dissolved.

**Conclusion**

A definite effect on the size was only found out for the rotation speed. The particle size became smaller with higher rotation speed during the synthesis. For the temperature there might be a dependency, but further measurements would be necessary. The concentration of cargo material does not seem to have a relevant effect on the particle size.

For the heat shrinking the production of capsules was accomplished, but Dextran did not enter the cavity but got caught in the walls. Also the shrinking effect was only partly verified.

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Supplementary Information

Experiment 1: Variation of the Dextran aq. amount
Experiment 2: Variation of the Dextran aq. amount
Experiment 3: Variation of the Dextran aq. amount
Experiment 4: Variation of the Dextran aq. amount
Experiment 5: Variation of the Dextran aq. amount
Experiment 1: Temperature

\[\text{Count} \quad \text{Size [\mu m]}\]

- 9°C
- 15°C
- 20°C
- 25°C
- 68°C
Experiment 2: Temperature
Experiment 3: Temperature
Experiment 4: Temperature
Experiment 5: Temperature
Experiment 1: Rotation speed
Experiment 2: Rotation speed
Experiment 3: Rotation speed
Experiment 4: Rotation speed
Experiment 5: Rotation speed
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