Colloidal micro- and nano-particles as templates for polyelectrolyte multilayer capsules

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Abstract

Colloidal particles play an important role in various areas of material and pharmaceutical sciences, biotechnology, and biomedicine. In this overview we describe micro- and nano-particles used for the preparation of polyelectrolyte multilayer capsules and as drug delivery vehicles. An essential feature of polyelectrolyte multilayer capsule preparations is the ability to adsorb polymeric layers onto colloidal particles or templates followed by dissolution of these templates. The choice of the template is determined by various physico-chemical conditions: solvent needed for dissolution, porosity, aggregation tendency, as well as release of materials from capsules. Historically, the first templates were based on melamine formaldehyde, later evolving towards more elaborate materials such as silica and calcium carbonate. Their advantages and disadvantages are discussed here in comparison to non-particulate templates such as red blood cells. Further steps in this area include development of anisotropic particles, which themselves can serve as delivery carriers. We provide insights into application of particles as drug delivery carriers in comparison to microcapsules templated on them.

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Keywords: Microcapsules, Colloidal particles, Templates, Polyelectrolytes, Multilayers

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1. Introduction

The significance of micro- and nano-particles ranges from early developments in the area of colloidal particles to a broad range of industrial and nanotechnology applications in physical, biological, and medical sciences. In this overview, we focus on the class of organic and inorganic particles which are used as templates for the preparation of polyelectrolyte multilayer capsules. The choice of the template for the preparation of these capsules is particularly important in different fields of application, with advantages and disadvantages pertaining to the template dissolution, the stability of the polymer shell, and tendency to aggregation. In addition, these particles can serve as drug delivery carriers themselves.

There are two important classes of templates, possessing either smooth or porous surface. The surface porosity of templates determines a number of parameters of polyelectrolyte multilayer capsules, including the easiness of preparation, polymer shell thickness, and efficiency of loading of molecules. For example, smooth particles usually need the addition of more layers of the polyelectrolytes, and the shell wall is quite thin, on the order of one to two nanometers. The interior of such capsules is typically clean and transparent, and after dissolution of the core particle, such capsules look like empty balloons. On the contrary, during the layer-by-layer (Lbl) adsorption on porous cores, the polyelectrolytes penetrate into the matrix, resulting in a substantially thicker polymer shell. In this case, the capsules look like sponges after the core dissolution. This differential property is very important for loading of molecules and for protection of encapsulated materials. For the capsules templated on porous particles, pores are utilized for loading producing relatively large capacity. Polyelectrolyte capsules based on smooth particles have a bigger inner volume, can easily encapsulate substances by heating and thereby shrinking of surface, and release of cargo from such particles can be more easily manipulated by external stimuli.

Colloidal particles used directly as carriers represent an important class of materials and stand on their own in various areas of research. We elaborate here on the applicability and potential role of particles as templates for the preparation of polyelectrolyte multilayer capsules on one hand, and as stand-alone carriers on the other hand.

2. Particles and templates

2.1. Synthesis of particles

A number of different particles are used as templates for the preparation of polyelectrolyte multilayer capsules. Information about particles most frequently used for the fabrication of polymeric capsules is summarized in Table 1. Templates can be divided into two major classes, namely smooth and porous particles. The smooth ones encompass, for example, melamine-formaldehyde, silica, and polystyrene. Most of them are commercially available in a large range of sizes: from approximately 100 nm to millimeters. Advantages of these particles include good stability (zeta-potential of ~35 mV) and excellent monodispersity. Specific modifications of the surface of these particles with amino-, carboxy-, mercapto-, maleimido-, hydroxy-, epoxy-, sulfo-, and other functional groups are available. The tailored properties of particles allow extending their potential application to medical diagnostics, pharmaceutical industry, biotechnology, molecular biology, analytics, etc. For drug delivery purposes, these types of particles are of limited use due to low loading efficacy on the functionalized surface. The most advanced approach consists in the loading of the hollow polymeric shell which is obtained when the solid core is decomposed, Fig. 1.

Porous particles constitute another type of solid particles. They have a number of advantages over their smooth counterparts, including facile synthesis, low cost of processing, and high porosity which leads to an increased loading capacity. The relatively big variation in size (polydispersity) and strong aggregative behavior are some disadvantages of this type of templates. Production of porous particles (calcium carbonate) is relatively easy, including the mixing of two salt solutions either in the absence or in the presence of special additives which are used to obtain certain surface modifications or a particular morphology.

Table 1

<table>
<thead>
<tr>
<th>Type of the particles</th>
<th>Size of the particles used as templates for polyelectrolyte capsules, microns</th>
<th>Size range, microns</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smooth particles</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Silica</td>
<td>0.5 [2] 1 [2], 1.9 [3], 4.5–5 [4–7]</td>
<td>2–5</td>
<td>HF, HF/NH₄F</td>
</tr>
<tr>
<td>Melamine–formaldehyde</td>
<td>1 [8], 2 [9,10], 3.2–3.5 [8,11,12], 4.2 [13], 5–5.7 [14–16]</td>
<td>3.2–5.7</td>
<td>HCl, N,N-dimethylformamide (DMF) or dimethyl sulfoxide (DMSO).</td>
</tr>
<tr>
<td>Polystyrene</td>
<td>0.64 [8] 4.4–4.6 [5,17,18], 9.6–10.6 [19,20]</td>
<td>4–20</td>
<td>THF</td>
</tr>
<tr>
<td>CaCO₃ cubic shape</td>
<td>2.5, 4–6 [21]</td>
<td>2.5–6</td>
<td>HCl, pH = 1</td>
</tr>
<tr>
<td>CaCO₃ cubic shape</td>
<td>4–5 [22]</td>
<td>4–5</td>
<td>HCl, EDTA</td>
</tr>
<tr>
<td>Calcium phosphate modified by PAH</td>
<td>158–300 nm [23]</td>
<td>0.16–0.3</td>
<td>HCl</td>
</tr>
<tr>
<td>Gold nanoparticles</td>
<td>13.5 nm [24]</td>
<td>0.14</td>
<td>Potassium cyanide</td>
</tr>
<tr>
<td>Micro gel dex-HEMA</td>
<td>150 [25]</td>
<td>150</td>
<td>NaOH solution</td>
</tr>
<tr>
<td>Alginate hydrogel microspheres</td>
<td>4.25 [26]</td>
<td>4.25</td>
<td></td>
</tr>
<tr>
<td>poly-β-L-lactic acid particles (PLA)</td>
<td>4–8 [10]</td>
<td>4–8</td>
<td>1:1 mixture of 1-methyl-2-pyrrolidinone acid and organic solvent (acetone)</td>
</tr>
<tr>
<td>Halloysite</td>
<td>Cylinder 50 nm × 300 nm [27]</td>
<td>0.05 × 0.3</td>
<td></td>
</tr>
<tr>
<td>Porous particles</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mesoporous silica</td>
<td>0.4 [28], 2–4 [1,29]</td>
<td>0.4–4</td>
<td>HF, HF/NH₄F</td>
</tr>
<tr>
<td>Calcium carbonate spherical</td>
<td>4–5 [18,22,30,31], 9 [32], 11.5 [33]</td>
<td>4–11.5</td>
<td>EDTA pH 6; acid solution pH &lt; 6</td>
</tr>
<tr>
<td>MnCO₃</td>
<td>1.9 [9] 2.5–2.8 [9,34], 3.6 [32], 5–5.5 [35,36]</td>
<td>2.8–5.5</td>
<td>EDTA pH 6; acid solution pH &lt; 6</td>
</tr>
<tr>
<td>Calcium carbonate elliptic</td>
<td>2–4 [22]</td>
<td>2–4</td>
<td></td>
</tr>
<tr>
<td>Hybrid templates</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Human erythrocytes</td>
<td>5–6 [37–42]</td>
<td>5–6</td>
<td>140 mM NaCl and 1.2% NaOCl</td>
</tr>
<tr>
<td>Virus-like particles</td>
<td>–4 [43]</td>
<td>–4</td>
<td></td>
</tr>
<tr>
<td>Liposomes</td>
<td>0.1–0.5 [44]</td>
<td>0.1–0.5</td>
<td></td>
</tr>
</tbody>
</table>
2.1.1. Calcium phosphate

Calcium phosphate synthesis usually involves mixing of two salt solutions to initiate crystal growth. Calcium phosphate precipitates form as a result of colloidal aggregation during mixing of CaCl₂ and Na₂HPO₄ in a buffer solution (TRIS pH 7.4) [45]. The PAH-modified nanoparticles of calcium phosphate were prepared at room temperature by rapidly pumping aqueous solutions of calcium lactate, (NH₄)₂HPO₄, and poly(allylamine hydrochloride)(PAH) in a volume ratio of 1:1:1 into 4 volume parts of water, forming particles with diameter of about 158 nm [23]. Another approach for the synthesis of smaller calcium phosphates nanoparticles (20 nm) was proposed by Y. Cai [46]: A CaCl₂ solution was dropwise added to a Na₂HPO₄ solution containing different concentrations of hexadecyl(cetyl)trimethyl ammonium bromide (CTAB) in magnetically stirred vessels at 20 °C.

2.1.2. Calcium carbonate (CaCO₃)

Synthesis of calcium carbonate templates is based on crystal growth of polycrystalline spherical vaterite particles precipitated by mixing of CaCl₂ and Na₂CO₃ solutions [47]. The nucleation and growth rate of

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**Fig. 1.** Transmission electron microscopy (TEM) image of a) catalase-loaded mesoporous silica (MS) spheres coated with (PLL/PGA)₃ multilayers, and b) following removal of the MS template; c) fluorescence images of (PLL/PGA)₃ microcapsules loaded with FITC-labeled catalase. The scale bar in the inset corresponds to 800 nm. PLL/PGA layers were assembled from a 0.05 M MES, pH 5.5 buffer. The MS spheres were dissolved using HF/NH₄F at pH = 5. Figure reprinted from [1]. Copyright Wiley-VCH Verlag GmbH & Co. KGaA.

**Fig. 2.** Bimodal microscopic images of CaCO₃ particles: Scanning electron microscope (SEM) images show the 430 nm vaterite containers after their synthesis (a), and the calcite single crystals after recrystallization (c). The 2-photon excited fluorescence images show a single vaterite container loaded with the Rhodamine Rh6G dye (b); after recrystallization, only a residual amount of dye attached to the calcite edges could be observed (d). The insets in (b) and (d) show fluorescence intensity profiles along the marked axes. Figure reprinted from [50]. Copyright Wiley-VCH Verlag GmbH & Co. KGaA.
the vaterite particles is determined by the supersaturation level of the dissolved amorphous CaCO$_3$ [48]. The final size of the vaterite particles depends strongly on the concentration of the reagents, the solubility of the salts, the reaction time, and the rotation speed during mixing. It was shown that at high salt concentrations (1 M), at stirring rates of up to 1500 rpm, and reaction time of about 2 min, the size of CaCO$_3$ particle was reduced to 3 μm [49]. Recently, size control of spherical vaterite particles in the range from 10 down to 0.4 μm (Fig. 2a) has been achieved in a mixture of water and ethylene glycol (EG) [50]. The presence of EG during synthesis of CaCO$_3$ particles diminishes the molecular diffusion, reduces the rate and the probability of nucleation, which leads to stabilization of the particles. Depending on the size of the particles, the pore size can be varied from 10 nm to 60 nm. Thus, with pore sizes well above 10 nm, the CaCO$_3$ particles are not nanoporous, but can be considered mesoporous [51]. In 430 nm vaterite particles, the average pore size is ~30 nm [52]. In 4 μm-particles it is ~35 nm, and 18 μm-particles have a pore size distribution ranging from 20 to 60 nm [51]. Thus, it appears that upon decreasing the size of the particles to less than 4 μm, the pore size reaches an average value of 30–35 nm. Biocompatibility tests for vaterite particles indicated no cytotoxicity and no influence on viability or metabolic activity [52].

### 2.1.3. Manganese carbonate (MnCO$_3$)

The procedure for manganese carbonate particles synthesis is similar to calcium carbonate synthesis described before, using salt solutions which contain appropriate ions [21]. The acidic manganese sulfate solution is added to NH$_4$HCO$_3$ in a volume ratio of 1:1; the stirred mixture is then aged at 50 °C for 16 h. The resulting MnCO$_3$ particles have round shapes, and their diameters range from 1.85 μm to 0.3 mm [9].

### 2.1.4. Cadmium carbonate (CdCO$_3$)

This type of particles is often prepared according to the method described by Janevovic and Matijevic [53]. A 10 M urea solution aged for 24 h at 80 °C is quickly added to a preheated 20 mM CdSO$_4$ solution. This procedure leads to monodisperse cubic CdCO$_3$ crystals with a linear size of about 2.5 μm. Temperature has been found to be critical for the shape of the CdCO$_3$ crystals. Thus, in the range of temperatures used (70–75 °C), large polydisperse spherical CdCO$_3$ particles (4–6 μm) are formed [21].

### 2.1.5. Synthesis of dextran-hydroxyethyl methacrylate (dex-HEMA) microgels

Dex-HEMA was slowly introduced with a pipette into a PEG solution under slow continuous stirring in a glass beaker. Upon formation of a homogeneous dispersion, 35 ml dimethyl aminoethyl methacrylate (DMAEMA) was added, and the mixture was slowly stirred for 1 min. In the next step, radical polymerization was started by addition of N,N,N,N′,N′-tetramethylethylene diamine (TEMED) (100 ml; pH neutralized by 4 M NaCl) and potassium persulfate (KPS). The mixture was stirred for 15 min, and then allowed to stand for 30 min. The resulting microgels were centrifuged/washed with 20 ml water (twice) and stored in 5 ml water at 20 °C [25].

### 2.1.6. Fabrication of alginate hydrogel microspheres

An emulsification method was used for the preparation of alginate hydrogel microspheres. Briefly, 50 g of 3 wt.% sodium alginate aqueous solution was dispersed in 75 g of isooctane containing 1.7 g of the surfactant SPAN 85 (sorbitan trioleate) using an ultrasonicator at 60% power for 5 min. A solution containing 0.9 g of Tween 85 (polyoxyethylene sorbitan trioleate) in 5 g of isooctane was then added to the emulsion and ultra-sonicated at the same power for another 5 min to achieve stable water/oil emulsion droplets. Following this step, 20 ml of aqueous solution containing 10 wt.% of calcium chloride was added. The microspheres were then rinsed three times with deionized water by centrifugation.

### 2.1.7. Formation of mesoporous silica particles

The reaction mixture for the synthesis of mesoporous silica particles was prepared directly in an autoclavable polypropylene bottle at –30 °C. A total of 19.6 g of CTAB followed by 10 g of solid Na$_2$SiO$_3$ were dissolved in 350 ml of distilled water, resulting in a clear solution. Then, 25 ml of ethyl acetate were quickly added under stirring. After 30 s, the stirring was stopped, and the mixture was allowed to stand at ambient temperature for 5 h; after this period of aging, the pH reached a constant value. During the aging process, organic solvents were allowed to evaporate through leaks in the cap of the bottle. The resulting solids were recovered by filtration of the warm reaction mixture, extensively washed with distilled water and ethanol, and dried at ambient temperature. The template was removed by calcination at 600 °C for 20 h in flowing air [54].

### 2.2. Stability and decomposition of particles

The most important parameter of colloidal systems is the stability of particles which depends on the degree of ion adsorption, and, therefore, on the zeta potential. The smooth particles melamine–formaldehyde, silica, and polystyrene have relative high zeta-potentials (~50 mV), and, therefore, high stability. The high surface charge prevents these particles from aggregation; furthermore, their morphology is unchangeable. In contrast to smooth particles, porous ones, for example CaCO$_3$, are not stable in aqueous solution or during extended steps of synthesis, and they transform to the more stable calcite structure [50]. Early studies have demonstrated the influence of aqueous and PBS buffer on CaCO$_3$ morphology (Fig. 2c). In order to prevent recrystallization, particles can be either transferred to pure ethanol or dried at 70 °C for 1 h and then kept as a powder.

In contrast to the stability of the aforementioned particles, the decomposable ones can be treated to obtain polymeric hollow carriers by immersion into appropriate solvents. These chemicals are listed in Table 1. Depending on the material of the particles, the dissolution process lasts from several minutes to several hours, and this is followed by thorough washing with water. When using chemicals for dissolution of particles, caution should be taken into account, since in certain cases these may include acidic media or might have toxic effects. Specifically, silica particles usually are decomposed in 0.3 M HF consisting of a mixture of HF and NH$_4$F. N,N-dimethylformamide (DMF)/dimethyl sulfoxide (DMSO) and tetrahydrofuran (THF) have been used for melamine–formaldehyde and polystyrene particles, respectively. Calcium carbonate particles have been found to dissolve by immersing them into acidic media, such as NaCl/HCl buffer (pH 2.0), as well as by complexation with the chelating agent EDTA in appropriate solution (0.2 M, pH 7.5) [31,55].

The gold core was removed and transformed in a gold cyanide complex according to Eq. (1).

$$2Au + 1/2O_2 + H_2O + 4KCN \rightarrow 2K[Au(CN)_2] + 2KOH$$ (1)

Red blood cells can be removed by exposure to a mixture of 140 mM NaCl and 1.2% NaClO.

### 2.3. Release from particles

The release of biomolecules, such as small drug molecules from particles [56], is governed by an interplay of drug desorption and carrier dissolution [57]. This process usually is very slow, especially in the absence of a payload-specific solvent, but could be increased when the carrier size is reduced. During the desorption/adsorption process, molecules get detached and reattached until a dynamic equilibrium is reached, causing small oscillatory modulations in the release curve [57–59]. Furthermore, the particles themselves can be degraded or dissolved by the surrounding medium, causing the payload to diffuse from the carrier.
The typical time-dependence of release from porous silica particles has been discussed earlier. The release is observed to be very fast during the first few hours (releasing up to 75%), and then the release curve reaches saturation, with around 10% of encapsulated material remaining bound for longer time. This has been demonstrated for the release of doxorubicin from silica particles [60]; Brilliant Blue F from porous silica particles [61]; Orange 2, Rhodamine 6G, and doxorubicin encapsulated in millimeter-sized silicagel particles [62], and for gentamicin released over a period of 20 days from mesoporous silica [63].

When the particles start to dissolve, the release is enhanced and reaches 100% when all carriers are dissolved. [64] This is commonly observed for biodegradable carriers such as the well-known gelatin, polyalginate capsules. One of the typical examples is poly(D,L-lactic-co-glycolic) acid PLGA particles: Continuous release of serum albumin from biodegradable PLGA particles during 60 days has been demonstrated [65]. Also, a number of particles based on carbonate and gelatin can dissolve at low pH in the stomach, and cargo is rapidly released.

2.3.1. Release by recrystallization at normal pH

The combination of dissolution and desorption processes is a unique property of slowly dissolving crystals, or it happens during recrystallization of the containers. This effect can be seen on the calcium carbonate particles in the vaterite phase. The vaterite particles (Fig. 3a) can dissolve or can get transferred into a crystal phase (to calcite, Fig. 3b), where the external layer of vaterite starts to ionize [55]. The simultaneous occurrence of the two processes (desorption and dissolution) offers the opportunity to realize the delayed burst release.

In previous work [52], the authors demonstrated the release based on this process and studied it for encapsulated cargo of different sizes: for high molecular weight tetramethyl-rhodamine isothiocyanate TRITC-dextran [52], for the low molecular weight model substance Rhodamine 6G [52,66], and for the drug photosensitizer “Photosens” (a mixture of sulfonated aluminum phthalocyanines AlPcSn, with n = 2, 3 or 4) [55]. The size of particles can vary between 400 nm and 4 μm. Typically, the recrystallization process starts when vaterite particles are immersed in an aqueous solution: This formation of particles can be explained by the dissolution and ionization of the external layer of vaterite [67].

Particles can be loaded with molecules of low molecular weight (e.g. Rhodamine Rh6G). The dye is slowly released until equilibrium is reached. An interesting feature is that after three to four days, the recrystallization process accelerates, raising the amount of dye molecules in the aqueous solution to 40%. The complete recrystallization of all vaterite spheres takes place within less than one week (Fig. 3c).

Immersion of the vaterite particles in ethanol induces slow release via adsorption–desorption. The total amount of released dye was reported to be around 22% after one day, and did not increase significantly during one week of monitoring. This points to the formation of the desorption–adsorption equilibrium without crystal phase transition. Colloidal stability of containers results from their high negative Z-potential. For particles loaded with high molecular weight molecules, no recrystallization was observed regardless of the media used (experiments lasted for one week). In this time-frame, only slow release is possible, while recrystallization is expected to end after one week, possibly due to strong attachment of TRITC-dextran to the surface of vaterite. This view is supported by the observed strong negative Z-potential, which is thought to prevent aggregation; in addition, TRITC-dextran slows down the rate of carrier dissolution. [68,69]. Thorough understanding of these mechanisms can be used to tune the release dynamics.

2.3.2. Release at low pH

Dissolution of calcium carbonate containers at pH < 5 was used for creating the matrix of encapsulated drug [70], or for simple release of the compound [66]. This pH-dependence of the dissolution of particles allows to adjust conditions for controlled and targeted cellular delivery. This is due to the fact that the microenvironment in tumors is typically more acidic than that in normal tissues [71]. Thus, during endocytosis, the pH level decreases in the endocytotic vesicles to around 5.0, in comparison to 7.4 in the cytoplasm. This can trigger the release of loaded or attached molecules because the acidic pH in these vesicles is maintained by proton pumps; therefore, pH-dependent release would be ideally suited for targeting viable cancer cells [72].

This pH-sensitivity could be functionally exploited for delaying drug release from particles, which is particularly relevant in the bloodstream (pH ≈ 7.4). More detailed data on release of low molecular weight molecules (“Photosens”) from vaterite microparticles was recently published [73]. The time course of release was determined starting from 5 min after immersion and was continued up to almost one week. In different experiments, the pH value of the ambient medium was varied, while Photosens was confined inside the carrier particles and was released during the transformation from vaterite to calcite particles, or amorphous calcium carbonate, except for residual amounts of the drug which reattached to the external surfaces. An interesting result here is the overall tendency of particles to dissolve quickly with decreasing pH, thus leading to formation of calcite crystals, and/or amorphous CaCO₃. This points to increasing solubility of calcium carbonate with decreasing pH, in good agreement with the observed 3.7-fold relative difference in solubility between vaterite and calcite [74].

The time interval until complete dissolution of the vaterite crystals decreases significantly upon decrease of pH. As such, the drug release is governed to a larger extent by the dissolution of the particles. At

![Fig. 3.](image-url)
intermediate acidity in the range of pH 6.5 to 5, the dissolution times are shorter, and the vaterite particles dissolve completely already after one day. The shortest time of the phase transition was observed to take place at quite a low pH range (5 to 4.5), where both sizes of vaterite particles dissolved within the first five minutes. This led to immediate (burst) release of cargo from particles.

The above-described processes can be interpreted as follows: Drug release is first controlled by desorption. Complete release can then take place during the phase transition. In contrast to drug delivery applications performed in an open environment, a complete dispersion of the loaded compound can be expected after the dissolution of the vaterite particles. The differences between the time-dependent release curves of large and small particles are twofold: a) the time for reaching saturation increases from ~1 to ~4 days, and b) the saturation level is lower for larger particles because of their potential for an enhanced re-adsorption due to more efficient re-attachment to newly formed calcite structures [55,66].

2.4. Microparticles as carriers of enzymes

Primarily, the role of particles has been considered here as templates for building polyelectrolyte multilayer capsules. However, the particles on which capsules are templated can also be used as carriers of enzymes. The polyelectrolyte multilayer shell can be constructed over such particles loaded with enzymes, providing an important functionality — protection of enzymes, just as in the case of capsules [75].

Capsules carrying enzymes in their pores have been prepared based on calcium carbonate matrix. Enzyme for the substrate can be added directly into a solution, although co-encapsulation of the substrate in liposomes, which are capable of carrying smaller molecules, has been shown applicable for initiating enzyme-catalyzed reactions. Peroxidase has been used in the above studies as the model system, while release of substrate from liposomes attached to the particles has been performed by ultrasound (Fig. 4).

Besides calcium carbonate particles serving as a reservoir for biomolecules, calcium phosphate cores are promising candidates for intracellular delivery of genetic material [76]. A number of studies have been devoted to the packaging of DNA, RNA, and oligonucleotides molecules in calcium phosphate nanoparticles [77]. The advantage of using calcium phosphate particles is that the size of these carriers can be adjusted on the ten-to-hundred nanometer scale which is important in cell transfection [78]. Furthermore, these carriers exert a protective role on the embedded material against the cellular environment, and allow for specific targeting functionalization of the calcium phosphate surface [79]. Designing artificial cells, initiating an enzyme-catalyzed reaction with protection of the enzyme from degradation in the surrounding medium, or drug delivery are some of the application areas of these approaches.

Fig. 4. Schematic representation of enzyme-catalyzed reaction in porous microparticles. Confocal transmission (middle row) and fluorescence microscopy images (bottom row) were taken at the indicated time intervals. Figure reprinted from [75]. Copyright Wiley-VCH Verlag GmbH & Co. KGaA.
2.5. Biomolecule-based particles

Particles are also ideally suited to the delivery of some of the most relevant biomolecules. For example, insulin [80] or hemoglobin can be directly loaded into particles by calcium carbonate [81,82] or manganese carbonate templating [83]. Such particles could also be suitable for biocompatible polymer coverage using the layer-by-layer (LbL) assembly approach. The advantage of microparticles of high deformability in combination with high loading capacity of hemoglobin, mimicking red blood cells [84], is the simplicity of manufacture as well as the fact that the particles themselves determine size and surface properties of the protein carriers.

3. Capsules

3.1. From particles to capsules: dissolution of templates

This overview covers the basics of capsule preparation and the role of particles in the properties inherited by capsules. Encapsulation efficiencies for capsules which are templated on either porous or smooth particles differ strongly. There are three basic methods of capsule formation (Fig. 5): i) co-precipitation whereby cargo molecules are loaded on the particles during their synthesis; ii) adsorption of cargo on pre-synthesized particles which are then used as template for the assembly of the polyelectrolyte shell; and iii) the loading of pre-formed hollow polyelectrolyte capsules.

Encapsulation during synthesis is widely used for molecules of high molecular weight, such as proteins. This method is based on the addition of cargo material to the salt solution reaction mixture with subsequent incorporation of the target molecules into the particle matrix. Porous particles like CaCO₃ have been widely used for loading of molecules by this technique and exhibited a relatively high efficiency as compared to the adsorption method [85]. However, co-precipitation might decrease, or increase, the size of particles, change the surface morphology and the shape of the particles which could then be significantly different from particles produced in the absence of loaded molecules. Work of She et al., comparing encapsulation of molecules by both methods, i.e. co-precipitation and adsorption, demonstrated that loading efficiency of TRITC-BSA was higher in the co-precipitation approach [58]; we have recently obtained similar results for encapsulation of the therapeutic enzyme l-asparaginase [85]. The efficiency of adsorption of

![Fig. 5. Schematic of loading based on smooth and porous particles, using the layer-by-layer (LbL) technique for capsule shell formation, followed by decomposition of the core material.](image-url)
molecules on a porous matrix depends on the surface area and the size of pores: The higher the surface area and the larger the pore size, the higher the number of molecules contained within the particles. The surface charge is another parameter which determines the amount of molecules to be loaded into the particles. Two fluorescent markers were loaded into the vaterite containers: Rhodamine 6G (480 Da) and TRITC dextran, whereby the attachment of the dextran group increases the molecular weight by one order of magnitude to ~4.4 kDa. In this way, the uptake of payload of different molecular properties could be quantitated: For the high molecular weight cargo, the loading efficiency was 157 ± 24 μg/mg, whereas for the low molecular weight substance Rhodamine 6G the loading efficiency depended on the number of washings. One to four repetitions resulted in efficiencies from 8 ± 2 μg/mg to (7 ± 2) · 10⁻² μg/mg. These efficiencies are of the same order as those of other substances loaded via adsorption to porous carriers [86-88]. Loading in pre-assembled capsules strongly depends on the type of cores used for the assembly of capsules. In the case of porous templates, the first polyelectrolyte layer determines the structure of the polymeric shell which is usually thick and appears as a polyelectrolyte network [31]. This polymeric matrix can be used for the loading of molecules by the adsorption technique, similar to the use of simple porous particles. In contrast to porous cores, the smooth ones display very different features because the polymers assembled onto the smooth surface form a thin polyelectrolyte shell that is only a few nanometers thick. The loading into polyelectrolyte capsules formed on smooth particles can be performed by changing physical parameters of environmental conditions. However, encapsulation methods based on variation of pH, ionic strength, and solvent are not easy to implement, since for all of them a certain compound (acid/base, salt, organic solvent) has to be added to the capsule suspension, which can induce deformations through a transient osmotic shock.

The applicability of the three methods described in the previous section is restricted to certain compounds and/or capsule systems. Furthermore, the distribution of the large (above 4 kDa) entrapped molecules would be higher in the capsule wall than in their interior [89]. Of high importance for biomedical applications (e.g. laser-induced release) is the response to thermal shrinking of microcapsules functionalized with gold nanoparticles (AuNPs). Heating of (PDADMAC/PSS)n (where n is the number of layers) microcapsules has been shown to induce the reorganization of loosely arranged polyelectrolyte layers into a denser structure [90]. Thermally increasing the entropy incites more efficient, stronger electrostatic interactions between the oppositely charged layers (enthalpy increase), accompanied by densification of the polymeric shell and subsequent reduction of its inner volume. The shrinking behavior can be adjusted by the total number of bilayers, while its direction is predetermined by an odd (swelling) or even (shrinking) layer number due to the interplay of hydrophobic and electrostatic forces [91]. While denser microshells tend to be mechanically more stable, their permeability decreases with increasing bilayer number [92].

Modifications of the particle surface, or surface functionalization, are different for smooth and porous particles. The difference between polyelectrolyte capsules based on porous particles and non-porous ones regarding their optical properties was demonstrated in several works [33,93,94]. Polyelectrolyte capsules modified by gold [33,93] or silver nanoparticles possessing plasmon resonance effects have been reported [94], revealing that the amount and distribution of metal nanoparticles in shells of polyelectrolyte capsules depend on the nature of the template. Capsules based on porous particles have better adsorption properties and exhibit higher surface-enhanced Raman spectroscopy (SERS) sensor effects [93]. If the number of nanoparticles used for adsorption also was controlled, capsules built on porous particles demonstrated that nanoparticles form larger elaborate aggregates [30,94,95] versus smaller aggregates on capsules with smooth surface [5,96], which provide other interesting optical properties, such as control of absorption in the near-IR range [33,94].

### 3.2. Release of cargo from capsules

Release of payload molecules from microcapsules, which can be initiated by various stimuli (pH, ionic strength, solvent, and temperature), substantially depends on the morphology of the core. These stimuli have been shown to alter the permeability of capsule shells in a controllable and reversible way by creating tiny pores in the polymeric structure and thus facilitating diffusion of molecules. Such methods are broadly used for small-molecule drug delivery. However, application of chemical approaches might be limited in cellular systems where drastic changes in the chemical composition of the solution are not a viable option.

Fig. 6 presents data of a comparative study on release from microcapsules templated on either smooth silica particles (Fig. 6a,b), or on porous calcium carbonate particles (Fig. 6c,d). In the first case (Fig. 6c), thermally shrunk microcapsules have been used. These capsules can be functionalized with nanoplasmonic gold nanoparticles which serve as centers of localized temperature rise and permeability change. A distinctive feature of such capsules is the possibility of complete control over spatial and temporal release profiles. Alternatively, capsules templated on porous calcium carbonate are often used for release which is achieved through dissolution of the core material upon decomposition of the surrounding biodegradable polymers (Fig. 6d). We note that multifunctionality of release is yet another aspect which receives increasing attention [97]. In this regard, multi-functionality and multi-compartmentalization are essential properties [98,99].

### 4. Complex morphologies of particles and capsules

#### 4.1. Multi-compartment particles and capsules

The multi-compartmentalization, which was mentioned in Section 2.4 and which represents a higher level of complexity, is a promising way to increase functionality of capsules destined for cargo transport into cells. Both particles and capsules can constitute multi-compartmentalized carriers, whose main advantage is simultaneous delivery of different molecules, targeting [100,101] using one carrier entity. The aforementioned encapsulation methods can also be adapted to the construction of a certain compartment of multi-compartment capsules: a) assembly from pre-formed different sub-compartments; b) synthesis of particles on which capsules are assembled; c) a combination of synthesis and assembly from pre-formed sub-compartments [102]. Of all multi-compartment capsules structures, the concentric and pericentric ones are most frequently used. Strategies for the construction of concentric capsules were demonstrated by Kreft and colleagues [103]. They demonstrated fabrication of the shell-in-shell capsules with two concentric calcium carbonate shells. These shells were independently loaded with biomolecules separated by a semipermeable polyelectrolyte membrane. Furthermore, this system has been tested as a reactor where the coupling of an enzymatic reaction through a semipermeable membrane was achieved via encapsulation of a substrate in one ball and an enzyme in the other one. Triggered release of the inner compartment of two-compartment capsules has been demonstrated [104]. The inner capsules were modified by metal nanoparticles, and upon near-infrared laser illumination the inner part of such dual-capsules was released. Three coupled enzymes were encapsulated into concentric multi-compartment CaCO₃ particles via co-precipitation [105]. These enzymes were incorporated in separated compartments together with two spacing compartments made of bovine serum albumin (BSA). Strong influence of the spacing between compartments on the reaction kinetics was monitored through confocal laser scanning microscopy (CLSM). The cascade reaction catalyzed by enzymes introduced in separate polymersomes, without artificial transport mediator located in the membrane, has been demonstrated by Kuiper et al. [106].

Pericentric microcapsules are another example of multi-compartment carriers. This type of complex capsules can be constructed by
surrounding a large container with small ones. One approach to manufacture pericentric capsules is to use polyelectrolyte-coated particles modified via electrostatic attraction between small and large counterparts [86]. Porous colloids, such as calcium carbonate cores, can also be applied for assembly of pericentric compartments due, in a large part, to their large surface-to-volume ratio and plenty of cavities [74,80]. By using pericentric structures made of calcium carbonate as inner core, filled with an enzyme surrounded by liposomes containing a substrate, an enzyme-catalyzed reaction was performed [74]. By disrupting liposomes via ultrasonication, the course of the reaction in large particle volumes was observed by CLSM. Detachment of small containers from the inner particles has been demonstrated to permit control of the release of small carriers via biodegradation of polyelectrolytes upon enzymatic disruption [80]. An overview of various morphologies of multi-compartment particles and capsules has been recently presented [107], while different ways of inwards build-up of concentric polymeric (polyamines and polycarboxylic layers) capsules templated on agarose or alginate hydrogels in an organic phase [108] have been shown. Generation of hybrid liposome–polymersome multi-compartment assemblies [109] has been also demonstrated.

4.2. Anisotropic capsules and anisotropic carriers

Of all parameters pertaining to particles as drug delivery vehicles, the shape has been shown to be a valuable characteristic which influences the internalization and further fate of the particles in the cell. Recent findings suggest that the geometry of particles could change their efficiency of uptake into cells in the phagocytotic process [110]. The carrier geometry of polymer particles was found to influence endothelial targeting in the vasculature, and the rate of endocytosis and lysosomal transport within endothelial cells [111]. Specifically, non-spherical (elaborated, rod-like) particles revealed higher intracellular transport as compared to that of spherical carriers [112,113]. Attachment of particles to specific sites of cells can be tuned not only via surface chemistry of vehicles, but also by changing the geometry of particles even at the nanoscale [114–116]. Thus, future trends in particle-directed drug delivery are toward novel methods of synthesis and elaboration of anisotropic particles which in specific cases resemble natural vehicles like bacteria [117]. Not only particles of certain shapes can be used as delivery system, but also capsules templated on specifically shaped particles would be beneficial to certain applications [118,119]. The flexibility of polyelectrolytes molecules allows reproducing a number of capsule shapes after decomposition of the initial solid cores [119–121]. Shchepelina et al. compared the morphology, mechanical properties, and permeability of hydrogen-bonded layer-by-layer (LbL) microcapsule shells assembled on squared cadmium carbonate particles against the same shells assembled on spherical silica [122]. The patterned template-assisted assembly of the cubic micro-particles driven by the competing capillary, Coulomb, and van der Waals forces was compared to the traditional spherical colloidal microparticles that had been studied by Lisunova et al. [123]. A set of shaped (spherical, elliptical, and squared) CaCO3 particles was synthesized through changing the stirring speed, time, pH value, and salt ratio (Fig. 7). The particles were further used for building polyelectrolyte capsules loaded with labeled dextran.
molecules which replicate the initial shape of CaCO₃ microcores after decomposition with EDTA [22].

Another promising concept for manufacturing anisotropic carriers is multi-compartmentalization under conditions where at least two particles/capsules interact with each other [98,107,124,125]. Such anisotropic hierarchy of nano- and microparticles/capsules makes it possible to generate properties of carriers that could not be achieved using conventional methods. Bio-inspired reactions occurring in the confined volume of particles/capsules have been carried out by applying multi-compartment approaches [107,117,125,126]. Advantages of the artificial compartmentalization strategy are the separation of large and small molecules, while at the same time carrying them in one entity with protection from the outside environment, and conduction of cascade reactions [127]. Furthermore, multi-compartment particles/capsules are predicted to be of potential use in theranostic applications [127]. Furthermore, multi-compartment particles/capsules are predicted to be of potential use in theranostic applications [127]. Furthermore, multi-compartment particles/capsules are predicted to be of potential use in theranostic applications [127]. Furthermore, multi-compartment particles/capsules are predicted to be of potential use in theranostic applications [127]. Furthermore, multi-compartment particles/capsules are predicted to be of potential use in theranostic applications [127].

5. Summary

In this overview, we highlighted methods of preparation and major physico-chemical properties of particles used as delivery carriers and particles serving as templates (cores) on which polyelectrolyte multilayer capsules are assembled. With regard to capsules, the choice of the template material determines, first and foremost, its physico-chemical and biological properties. Porous particles are presented, as well as those possessing smooth surface. Non-particle, hybrid composites (red blood cells, virus-like particles, and liposomes) are also briefly discussed. It is important to emphasize that properties of microcapsules, which are built on such templates, substantially depend on the templates on which the former are assembled, as also overviewed here. Finally, reference is made to emerging trends in the development of particles and capsules, the production of multicomartment and anisotropic particles, and particles of controlled size, porosity, and monodispersity.

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