The Synthesis of Alginate Microparticles and Nanoparticles

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Abstract

Alginate is a natural polysaccharide that is widely used as a component of pharmaceuticals and in food industry. Alginate particles can be used for encapsulation of substances with the necessity of prolonged release. They can also provide appropriate microenvironment for cells. Here the methods of the synthesis of alginate beads, micro- and nanoparticles are reviewed with special attention to the calcium alginate ones. The results from publications that did not deal with alginate particles but, to our opinion, could be applied in this field are also included in order to give an outline for possible future research. The suggested applications of the particles are mentioned as well. The two main methods for the synthesis of calcium alginate particles are internal and external gelation, but the external gelation techniques can be themselves subdivided into several subtypes. Currently, a technique being able to produce alginate nanoparticles with any desirable size does not exist. We analyze the possibilities of employing aerosolization method for this purpose. The potentials to overcome the problem of burst release of the encapsulated substances by means of cyclodextrin inclusion complexes and employing additional crosslinking agents are also discussed. The clinical application of alginate nanoparticles is still limited because of the burst release of encapsulated drugs and the poor size control of the particles formed. Further research must concentrate on overcoming these problems and on topical application of alginate particles without entering bloodstream rather than on investigation of model drug release in vitro without taking the above-mentioned problems into account.

Keywords: Calcium alginate; Alginate beads; Alginate microparticles; Alginate nanoparticles; Drug encapsulation; Particle size control; Aerosol

Introduction

Alginites are polysaccharides. The commercially available ones come from brown algae. They are linear copolymers of \((1\rightarrow4)\)-linked units of \(\beta\)-D-mannuronic acid and \(\alpha\)-L-guluronic acid. The molar ratio between them and their distribution along the commercial polymer depend on the algal source, its location, age, collection season and extraction technique. Guluronic acid residues can form so-called egg-box complexes with calcium ions or some other divalent metal cations leading to gel formation (Figure 1). The name ‘egg-box’ is used because, if depicted schematically, the cations look like eggs situated inside puckered boxes formed by four guluronic acid residues of two superimposed chains. Mannuronic acid residues have much less affinity to metal ions [1]. Barium ions have more affinity to alginate than the calcium ones. If reacted with calcium alginate at lower concentrations, they create new gelling junctions. At higher concentrations, barium ions also displace calcium ions from existing junctions [2].

![Figure 1: The egg-box model.](image-url)
To extract alginate, algae are usually washed with organic solvents and water, dried and milled. After acid pretreatment, alginate is solubilized with Na$_2$CO$_3$. The crude extract is concentrated, dialyzed against water and then freeze-dried or precipitated with ethanol. Acidification or treatment with Ca$^{2+}$ can be used instead. Brown algae are abundant in nature; however, the possibility of cultivating them exists as well [1].

The viscosity of aqueous sodium alginate solution rapidly increases with its concentration. For example, the addition of 10% of alginate to water leads to a ~100-fold increase in viscosity. But the poly electrolyte nature of alginate has little effect on its hydration, and in the above example less than 3% of the water molecules present in solution is involved in alginate hydration. Such a large viscosity increase is determined by the polysaccharide network, with large bulk-like water pools present between the polysaccharide chains [3].

Sodium alginate may act as a mucoadhesive polymer. A comparative study of adhesion between buccoadhesive compacts and pig buccal mucosa or sodium alginate solution revealed that the results were of similar performance [4]. Sodium alginate was proposed as a mucoadhesive component of a nasal gel [5] or in buccal patches containing salbutamol sulfate [6]. Sodium alginate conferred in situ gelling mucoadhesive properties and retarded drug release from liquid rectal suppositories. These suppositories were successfully tested on Guinea pigs to alleviate symptoms of histamine-induced bronchospasm [7]. Sodium alginate was also evaluated as an excipient in salbutamol sulfate sublingual films [8] and tablets [9]. However, drug release was found to be too slow in the films [8] or too rapid in tablets [9]. Only salbutamol sulfate tablets formulated from granules containing mastic and sodium alginate excelled commercial tablets in the terms of drug release when tested on rabbits [10]. In combination with hydroxypropyl methyl cellulose and propylene glycol sodium alginate was used in the formulation of terbutaline sulfate sublingual films [11]. Sodium alginate could be also used as a component of plugs for water-soluble parts of crosslinked gelatin capsules containing pellets with encapsulated salbutamol sulfate. The plug absorbed the surrounding fluid, and began to release the drug through the swollen matrix and was finally ejected out of the capsule by erosion of the material. The usability of the system was shown on rabbits [12].

Chitosan-alginate complex was proposed as an excipient for orodispersible tablets, and their disintegration time was so short that it was even referred to as a ‘super dis-integrant’ [13]. One optimized formulation containing the excipient for 5-fluorouracil tablets, suitable for trans buccal and rectal drug delivery, contained this chitosan-alginate complex along with the same components un-complexed in order to avoid burst release and to improve the mucoadhesive properties [14].

Sodium alginate itself also has a therapeutic effect. When admixed to foods for diabetic human patients, it decreased gastric emptying rate and rises glucose in blood, serum insulin and plasma C-peptide levels [15]. Orally administered, sodium alginate significantly alleviated small intestinal enteritis in rats, caused by treatment with the anti-inflammatory drug indomethacin, and this relief seemed to be independent of the sodium alginate viscosity administered [16]. Oral disposal of sodium alginate to rats with colitis led to a significant reduction of colonic damage, decreased lesion formation [17,18] and inhibited mucosal injury [17].

Orally administered alginate oligosaccharide obtained from hydrolysis of sodium alginate by *Bacillus subtilis* improved histopathological and biochemical parameters of mice having ovalbumin-induced asthma in a dose-dependent manner [19]. Rats fed with sodium alginate drank more water, and their urine volume and pH rose sharply. In contrast, calcium alginate caused very little changes in the same parameters [20].

Commercially available calcium alginate swabs were used for sampling nasal flora for subsequent DNA extraction [21]. The mucoid exopolysaccharide produced by the pathogenic bacterium *Pseudomonas aeruginosa* is alginate, but it has low immunogenicity if it is not conjugated with a carrier protein. Even in the form of conjugate it is non-toxic if administered intraperitoneally to mice or guinea pigs and non-pyrogenic if administered intravenously to rabbits [22]. When alginate beads with encapsulated tumor cells were implanted to mice, a process of angiogenesis was observed in the implants zone. The beads were prepared from commercially available alginate [23]. It may be considered as a further proof of alginate biocompatibility.

Sodium alginate was also shown to be beneficial for agriculture. Being administered as dietary supplement to the white shrimp *Litopenaeus vannamei*, it acted as an immunostimulant and improved its resistance against the attack of *Vibrio alginolyticus* bacterium [24]. Sodium alginate digested with alginate lyase promoted root elongation of rice, carrot [25], lettuce [26] and barley plants [27,28] even under hypoxic conditions [28]. It was hypothesized that digested alginate might initiate some signal transduction pathway [27,28]. Under hypoxic conditions digested alginate also caused enhancement of the activity of the enzymes regenerating NAD$^+$ [28].

Alginate oligomers promoted the germination of unhulled rice and Komatsuna seeds as well as tobacco callus differentiation. The mixture of oligomers was assumed to contain so-called oligosaccharides, an oligosaccharide inducing unusual proliferation and/or differentiation of plant cells. There are several kinds of oligosaccharides. They act as a chemical signal for the stimulation of hormone synthesis [29]. The promoting effect of alginate oligosaccharides on root formation and growth in rice was mediated by endogenous indole-3-acetic acid [30].

But proliferation of the microalga *Chlamidononas reinhardtii* was repressed by the same oligomers [29]. It should be noted that only digested sodium alginate shows this effect. And the possibility of alginate degradation by the lyses of soil bacteria is assumed [28]. Therefore, it is speculated that these active substances can be formed from alginate under natural conditions.

Another notable advantage of alginate is its ability to bind micronutrients. Some important Mn, Cu, Zn and Mo fertilizers are MnCl$_2$, CuSO$_4$·5H$_2$O, ZnCO$_3$, and Na$_2$MoO$_4$·2H$_2$O, respectively [31].
Manganese can be complexed with alginate by addition of MnCl₂ to the gelling solution of CaCl₂ or BaCl₂, and slow release of manganese ions from the beads into physiological saline has been reported [32] because the affinity of alginate to Ca²⁺ is higher than to Mn²⁺ [33]. The beads containing Ba and Mn could be used for manganese-enhanced magnetic resonance imaging and were tested on rats [32]. The use of CuSO₄ [34] or CuCl₂ [35] as a gelling solution led to copper alginate hydrogel being able to release copper ions into simulated body fluid [34] or into phosphate buffer (pH 6) [35] in a prolonged manner. The use of basic zinc carbonate for zinc alginate hydrogel formation using internal gelation method has also been reported. The hydrogel was active against E. coli [36]. Molybdenum can be adsorbed by preformed calcium alginate beads preferably in the form of H₂MoO₄ or [Mo(H₂O)₆]³⁺ at pH 2 and released back up to 50% into 0.1M HCl. If radioactive molybdenum is used, the method is suitable for radiotherapy [37]. Copper ions from CuCl₂ may also be adsorbed onto preformed calcium alginate beads [38]. Copper alginate shows activity against Staphylococcus aureus, Staphylococcus epidermidis, Staphylococcus pyogenes and E. coli [34].

Alginic microparticles and nanoparticles are usually used to encapsulate and carry various substances, and the goal of many studies is to achieve sustained release of them. It should be noted, however, that the results of experiments dealing with the release of poorly water-soluble drugs might often be misinterpreted, because the drug not found in the solution is assumed to remain encapsulated in the particles. However, it may decompose after the release or simply precipitate out. Supersaturated solutions with varying extent of supersaturation can also be formed, making the results irreproducible. Special care must be taken in the case of putting the particles into a dialysis bag, because the films act as an additional diffusion barrier. If centrifugation is used for separation of the medium with the released substance from the nanoparticles, the pressure generated during the process can disturb the equilibrium. It can also make difficult resuspending the nanoparticles in the fresh portion of medium for further incubation [46].

For consistency, throughout this review we will use the following terms (even if different names for them were used in the respective publications):

- Encapsulation efficiency: the percentage of the substance that was encapsulated (i.e. not lost).
- Loading efficiency (expressed in percent): the ratio of the weight of the successfully encapsulated substance regarding to the total weight of the particle. Some authors calculate loading efficiency using different formulae, but we will give their values without a special discussion. We have rounded encapsulation and loading efficiencies as well as zeta potentials to the nearest integer values.

- Nanoparticles are considered smaller than 1µm. Microparticles have size from 1 to 1000µm. Beads have size in a millimeter range. We have rounded bead size to the first decimal place.

Also, some authors term their particles ‘microcapsules’. However, we will use this term only if they have demonstrated or at least assumed the presence of a liquid core in their particles. In other cases, we will refer them to as microparticles.

**Alginate-chitosan particles**

Since alginate is a polyanion and chitosan is a polycation, they can form a polyelectrolyte complex upon mixing, provided both of them are charged, i.e. at suitable pH. This mixture can spontaneously form particles. The pKₐ of alginate carboxyl group is close to 5, and that of the ammonium group of chitosan is about 6.2 [1].
Alginate-chitosan nanoparticles were prepared by dropwise addition of a chitosan solution containing glutathione into an alginate solution at pH 4, under stirring. If prepared at 0.75 alginate: chitosan ratio, the formed nanoparticles with encapsulated glutathione had the following characteristics: size 361 nm, polydispersity index 0.33, zeta potential 27 mV [47, 48]. At 1.5 alginate: chitosan ratio, the values were 212 nm, 0.4 and 23 mV, respectively, although the storage stability decreased, making these nanoparticles less suitable for application. In the same way, the pH increase from 5.0 to 6.5 and further caused aggregation [48]. The encapsulation efficiency was 27% [47] or 80% at ratio 0.75 and fell to 1% at ratio 1.5 [48]. The respective investigations were aimed to achieve the synthesis of mucoadhesive nanoparticles with an encapsulated NO donor needed for treat important diseases because of the multifaceted role of NO in vivo. Therefore, encapsulated glutathione was nitrosated inside the nanoparticles by adding sodium nitrite to the solution. S-nitrosoglutathione decomposition at 400 μM was delayed by its encapsulation in the nanoparticles. At 18 μM encapsulated S-nitroso glutathione was not cytotoxic to cultured Chinese hamster lung fibroblast cells (V79), whereas free S-nitroso glutathione was slightly cytotoxic at the same concentration. This assay could enable the use of these anti-microbial nanoparticles in pharmaceutical applications such as wound healing without severe side effects [47, 48].

Later, the same technique was used by the same group to encapsulate mercaptosuccinic acid and nitrosate it inside the particles. In this case, the hydrodynamic size of the nanoparticles was ~750 nm. The encapsulation efficiency was 89%. Burst release of NO in aqueous solution was followed for 4 hours, although the release in the normal mode continued for 6 hours more. These nanoparticles were assayed for topical application for bovine mastitis. The minimal inhibitory concentration of the nanoparticles for Staphylococcus aureus determined in vitro was 125–250 μg/mL. The number of colony forming units was 10-fold and 1000-fold lower after bacteria were incubated with nitrosated nanoparticles at 500 μg/mL for 4 and 7 hours, respectively, compared with bacteria growth in the presence of empty nanoparticles at the same concentration and time. The CFU drastically decreased further upon the addition of a second dose of nitrosated nanoparticles. For E. coli the minimal inhibitory concentration exceeded 2000 μg/mL, i.e. these nanoparticles were inefficient against this bacterium.

The 50% cytotoxicity concentration of the nanoparticles for cultured HEP-2 cells was 640 μg/mL. Chitosan nanoparticles without alginate at the same concentrations of the acid released more NO at higher rates. Nevertheless, it was concluded that NO-releasing nanoparticles might be used to combat bacteria for treating and preventing bovine mastitis [49].

Spherical alginate-chitosan beads with encapsulated lemongrass oil having size of 1.8–2.1 mm displayed significant antibacterial and antioxidant activity. For unencapsulated oil the same activity was observed only at higher concentration. This beneficial action was attributed to the strong interaction between chitosan and the oil. This kind of beads has potential applications as a greener agent for medical purposes [50].

The following advantages of alginate-chitosan particles can be underlined:

- Chitosan can enhance drug bioavailability by its capacity of infiltration into the mucus layer of the small intestine with subsequent opening of tight junctions of epithelial cells [51].
- Unlike calcium alginate, alginate-chitosan polyelectrolyte complex cannot be disintegrated by chelatoring agents.

Their main disadvantage could be the necessity to use an acidic solution of chitosan because of its insolubility at neutral pH.

**Preparation of alginate particles without employing gelation**

Now we describe the techniques for preparation of particles from bulk sodium alginate or its solution as well as spontaneous formation of particles of modified alginic acid in water. The resultant nanoparticles are usually intended to be ready to use. However, dry sodium alginate particles can be later treated with CaCl₂ solution in order to convert them to calcium alginate particles.

There exists a patented technique for producing alginate, cellulose, starch or collagen particles from bulk substances by ball milling with the possibility to control particle size from 100 nm to 50 μm. The resultant nanoparticles containing therapeutic proteins have shown efficacy in treating solid tumors, single dose vaccination, and oral delivery. For instance, tumor-bearing mice that received these nanoparticles containing Texas red and cisplatin showed significant tumor size diminishing. If the same nanoparticles were coupled to dendritic cell-binding peptide and contained encapsulated pneumococcal surface protein A, together with an adjuvant, they were effective to combat the bacterial load of the mice that was reduced (in the terms of infected tissue volume) after exposition to nanoparticles. The nanoparticles produced by the same milling technique were also used to induce passive immunity against anthrax toxin in mice by means of oral delivery of monodonal antibodies developed versus anthrax toxin [52].

Another technique consists in dropwise addition of pure ethanol or acetone to 1% sodium alginate in water containing drug solution in dimethyl formamide. Mixing [53] and cooling down to 3-5 °C is needed during the process. At low mixing speed aggregation was observed [54]. These microparticles can be separated by filtration, washed with the same solvent and dried on air in a heating oven [53] or in a desiccator [54]. Using nitrofurazone as an example of an encapsulated drug, the loading efficiency and yield of microparticles decreased as the particle size increased from 5 to 30 μm with ethanol dripping rate increasing. The presence of 0.1% ammonia [54] (pH 8.9) and of a surfactant was needed in order to avoid particle aggregation in the case of nitrofurazone. Other drugs, viz. acridone, tetracycline, dibazole and metronidazole were encapsulated in the same way (but without ammonia), although the encapsulation conditions needed to be optimized for every drug separately. The yield varied from 31% for metronidazole to 77.5% for tetracycline, and the loading efficiency varied from 2% for metronidazole to 43% for nitrofurazone [53]. In a later publication, the same group reported that in the case of nitrofurazone the...
yield of microparticles was 81% with a loading efficiency of 34%. Spray drying instead of filtration was recommended to increase the yield [54]. The stability of nitrofurazone-loaded microparticles resuspended in water was reported to increase with pH [53]. At 1% and 2% of particles (nitrofurazone concentration was 0.34% and 0.68%, respectively) these solutions were more active against E. coli, P. aeruginosa, P. vulgaris, S. aureus and B. subtilis than aqueous nitrofurazone solution having drug concentration less than 0.02% because of its insolubility. In the case of Candida albicans the same solutions of microparticles excelled in antifungal activity nitrofurazone solutions in DMSO with the concentrations of 1% and 2% [54]. The ability of the particles to form stable suspensions and to enhance drug solubility in water broadens the field of drug application [53,54]. The encapsulated drugs are expected to be more stable under ambient conditions [53].

Spray drying the sodium alginate solution containing the payload (caffeine-loaded peptidic nanoparticles) yielded microparticles having size of about 4µm. The crosslinking with CaCl₂ solution increased their mean size to 7.4µm but decreased their shrinkage and slowed down the release of caffeine into simulated gastric fluids. The particles are potentially bioactive because of the presence of antioxidant peptides [55]. Spray drying the solution containing sodium alginate, pectin and gentamicin sulfate at inlet temperature of 90 °C was used for wound dressing preparation. The volume diameter at the 50th percentile (spanning from 310 to 1003nm for various samples), the width of particle size distribution, water content and drug release rate increased with nozzle spray mesh diameter and with feed solution concentration at constant ratio of the components. Flowability of the powders, the adhesive strength of the gel formed from them in contact with simulated wound fluid as well as its activity at 0.25mg/ml of gentamicin sulfate against Staphylococcus aureus and Pseudomonas aeruginosa showed the opposite tendency. Antimicrobial activity was expressed as the diameter of the zones of clearance around the two samples spotted on agar plates with the spread bacterial culture after 24 hours. For Staphylococcus aureus the activity of two samples was also tested in culture medium after 3, 6, 9 and 12 days of incubation. The particles were mainly spherical, but other shapes appeared when either feed concentration or mesh nozzle increased, and further increase led to large collapsed particles. All the particles were composed of smaller aggregated particles. The encapsulated drug was being released in simulated wound fluid in Franz-type diffusion cells for up to 5 days. Loading efficiency was around 24-27% with an encapsulation efficiency between 70 and 83%, and for all the samples initial burst release was observed. At around 24-27% with an encapsulation efficiency also decreased from 68% to 46%. Mean nanoparticle hydrodynamic diameter also decreased from 559nm to 305nm, and particle formation rate was sped up when substitution degree increased. An unimodal particle size distribution was revealed. In simulated gastric fluid they retained spherical shape and released ~40% of the encapsulated vitamin for 3 hours. But in simulated intestinal fluid they became irregularly shaped, their hydrodynamic diameter was 757nm and burst release of 40% of the vitamin occurred, with 60% of the vitamin released after 7 hours. The nanoparticles can be used as oral carriers for liposoluble nutraceuticals [57].

The great disadvantage of sodium alginate particles is the very limited possibility of prolonged drug release because of sodium alginate solubility leading to fast disintegration of the particles. The particles of modified alginic acid offer convenient manipulation, but the need of its prior chemical functionalization limits the applicability of the technique for non-specialized, e.g. biomedical laboratories.

**Production of Alginate Particles Using Other Particles as Cores**

In this method, sodium alginate is physically adsorbed or covalently linked to the surface of other particles. The particles can also be formed already capped with alginate. In some cases subsequent gelation with CaCl₂ is carried out. For CaCO₃ particles there is a possibility of their generation simultaneously with alginate gelation.

Dropwise addition of chitosan nanoparticles with encapsulated bovine serum albumin modified with rhodamine isothiocyanate to sodium alginate solution at controlled pH yielded negatively charged nanoparticles having hydrodynamic diameter of several hundred nanometers depending on the solution composition. The highest diameters were registered in water. The nanoparticles successfully delivered the protein into cultured cells, with the localization depending on cell type. Significant increase in peroxide production by HCEC cells was observed at 300 and 600µg/ml of empty nanoparticles after exposure for 4 hours. However, there was almost no superoxide production after either 4 or 24 hours of exposure. The metabolic activity of LN229 and MCF-7 cells remained unchanged for up to 72 hours of incubation with the empty nanoparticles. But MDA-MB-231 and HEC cells displayed significantly decreased metabolic activity at nanoparticle concentration above 180µg/ml after 72 hours of exposure, but not after 24 hours. Similar survival decrease at these concentrations of nanoparticles was observed for A549 cells. Dose dependencies acquired after 24 or 72 hours of exposure were almost the same. Survival of HT29 and CaCo2 cells was significantly increased only after exposure to 600µg/ml of the nanoparticles for 72 hours. The nanoparticles have potential applicability as nanocarriers in cancer therapy [58]. A similar technique was used for enoxaparin encapsulation. The proposal was...
to evaluate nanoparticles loaded with this low molecular weight heparin for its oral delivery, controlled and prolonged release in order to improve patient compliance. In this, chitosan nanoparticles were covered with sodium alginate (applied in phosphate buffer) and treated with CaCl₂. Parameters of the optimized formulation were as follows: average size 335nm, spherical, polydispersity index 0.37, zeta potential ~31mV, encapsulation efficiency >70%, drug release in simulated gastric fluid for 2 hours 2%, in simulated intestinal fluid for 14 hours ~60%. Degradation and erosion of nanoparticles was identified as a possible drug release mechanism. The pharmacokinetic parameters of the drug given orally to fasted rats through cannula in a dose of 50mg/kg body weight were improved. Nevertheless, those of intravenously administered free enoxaparin at 1mg/kg were better. 75% of the encapsulated drug applied at 2mg/ml reached across the intestine to the serosal fluid for 90 minutes, as shown in vitro by means of everted intestinal sac model. 900 IU of orally administered encapsulated drug reduced thrombus formation by 59% compared with buffer. Significant uptake of the nanoparticles by the intestinal mucosa for 1 hour was shown by administration of nanoparticles loaded with fluorescein isothiocyanate instead of the drug through gastric cannula to fasted rats. Therefore, the nanoparticles proved their utility as oral delivery vehicle for enoxaparin. Such a vehicle is a foremost requirement for non-invasive and non-hospitalized treatment of vascular disorders (deep vein thrombosis, pulmonary embolism and venous thromboembolism). But subcutaneously injected free drug was even better [51].

Microspheres loaded with dermatan sulfate were formed by its complexation with chitosan with subsequent addition of the resulting solution to sodium alginate solution followed by spray-drying. The spray-dried microspheres were collected in CaCl₂ solution in 90% ethanol and then vacuum-dried. Their properties (including shape) significantly depended on buffer, pH and polyelectrolyte ratio, because these parameters influenced polyelectrolyte complex formation. The objective of the study was to achieve sustained delivery of dermatan sulfate from microspheres intended for its site-specific delivery. The particles had size around 2.3 µm and zeta potential from ~27 to ~48mV. Encapsulation efficiency was up to 96% with loading efficiency of around 4%. They released dermatan sulfate in Tris buffer for more than 10 days. After incubation for 48 hours at 10mg/ml in culture medium they increased growth of cultured mouse fibroblasts (NIH 3T3) up to 170-200%. Free dermatan sulfate at 400µg/ml had the same effect. Therefore, released dermatan sulfate was structurally intact and retained its bioactivity. The microspheres synthesized at pH 5 had slightly higher stimulatory effect than those synthesized at pH 3.5. A clear dose dependence of the stimulatory effect was observed at 1-10mg/ml of the microparticles. If chitosan was omitted, the microparticles exhibited rapid burst release, and partially dissolved microspheres fused and formed a network. The microspheres have great potential to be used in tissue engineering applications because dermatan sulfate is an emerging therapeutic agent for tissue regeneration [59].

Another unusual technique was used for the encapsulation of fungicide prochloraz. To the oil-in-water emulsion of prochloraz stabilized by surfactant cetyltrimethylammonium bromide prehydrolized tetraethyl orthosilicate was added. It resulted in its further hydrolysis and silica shell formation around the pesticide-containing drops. After a drying step the silica shell was further modified with amino groups and coupled with alginate using carbodiimide chemistry. The resulting microcapsules were dried again and treated with metal chlorides in solution. This was done in order to use them in agriculture as a nanofertilizer and nanopesticide at the same time. Their size (after drying and redissolving) was ~2.5µm with homogenous distribution. Fungicide loading efficiency was about 30% (w/w). Prochloraz release for 60 days in 30% (v/v) methanol in water was shown. The degradation rate of free prochloraz increased with temperature and was also high under strong acidic and alkaline conditions. In contrast to this, the microcapsules were insensitive to pH, protected the encapsulated fungicide from decomposition under ultraviolet irradiation, and temperature increase just caused some release alteration. They are expected to increase disease resistance of plants. The delivery system may be extended to other photosensitive or pH-sensitive pesticides [60].
weight was observed. The stimulatory effect decreased with nanoparticle concentration. The aim of the study was to elucidate the stimulatory effect of silver nanoparticles at low concentrations on wheat growth at early stages of ontogenesis. The nanoparticles were referred to as an example of a prolonged action form of a biogenic metal produced by means of ‘green nanochemistry’ and acting as a plant growth stimulant [61]. We have produced calcium alginate nanoparticles with gold core (Figure 2). At first, gold nanoparticles were synthesized by tetrachloroaurate reduction with polyethylene glycol using a method modified from [62]. After physical immobilization of sodium alginate onto their surface and resuspending into water CaCl₂ solution was added. The reaction protocol must be followed very thoroughly in order to avoid extensive aggregation and destabilization of the nanoparticles. The size of the nanoparticles can be adjusted by means of auto metallography. Selection of other volumes of water and CaCl₂ solution allowed us to produce stable aggregates [63].

An interesting dependency of the thickness of a layer of physically adsorbed sodium alginate on magnetite (Fe₃O₄) nanoparticles expressed as particle size on experimental conditions was revealed. The average diameter of the nanoparticles dripped onto a TEM grid was 70nm. But if the same nanoparticles (without their separation from alginate solution) were sprayed onto the grid, their average size became 14nm. Aerosolization of the same core magnetite nanoparticles mixed with sodium alginate solution of another concentration into CaCl₂ solution led to nanoparticles with average diameter of 8nm. If chitosan and sodium tripolyphosphate were used instead of alginate and CaCl₂, the sizes of the resulting nanoparticles were also unequal in the three cases. The nanoparticles were intended for drug delivery or hyperthermia therapy of cancer [64].

By mixing solution containing CaCl₂,doxorubicin hydrochloride and a plasmid encoding p53 with another solution containing sodium carbonate and sodium alginate, nanoparticles containing simultaneously DNA and doxorubicin were prepared. Due to the existence of alginate the growth of CaCO₃ based coprecipitates could be effectively retarded. The resulting nanoparticles had size about 145nm and zeta potential ~ -14mV, and these parameters for the nanoparticles without alginate were 300nm and ~ -25mV, respectively. The encapsulation efficiency was 89% for DNA and 84% for doxorubicin. Loading efficiencies were 10% and 1%, respectively. About 40% of encapsulated doxorubicin were released in water for 48 hours with initial burst release. Proliferation of HeLa cells exposed to the nanoparticles at 37 °C for 48 hours was completely inhibited, and apoptosis was induced. Therefore, they have promising applications in cancer treatments. Cell inhibition rate was dose-dependent at 0.1-0.4μg/ml of doxorubicin. It was always higher for encapsulated doxorubicin. The nanoparticles containing 2μg of DNA displayed slightly higher cell inhibition rate than those containing 1μg of DNA. At 0.2μg/ml of doxorubicin most HeLa cells became rounded or detached from the plate, and blebbing occurred. The nanoparticles without alginate led to lower cell inhibition rates. If the nanoparticles contained another plasmid (pGL3-Luc) and did not contain doxorubicin, no cytotoxicity was found [65]. Layer-by-layer absorption of alginic acid and protamine in water or 0.5M NaCl (pH 5.5) onto spherical calcium carbonate microparticles produced in the presence of alginic acid and having size from 2 to 12μm with subsequent dissolution of the CaCO₃ core in HCl or EDTA was also reported. The microcapsules synthesized in this way could be used for the encapsulation of polymers and their pH-dependent release, as demonstrated for dextrane conjugated with fluorescein isothiocyanate [66].

Although not stated in the respective publications, the great advantage of this method is the improved particle size control. For example, the techniques of synthesis of metal nanoparticles with predictable size already exist. Another advantage is the possibility of encapsulating the substances into some types of core particles instead of encapsulation into the alginate layer. In this case the alginate layer acts as an additional diffusional barrier. But the existence of a core particle limits versatility and the possibility of food or biomedical application of these particles. However, this is a convenient method of generating micro- or nanoparticles by chemical dissolution of some types of core particles.

**Emulsification-Internal Gelation**

Figure 3: Alginate particle prepared using emulsification method. 1 – encapsulated drug; 2 – alginate core; 3 – empty micelle.
In this method, an aqueous solution of sodium alginate containing an insoluble calcium salt (usually carbonate) is emulsified into an oil phase (Figure 3). Upon addition of an acid, calcium ions are liberated inside the alginate drops converting sodium alginate into calcium alginate, i.e. to a hydrogel, and the drops become being particles.

The technique of emulsification-internal gelation was useful for glucose oxidase encapsulation. A water-in-oil emulsion consisting of mineral oil, surfactant Span 80, acetic acid and sodium alginate solution – previously ultrasonicated with calcium carbonate – was prepared. Then water was added, the emulsion was stirred, and the formed beads were rinsed with Tween 80 and water. The resulting empty beads were incubated in buffered enzyme solution followed by chitosan coating. In the wet state the microparticles had volume-based mean diameter of 138μm. After lyophilization, storage at -20 °C and rehydration the immobilized enzyme lost 10% of its activity, and the uncoated beads collapsed. Loading and encapsulation efficiencies were 10% and 80%, respectively, and depended on enzyme loading pH. Loading efficiency increased with enzyme concentration, while encapsulation efficiency did not. Both parameters decreased with chitosan concentration, but the activity of the immobilized enzyme reached a maximum peak at 0.5% chitosan. It was concluded that these nanoparticles had the potential to serve as a useful and efficient delivery system for therapeutic proteins. Interestingly, at high enzyme concentrations the immobilized enzyme exceeded in activity the free one [67]. In a related report, no coating was applied, and canola oil was used as oil phase. Dry CaCO₃ powder was ground prior to application to decrease grain size from 30μm to 2.5μm. CaCO₃ substitution for oxalate or tartrate was unacceptable because too low pH was required. Otherwise, phosphate was rejected because of too large grain size, and citrate produced large beads with a high variability in size distribution. At CaCO₃/alginate monomer ratios higher than ¼, residual insoluble calcium salt was observed inside the particles. The alginate particles had mean diameters from 50 to 1000μm. The use of low guluronic and low or medium viscosity alginate gave a narrowest particle size distribution, but high guluronuc content and homopolymer blocks led to stronger gels. If Span 80 concentration increased up to 1%, the mean particle size decreased asymptotically. The particles were stronger and more spherical when using larger amounts of acetic acid while limiting the pH reduction by enhancing the buffer capacity of sodium alginate solution using carbonate buffer. Further reduction of polydispersity was achieved by dropping the oil phase into the acidic buffer solution in water. The synthesis can be carried out under sterile conditions, and its potential application may be cell immobilization [68]. A similar protocol did allow the encapsulation of mouse insulinoma 6 cells. For this, to the sodium alginate solution containing a suspension of CaCO₃ and cells in their culture medium mineral oil was added followed by the addition of acetic acid previously mixed with mineral oil. After that, a buffered saline solution mixed with culture medium was added to neutralize the pH. All the experiment was carried out under agitation, and its rate determined the particle size. It could be made from ~200 to 2000μm (circularity was 0.9), the distribution was close to normal but very wide. The beads were separated by centrifugation, washed and separated by filtration. CaCO₃ was not completely dissolved, and its grains remained inside the alginate beads. The beads had lower compressive stress resistance than those produced by extrusion using a commercial encapsulator. MOPS buffer provided better cell survival than HEPES buffer under optimized conditions (pH, buffer concentration, emulsification and acidification time). Cell division inside the beads was observed, and the encapsulated cells produced insulin at higher concentrations than single cells. The emulsion process is potentially suitable for cell therapy and transplantation purposes [69]. The use of paraffin oil has also been reported. In this case, a homogenized CaCO₃ suspension in a solution containing sodium alginate, chitosan and drug (salbutamol sulfate) was added into liquid paraffin oil containing Span 80. After emulsification, liquid paraffin containing glacial acetic acid was added, and stirring was continued until CaCO₃ was solubilized. Then CaCl₂ solution containing Tween 20 was added to recover the beads by decantation. Beads were washed with the same solution and then with acetate buffer. The parameters for the optimized formulation were as follows: spherical, size 591±40μm, encapsulation efficiency 95%. These values increased conforming to increasing polymer concentration. Cumulative drug release after 2 hours at pH 1.2 was 7%, after subsequent 3 hours at pH 6.8 (phosphate buffer) 10%, and after 5 hours at pH 7.4 (phosphate buffer) 99.6%. It was concluded that the hydrogel particles were the potential system for colon delivery of salbutamol sulphate via oral route for chronotherapy of nocturnal asthma [70].

However, in another publication [71] it was stated than salbutamol sulfate absorption site was not colon, moreover, it would degrade in colon, although no data was presented. Using the same technique as that in the previous report, alginate microparticles having mean size from 4 to 7μm (depending on recovery protocol) with immobilized insulin were synthesized. The particles were recovered from the oil phase by washing it with acetate buffer and separated by decantation. The remaining particles were recovered by centrifugation. The difference between the recovery protocols were centrifugation rate and additives to the buffer (CaCl₂, Tween 80, solvents). Only the protocol employing acetone, isopropanol and hexane returned the secondary structure of insulin. The encapsulation efficiency was more than 75% with the particle recovery yield around 70%. The particles were uniform and spherical with a rough surface. It should be noted that all the formulations showed burst release of all the encapsulated insulin for 5 min in simulated gastric solution without enzymes. But the technology was claimed as promising for insulin delivery due to the gentle methodology of producing microparticles [72]. In order to evaluate the effectiveness of insulin-loaded nanoparticles on its permeability across intestinal cells, another encapsulation technique was used. Here the alginate solution contained dextran sulfate, poloxamer 188 and ultrafine calcium carbonate suspension. Sorbitan monooleate was used for emulsification of this solution into paraffin oil under stirring and ultrasonication. Insulin solution in HCl was added simultaneously with acetic acid solution in oil under the same conditions. Nanoparticles were washed and coated with chitosan, polyethylene glycol and albumin. At the time of
coating they were treated with CaCl$_2$. Their average size was 300nm with polydispersity index 0.17, zeta potential 29mV, encapsulation efficiency 31% and loading efficiency 6%. The nanoparticles released up to 35% of insulin at pH 1.2 for 2 hours. At pH 5.5 burst release was observed. Dye-labeled nanoparticles applied at 0.5mg/ml were found at the cell membranes and probably inside Caco-2 and HT29-MTX cells co-cultured in a 90:10 proportion. Nevertheless, at 0.1-1mg/ml they had relatively low cytotoxicity for separately cultured AGS, Caco-2 and HT29-MTX cells. At 0.5 mg/ml the nanoparticles enhanced insulin permeation through a monolayer of Caco-2 and HT29-MTX cells co-cultured with Raji B cells (a model of intestinal wall) from the apical to basolateral direction at pH 5.5 and especially at pH 7.4. Insulin permeability was demonstrated to be mediated by clathrin endocytosis. The clinical potential of the nanoparticles for the oral delivery of insulin and therapy of type I diabetes mellitus was envisaged [73].

Internal gelation using canola oil containing acetic acid and surfactant Span 85, and sodium alginate solution containing CaCO$_3$ nanoparticles (116-244nm depending on the conditions, zeta potential 14mV) capped with surfactant AOT carried out in an electro dispersion reactor allowed size control of the microparticles by peak voltage increase in the range of 412 to 10µm. Further peak voltage increase did not decrease the bead size substantially, i.e. the method is currently limited to microparticles. Dried particles aged in CaCl$_2$ solution had a diameter of about 5µm. The flow rate ratio also significantly influenced the droplet size. The particles were highly porous because of CO$_2$ release during their formation and were spherical with a wrinkled surface. Only a slight amount of CaCO$_3$ was required to induce gelation, viz. 0.01M for 4% (w/v) alginate, although much higher concentrations were used in practice. The microparticles appeared suitable for encapsulation of pharmaceutical compounds [74].

A technique of internal gelation without employing emulsification has been reported. Mixing CaCl$_2$ with DM-nitrophen™ leads to a complex. This complex has a K of 5nM that dramatically increases after UV photolysis. Bulk hydrogels prepared in this way were intended for patterned microscale 3D cell culture, and their applicability for human cell encapsulation was demonstrated. But the high cost of DM-nitrophen™ limits its applicability [75].

In a previous review [76] the following advantages of emulsification-internal gelation method were stated:

- Particle size can be easily controlled from a few nanometers (potentially) to millimeters by controlling oil viscosity, sodium alginate type and concentration, surfactant concentration, and emulsification conditions, including mixing.
- The particles are homogeneous.
- Low shear protects fragile encapsulants.
- Scale-up potential is almost unlimited without the loss of particle size control.
- The following disadvantages were pointed out:
  - High porosity of the particles.
  - Possible loss or inactivation of encapsulated proteins during formulation and washing steps.

Additional concerns:
- If mineral oils (paraffin oil, isooctane, silicon oil, etc.) are used, they may cause environment-related problems.
- Acid concentration must be carefully calculated to avoid overdosing [76].

External Gelation

In this method a soluble calcium salt (usually chloride) induces alginate gelation. But simple mixing the solutions of sodium alginate and CaCl$_2$ yields bulk hydrogel without any particles. Therefore, special techniques are used.

Emulsification coupled with external gelation

Here sodium alginate solution containing a drug being encapsulated is emulsified in an oil phase. The emulsion droplets formed as act as templates for the formation of spherical particles. Then CaCl$_2$ solution is added converting the drops to particles [77].

Surfactants are usually used to stabilize the emulsions. However, sometimes they are not employed. The addition of sodium alginate solution to sunflower oil with subsequent homogenization by mixing, addition of CaCl$_2$ solution and acetic acid and another homogenization step led to beads. They were separated by centrifugation, washed with hexane and water followed by treatment with isopropanol: water mixtures with increasing concentration of isopropanol up to 100%. The last step was supercritical drying. The loading was carried out by bead incubation at 40 °C at 120 atm with ibuprofen for 24 hours. Bead characteristics were as follows: loading efficiency 30.5%, pore size 35nm, surface area 359.5m$^2$/g. High loading efficiency makes these aerogel beads efficient drug carriers [78].

In a simple technique employing a surfactant, CaCl$_2$ solution was added to that of sodium alginate containing 5-25% PEG-1500 or 0.5-5% Tween 80 at pH 5 under sonication. The nanoparticles were separated from the reaction mixture by centrifugation after 24 hours [66,79]. They had average diameter of 210-540nm but were prone to collapse during drying [66]. The amount of PEG-1500 and Tween 80 was about 64mg and 19mg, respectively, per one gram of lyophilized nanoparticles independently of the initial concentrations of these stabilizers in sodium alginate solution. In the presence of Tween 80 some hydrophobic regions were formed inside the nanoparticles. The nanoparticles containing polyethylene glycol on the surface are expected to have low immunogenicity, and those having Tween 80 are expected to cross blood-brain barrier [79].

Nanoparticles were also produced by dropwise addition of sodium alginate solution containing a plasmid encoding enhanced green fluorescent protein to the solution of AOT in toluene. Then CaCl$_2$ solution was added. After washing with acetone and water the nanoparticles were separated by centrifugation. They had a core consisting of sodium alginate, and a membrane consisted of calcium alginate. Their morphology depended on the position on
the respective point on the phase diagram, and that was in turn determined by the toluene: alginate: AOT ratio. Only in one region separated and spherical nanoparticles having size 55-100nm (average size 80nm) were formed. Transfection of cultured NIH 3T3 cells performed using the nanoparticles (at 2μg/ml of encapsulated DNA) and the complex of DNA with polyethyleneimine has shown comparable results after 48 hours, but the nanoparticles performed more slowly. The nanoparticles were proposed as non-viral vectors for gene therapy. Once internalized by non-phagocytic cells via endocytosis pathways, they could undergo quick erosion and elicit osmotic swelling, hence facilitating endosomal escape of gene to the cytoplasm [80].

Alginate microparticles can be used as a source of Ca²⁺ ions causing gelation of sodium alginate solution (in phosphate buffered saline) they were placed into [81]. The latter solution may contain peptides [81,82], oligonucleotides [81] or cells to be delivered by subcutaneous injection close to a tumor site. The solution had a volume of 150μl and contained ~10⁶ microparticles. Gelation in vitro occurred for several minutes. The gels remained intact in mice for more than 21 days and than dissolved [81]. The peritumoral gels carrying antigen-pulsed dendritic cells accumulated CD4⁺ T-cells but did not attract a significant amount of CD8⁺ T-cells, and the effect on established tumor growth was limited. If the gel also contained interleukin-15 super agonist, it could be released for over 2 weeks in vitro (in RPMI growth medium) and for 5-7 days in vivo. This gel retarded tumor growth and caused transient reduction of the size of larger tumors. This effect was mainly caused by the cytokine, with dendritic cells providing only a minor effect. The average suppression of tumor growth caused by the above-mentioned gel, the same gel without the cells but containing the cytokine and Toll-like receptor-9 agonist CpG and the gel containing only the cytokine but injected two times was equal. Local immunotherapies may be used for treatment of unresectable tumors and sites of surgical resection to combat local recurrence. Local immune response might make the tumors to become in situ vaccines themselves [82]. The microparticles with the average diameter of 20μm were synthesized as follows. Sodium alginate solution in phosphate buffered saline was added dropwise to the solution of Span 80 and Tween 80 in isooctane. After homogenization of the resulting solution CaCl₂ solution was added followed by another homogenization step, centrifugation and washing with isooctane and water [81].

A close technique can be used to form alginate nanocapsules. Nanocapsules are vesicular systems having oily or aqueous core surrounded by a polymeric membrane. The drug is mixed with an organic solvent, and the mixture is slowly added to aqueous solution of sodium alginate containing a surfactant, and an oil-in-water emulsion is formed under sonication. Finally, CaCl₂ solution is slowly added, and resulting calcium alginate forms a drop membrane. This method is reviewed in [77].

Sometimes liquid paraffin can be used as an oil phase. The solution containing guar gum, sodium alginate, Tween 80 and drug (salbutamol sulfate) was added dropwise to liquid paraffin followed by the addition of CaCl₂ solution and incubation. Another preparation was produced by dropwise addition of a solution containing chitosan, sodium alginate, the drug and acetic acid to liquid paraffin followed by addition of diocetyl sodium sulfosuccinate and of glutaraldehyde solution in toluene. All the experiments were carried out under stirring. The microparticles produced were filtered, washed with petroleum ether and dried. They were spherical with average size of about 438μm. Encapsulation efficiency was more than 95%. Their incubation in simulated gastric fluid for 2 hours and subsequently in simulated intestinal fluid for 10 hours caused gradual release of more than 85% of the encapsulated drug. Salbutamol sulfate is used in the management of bronchopulmonary diseases involving bronchospasm, and its sustained release dosage forms are needed to reduce frequent dosing (3-4 times a day) [83]. Interestingly, a close technique without the use of a surfactant was reported. Salbutamol sulfate and liquid paraffin were added to a solution containing sodium alginate, hydroxypropyl methylcellulose and starch. Homogenization and stirring led to a stable emulsion that was extruded through a needle into CaCl₂ solution and incubated in it followed by filtration, washing and drying. The beads were mixed with magnesium stearate and manually filled into empty hard gelatin capsules. The capsules were administered to fasted rabbits through plastic tubing followed by flushing of water. They remained in the stomach for more than 6 hours. Their floating time in simulated gastric fluid in vitro was also more than 6 hours. The capsules used for the experiment with rabbits also contained BaSO₄. Encapsulation efficiency and bead size increased with hydroxypropyl methylcellulose and starch amount increase. This might be due to emulsion viscosity increase. Bead density decreased with hydroxypropyl cellulose amount increase and starch amount decrease. Drug release after 10 hours of capsule floating in simulated gastric fluid decreased simultaneously. For the optimized formulation encapsulation efficiency and drug release were 70% and 57%, respectively, with the average bead diameter of 1.9mm and density of 0.85g/cm³. Bead surface was rough and had small pores. These stomach-specific floating systems might be beneficial over the conventional oral dosage forms (prone to tolerance development after repeated administration) because of prolonged gastroretentive delivery of drugs with improved bioavailability, therapeutic efficacy and patient compliance [84].

In order to ensure production of calcium alginate nanoparticles without damaging the biomolecules being encapsulated, reversed phase inversion temperature method has been proposed. In this case, the mixture of aqueous sodium alginate solution, decane and tetra ethylene glycol monododecyl ether employed as a surfactant was equilibrated at the temperature slightly below the phase inversion temperature of this system. Then the mixture was quickly transferred to a temperature well above it leading to formation of water-in-oil emulsion, and CaCl₂ addition to this emulsion led to the formation of nanoparticles. The most stable particles were produced if CaCl₂ was added as a simple solution, not as water-in-oil emulsion. All the synthesis was carried out under stirring. In order to separate the nanoparticles, decane was added leading to precipitation of micrometer-sized structures with an irregular shape. The nanoparticles remaining in solution were spherical and homogeneous with mean radius of 50nm and size distribution ~20-120nm in radius. The yield of nanoparticles was ~18%. The

amount of the surfactant and decane remaining in the sample was below 0.01 wt.% and 0.2 wt.%, respectively. The procedure has potential to be scaled up and to be adapted for use with pharmaceutically acceptable oils. The size of the resulting particles makes them suitable for delivery of biomacromolecules to cells and certain mucosal tissues [85].

In an approach developed at the same time by another group, both sodium alginate (containing bovine serum albumin as a model protein) and CaCl₂ aqueous solutions were emulsified in isopropyl myristate using diocetyl sodium sulfosuccinate as a surfactant. CaCl₂ microemulsion was poured dropwise into the alginate one followed by ultrasonication. The nanoparticles were separated by centrifugation and washed with acetone. The size of the particles was about 500nm, but particles smaller than 100nm were also present. Loading efficiency was 40%. The nanoparticles showed burst release of the encapsulated protein into phosphate buffered saline for 15 min, followed by a plateau and than by sustained release of all the remaining protein for more than 8 hours. The released protein did not suffer covalent aggregation or degradation by fragmentation. The nanoparticles were intended for oral protein delivery, and the components for their preparation were pharmaceutically acceptable [86].

A possibility to produce alginate particles without using water has been reported. This technique involves addition of mixture of drug (salbutamol sulfate), sodium alginate, CaCl₂, and PEG-400 into molten PEG-6000. All the reagents (except PEG-6000) were applied as powders. The resulting mixture was cooled rapidly at -20 °C and then powdered in a mortar followed by sieving. Particle size range for different formulations was 6.75-9.8μm. They had rough surface [87]. Interestingly, particle size increase with sodium alginate concentration increase reported for encapsulation of the same drug by dripping method [88] was observed here as well. Drug release in phosphate buffer decreased with sodium alginate concentration increase and increased with PEG-6000 proportion decrease. This was explained by hydrogen bonds between alginate and polyethylene glycol. All the formulations tested showed burst release with subsequent release of the remaining drug for several hours. They were intended for pulmonary drug delivery by means of a dry powder inhaler [87].

The advantages of the method are the same as those stated above for emulsification-internal gelation, except particle homogeneity. The additional advantages are easier formulation because of no need for finely dispersed insoluble calcium salt, decreased porosity and the absence of acid. The disadvantages are the use of oil and usual need for surfactants with questionable biocompatibility. Therefore, these nanoparticles have rather low chances for food applications, but limited agricultural or even biomedical use is possible. It should be noted, however, that surfactants make the nanoparticles less prone to aggregation. Therefore, the nanoparticles produced by this method must have enhanced stability.

Microfluidic devices

The advances in microfluidics paved the way to its application in nanotechnology, the synthesis of calcium alginate nanoparticles not being an exception. The parts of microfluidic chips can be manufactured in the laboratory from plastic by means of commercially available computer-controlled laser cutters.

For the synthesis of alginate microparticles having diameters in the range 50–2000µm with a variation less than 5% a microfluidic chip has been invented. A stream of sodium alginate solution was focused by two sunflower seed oil streams leading to emulsion that was dripped into CaCl₂ solution, and microparticles formed and precipitated at the interface between the phases. The emulsion size increased when the aqueous flow increased. When the oil flow increased, the emulsion size decreased. Increasing both oil and alginate solution flow rates while maintaining the flow rate ratio resulted in smaller emulsions. Gold nanoparticles were encapsulated into the alginate microparticles at the time of their synthesis. The microparticles were freeze-dried and than vacuum-dried, but most of them retained spheroid shapes. The approach will provide many potential uses for pharmaceutical applications such as targeted drug delivery and DNA transfection [89]. In another microfluidic device liquid paraffin containing Span 80 was used as the continuous phase and sodium alginate solution containing Na₂SO₄ served as the dispersed phase. The droplets of the mixture were collected in BaCl₂ solution. This led to the formation of barium alginate microspheres with BaSO₄ nanoparticles inside. The microspheres were washed and stored in NaCl solution. They were spherical with the diameter 229-284 μm (average 250μm). Their size could be modulated by adjusting the flow rate ratio of the two phases. The distribution of BaSO₄ nanoparticles was denser on the edge of the microspheres than in their interior. The aggregation of the nanoparticles was not observed. BaSO₄ dispersion density increased with Na₂SO₄ concentration increase, but at 0.4M Na₂SO₄, the precipitate appeared on the microsphere surface making it brittle. The microspheres were suggested as an embolic agent synthesized by means of droplet microfluidics and intended for transcatheter arterial embolization. In this method an embolic agent is injected through a microcatheter guided by a radiographical instrument in order to cause artery occlusion and to prevent nutrient supply to the tumor. At 5mg/ml the microspheres were not cytotoxic to HepG2 cells. After injection of them at ~30mg/ml to the renal artery of healthy rabbits interlobular and arcuate arteries became almost completely filled with the microspheres possessing excellent X-ray visibility. These microspheres showed better and longer lasting X-ray visibility after embolization if compared with commercially available calcium alginate microparticles that did not contain BaSO₄. The described microspheres present exciting opportunities for potential clinical applications in transcatheter arterial embolization therapy and for noninvasive tracking and mapping of the fate of the embolic agents in vivo [90].

Needless to say about the great advantages of microfluidic devices. They simplify scaling up while minimizing batch-to-batch variations and labor requirements. Moreover, they can be part of more complex devices and enable synthesis of nanoparticles in the field. The possibility to synthesize nanoparticles in small quantities is promising for personalized medicine. However, not every specialist can invent them, and it would be impossible for many poorly equipped or non-technical (e.g. purely biomedical) laboratories to construct them.
Water-in-oil-in-water emulsions

In more complex variant of the technique, the water-in-oil emulsion is further emulsified in water containing another surfactant, and then CaCl₂ solution is added. Calcium alginate nanoparticles (500-700nm, zeta potential around -13mV) stabilized by AOT and polyvinyl alcohol were synthesized by gradual addition of calcium chloride solution to water-in-oil-in-water emulsion of sodium alginate containing doxorubicin or rhodamine being encapsulated. Loading efficiency for these drugs was 5% and 4%, respectively. The nanoparticles enhanced doxorubicin cytotoxicity to cultured cancer cells in a dose-dependent manner (at 0.01-0.1µg/ml for MCF-7 cells and at 0.1-10µg/ml for NCi-ADR/RES cells) by extending its way to nucleus overcoming P-glycoprotein-mediated efflux without the use of additional P-glycoprotein inhibitors. At highest concentration of the encapsulated doxorubicin no viable cells were observed. Overexpression of P-glycoprotein results in the development of multidrug resistance of cancer cells, but drug encapsulation in nanoparticles overcomes this effect. Increased therapeutic efficiency was associated with an increase in the level of cellular and nuclear drug accumulation [91]. Similarly produced nanoparticles enhanced activity of encapsulated methylene blue in photodynamic therapy of cultured cancer cells (at ~0.5-20µM in a concentration-dependent manner for MCF-7 cells) by promoting its nuclear delivery and increasing production of reactive oxygen species. Decreased viability of 4T1 cells was observed only at the highest drug and light doses (at 0.6µM and 2400µJ/cm² but not at 0.3µM or at 1200µJ/cm²). The parameters of the nanoparticles were as follows: average diameter 79nm, surface charge 19mV, encapsulation efficiency 82%. Methylene blue can be used as a photosensitizer in photodynamic therapy of cancer, but its clinical use has been limited because of insufficient therapeutic efficacy. The respective study was aimed at enhancement of its efficiency by means of encapsulation in nanoparticles [92]. Lyophilized nanoparticles produced using the same technique and by the same group increased the efficiency of photodynamic therapy based on 8µM free or encapsulated methylene blue and directed against 4 cultured cancer cell lines under ambient conditions. They also reduced mammosphere formation by cancer stem cells present in the culture of MCF-7 cells under normoxia and hypoxia. It was achieved by switching the mechanism of radical generation to a type I reaction requiring much less molecular oxygen. This is significant because of a usually hypoxic tumor microenvironment. The loss of efficacy at low concentration of oxygen is a major limitation of photodynamic therapy, and it can be overcome by promoting type I chemistry by methylene blue encapsulation in nanoparticles composed of polymer and surfactant that are both anionic. The nanoparticles were spherical with a hydrodynamic diameter of 275nm, zeta potential of ~20mV and loading efficiency of 7%. Aggregates were separated by centrifugation [93]. The AOT-alginate nanoparticle system is a promising delivery vehicle for anticancer photodynamic therapy with methylene blue [92,93]. The diameter of the freeze-dried nanoparticles with simultaneously encapsulated doxorubicin and methylene blue was 40nm if measured by means of atomic force microscopy and ~75nm if measured using dynamic light scattering. Aggregates might form, but they were separated by centrifugation. Drug loading was 7% and 13%, respectively. Their administration via a lateral tail vein to mice with drug-resistant mammary adenocarcinoma at 4 and 8mg/kg, respectively, followed by light exposure slowed down the tumor growth and increased mice survival. 15% of tumor cells were apoptotic compared with 1% if nonencapsulated drugs were administered. The nanoparticles also damaged tumor neovascularization. Nanoparticle-mediated combination of chemotherapy and photodynamic therapy has significant therapeutic potential against drug-resistant tumors, for it could overcome drug resistance by invoking multiple anticancer mechanisms. Encapsulation of the drugs in alginate nanoparticles may result in significantly enhanced intracellular accumulation and production of reactive oxygen species [94]. Encapsulation efficiency of methylene blue could be increased up to 99.8% by sodium alginate and AOT concentration increase as well as by decrease of the volume of the AOT phase. But it decreased with the amount of drug added to the formulation. Many other ‘model drugs’ could also be encapsulated, for example verapamil, clonidine, diflucan and fluorescein. The encapsulation efficiency and release profile strongly depended on drug composition. The release of weakly acidic diclofenac in phosphate buffered saline was faster than that of acidic doxorubicin and verapamil, but even the release of 70% of encapsulated diclofenac lasted for 7 days with initial burst release. The encapsulation efficiencies for these drugs were 6%, 60% and 77%, respectively. Two drugs could be encapsulated simultaneously, but the release became faster. The zeta potential of the nanoparticles was in the range of ~25 to ~35mV. Surfactant-polymer nanoparticles would be useful for sustained delivery of water-soluble drugs, maybe together with imaging agents [95]. Similar technique was used for encapsulation of pesticide imidacloprid in order to investigate the ultimate effect of this pesticide in the form of nanoformulation. The nanoparticles had the following characteristics: size ~150nm (aggregates were present), encapsulation efficiency 99%, loading efficiency 2.5%. Increasing the concentration of AOT was found to increase particle size. Increasing the concentrations of both sodium alginate and AOT led to encapsulation efficiency increase, but the effect was complex. The encapsulated pesticide was less cytotoxic against cultured Vero cells at 5 and 10µg/ml if compared to free imidacloprid. But it was more effective against leafhoppers after 11-15 days of spraying 150ml of its 0.145-mg/l solution on ~10 Okra (bhindi) plants growing on a total area of 1m². Nano formulations of pesticides will reduce the amount of pesticide needed, decrease the environmental pollution, side effects and the amount of pesticide residues in food. They will be beneficial for economy, be of increased safety for the farmer or pesticide applicator and offer targeted delivery [96].

As can be seen, the double emulsion technique is much less popular than those involving water-in-oil emulsions. They share the same disadvantages, but the former one is more labor-consuming and requires two surfactants. However, the technique may provide somewhat better particle size control, because the maximum size of the aggregates is limited by the size of the micelle.
Dripping method

The simplest way to produce particles using external gelation is dripping one solution into another one (Figure 4). When a drop of sodium alginate solution faces Ca\(^{2+}\) ions, gelation immediately occurs onto its surface preventing its dissolution. Then Ca\(^{2+}\) ions slowly diffuse inside the drop to induce gelation in all its volume, and the drop becomes a particle.

Figure 4: Dripping method. 1 – sodium alginate solution; 2 – CaCl\(_2\) solution (or vice versa; a drug can be in either solution); 3 – alginate particle; 4 – stirrer.

Dripping CaCl\(_2\) into sodium alginate

If CaCl\(_2\) solution is being dripped into sodium alginate solution, so-called pre-gel is initially formed. This pre-gel is a colloidal solution (i.e. nano- or microparticles) of calcium alginate hydrogel that has not separated into a new bulk phase yet. This separation (i.e. aggregation) can be prevented by addition of chitosan or other stabilizers. However, the optimization of reaction time (before separation of the particles) and of concentrations of CaCl\(_2\) and sodium alginate makes possible the synthesis of nanoparticles even without stabilizers.

The nanoparticles prepared by dropwise addition of calcium chloride solution to the sodium alginate solution under continuous ultrasonication had hydrodynamic diameter from 210 to 540nm. In the presence of stabilizers (Tween 80 or PEG-1500) smaller nanoparticles having improved stability were produced. The size distribution and stability of the nanoparticles were also affected by ageing in the supernatant. Using these nanoparticles, bovine serum albumin encapsulation efficiency of more than 99% was demonstrated. The protein was admixed to alginate solution before the preparation of nanoparticles. The nanoparticles with encapsulated protein were claimed as perspective vehicles for hydrophobic drugs because they were composed only of components listed in the FDA Inactive Ingredients Database [97]. Another report in which a close technique was used dealt with insulin admixed to CaCl\(_2\) solution, and the nanoparticles were covered with chitosan. The main objective of the investigation was to develop insulin-loaded chitosan-coated alginate nanoparticles with small size in order to improve insulin bioavailability after oral administration. The parameters of the nanoparticles were as follows: average size 104nm, encapsulation efficiency 78%, zeta potential 4mV. These values increased with the amount of chitosan. 27% of insulin was released in simulated gastric fluid for 2 hours with initial burst release. When transferred in simulated intestinal fluid, the nanoparticles showed sustained and prolonged insulin release. Administered orally through a feeding needle at 50 or 100IU/kg body weight, the nanoparticles demonstrated significant reduction in glycaemia in diabetic mice for at least 9 hours with ~8% insulin bioavailability. The nanoparticles were firmly attached to the mouse intestinal lumen, as shown ex vivo using freshly excised small intestinal tissue. The acute toxicity study revealed safety of the nanoparticles administered orally at 150mg/kg. No mortality, no hepatotoxicity (tested as serum levels of alanine aminotransferase, aspartate aminotransferase, glutamate oxaloacetate transaminase and lactate dehydrogenase) and no nephrotoxicity (tested as serum and urine creatinin as well as urine protein, urobilinogen, blood, ketone, bilirubin and glucose levels, pH and specific gravity) were observed. Pathohistological study also showed no damage to liver and kidney tissues. Hence, the nanoparticles might be employed as a promising carrier for oral delivery of therapeutic proteins [98]. Sometimes this encapsulation technique can be used even without ultrasonication. An optimization study in which sodium alginate was tested in the concentration range 0.07-0.75% (w/v) and CaCl\(_2\) in the range of 0.025-0.15% (w/v) revealed their concentrations of 0.3% and 0.1%, respectively, as optimal, i.e. leading to spherical nanoparticles without aggregates or bulk gel. The nanoparticles were incubated in their original solution under stirring followed by centrifugation and freeze-drying. In incubation time and the stirring speed were also optimized. The nanoparticles coated with poly-L-lysine were used for bovine serum albumin immobilization. The protein was admixed to sodium alginate solution. Encapsulation and loading efficiencies increased with the protein concentration in sodium alginate solution up to 2mg/ml and reached 76% and 73%, respectively. The characteristics of protein-loaded nanoparticles were as follows: average size 50nm, polydisperisty 0.36 and zeta potential ~24mV. Protein release in phosphate buffered saline accelerated with time until 3 days, and finally ~60% of it was released. The nanoparticles were suggested as a carrier for drug and vaccine delivery, and their properties strongly depended on concentrations of both sodium alginate and CaCl\(_2\) [99]. A close technique was used to produce nanoparticles loaded with zinc-insulin complex. The nanoparticles were covered with chitosan that was added dropwise to the resulting solution. They were spherical but had rough surface. Their size increased with decrease of alginate: chitosan mass ratio from 764 to 2209±496nm. pH decrease from 5.2 to 4.7 slightly decreased the mean particle size, but further decrease to 4.2 displayed the opposite effect. The formation of new chemical
entities inside the nanoparticles was revealed, i.e. the nanoparticles with encapsulated insulin consisted of the complex and not of the physical mixture of the components [100]. In another case, insulin modified with methyl polyethylene glycol was added to sodium alginate solution, and then CaCl₂ and chitosan were applied. The nanoparticles were incubated in the reaction mixture under stirring followed by centrifugation. At pH 1.2 the release of encapsulated insulin began after 20 minutes, at pH 5.6-7.2 65-90% of insulin were released for 5 minutes. Therefore, the encapsulated insulin would mainly release in small intestines with minimal loss in the stomach [101]. The characteristics of chitosan-coated nanoparticles with encapsulated pesticide acetamiprid prepared in a similar way but with pH adjustment of the solutions were as follows: size 30-40nm, hydrodynamic diameter 201.5nm, polydispersity index 0.39, zeta potential ~32mV, encapsulation efficiency 62%, maximum release 93%. The nanoparticles were found to be superior to commercial formulation in terms of controlled release of the insecticide in soil and protecting it from degradation. The pesticide release rate increased with pH up to pH 10. This formulation could be used in any type of soil. It could reduce the frequency of application of pesticides by controlling the release and subsequently reduce harm caused to health by handling of pesticides or through residues in food or environment. The cost of pesticide application would also be diminished because less pesticide would be needed, and alginate and chitosan are inexpensive [102]. Chitosan-coated alginate nanoparticles with encapsulated herbicide parathion dichloride were prepared similarly. Interestingly, the particles were not removed from CaCl₂ solution after preparation. Their characteristics were as follows: spherical, size 197-305 nm, hydrodynamic size 635nm, polydispersity index 0.52, zeta potential ~23mV, encapsulation efficiency 74%. The encapsulation decreased parathion soil sorption and was therefore expected to greatly improve its herbicidal action. The formulation was shown to be promising for agricultural use, offering increased duration of herbicidal action, while reducing problems of environmental toxicity. The biodegradability of the nanoparticles is an additional advantage [103]. Likewise, prolonged release of herbicides imazapic and imazapyr from alginate nanoparticles covered with chitosan was shown. The aim of the work was to prepare and characterize nanoparticles containing these two pesticides simultaneously for use in agricultural weed control. The nanoparticles (average size 378 or 247nm (aggregates were present), polydispersity index ~0.3, zeta potential -30mV, encapsulation efficiency 62% for imazapic and 71% for imazapyr) were produced using the same technique as in the above cases. The particles were not separated from their original solution, and remained stable in it for at least 30 days. At concentration equivalent to 400 g/ha the encapsulated herbicides retained their activity against Bidens pilosa in soil, if applied pre-emergence. The parameters analyzed were germination efficiency, height, root length and green mass. The nanoparticles showed reduced genotoxicity to Chinese hamster cells treated with 0.1mg/ml of the herbicides for 1 hour, as shown by comet tests, and to Allium cepa roots treated with 0.5mg/ml of nanoparticles for 24 hours, as shown by relative alteration index determined using optical microscopy. At 75g/l of imazapic and 25g/l of imazapyr they also displayed reduced negative effect against soil bacteria 7 days after treatment, as revealed by real-time polymerase chain reaction (qPCR). After 30 days the result with the nanoparticles showed the greatest similarity to the negative control (untreated soil). The formulation offers an effective alternative for weed control in agricultural cultivations. The encapsulation of two herbicides in one carrier system to improve their activity and to reduce the impact on humans and environment was proven to be a good strategy for sustainable agricultural production, although scaling up the technology needs more research [104]. A close technique was used for encapsulation of antitubercular drugs separately [105] or in several combinations [106]. The study was aimed at evaluation of the chemotherapeutic potential of econazole encapsulated in alginate nanoparticles alone or in combination with other drugs against murine tuberculosis [106]. Chitosan-coated nanoparticles were recovered by centrifugation after overnight incubation in the original solution [105-107]. They had an average size of ~230 nm and a polydispersity index of 0.44. Encapsulation efficiency was more than 70%. Interestingly, the loading capacity was claimed as 480-730mg of drug per 100mg of alginate [105-107]. Oral treatment of mice infected with Mycobacterium tuberculosis H37Rv with either free or encapsulated drugs resulted in total clearance of bacilli in lungs and spleen compared with untreated control, but the required dosing frequency was reduced by 15-fold [105,106]. It was also valid if the nanoparticles were administered by nebulization to guinea pigs infected by the same bacilli. The aerodynamic characteristics of the droplets were as follows: 81% of them were in the size range 0.4-2.1µm; mass median aerodynamic diameter 1.1µm; geometric standard deviation 1.7µm. The nanoparticles did not display hepatotoxicity, as shown by the levels of serum bilirubin, alanine aminotransferase and alkaline phosphatase [107]. The doses were as follows: isoniazid 10mg/kg, rifampicin 12mg/kg [105-107], pyrazinamide 25mg/kg (orally) [105,106] or 15mg/kg (oral if free and nebulized if encapsulated) [107], ethambutol 16 mg/kg [105,106], econazole 3.3mg/kg body weight [106]. For a mouse weighting 20g, 143mg of dried nanoparticles was a therapeutic dose. Free drugs were administered in 50µl 1:4 (v/v) methanol: distilled water [105]. For a 400-g guinea pig the dose was 21.3mg of dried nanoparticles. The drugs for nebulization were suspended in 4ml of physiological saline [107]. Free drugs remained at detectable level in tissues for 12-24 hours after a single administration. In contrast, encapsulated drugs remained at least for 10 days. Alginate nanoparticles can serve as an ideal carrier for the controlled release of antitubercular drugs [105-107], moreover, they are inhalable [107]. It should be noted, however, that other authors did not recommend chitosan-coated nanoparticles produced using the same technique for pulmonary drug delivery by nebulization because of their too big size (556nm, polydispersity 0.29) and instability [108], although this technique has been reported to be able to generate much smaller nanoparticles, as discussed above. The nanoparticles were unstable in 10% fetal calf serum, simulated lung fluid and even in 0.9% saline, despite zeta potential of 29mV. Human alveolar type 1 epithelial cells at 1600cells/cm² in cubated for 24 hours with medium containing these nanoparticles at 2000µg/ml showed ~80% viability, and their DNA content

decreased to 76%. At 100-1000µg/ml significant dose-dependent cellular uptake of nanoparticles with encapsulated indocyanine green after 2 hours of incubation was shown. The nanoparticles released 27% of encapsulated bovine serum albumin in water through a dialysis bag within 2 days and more than 80% for 21 days. The goal of the study was to determine the most promising formulations for further development as carriers for pulmonary delivery of biological agents [108]. In the same way, alginate nanoparticles coated with quaternized chitosan were prepared. The reaction mixture was equilibrated overnight, and then the nanoparticles were separated by ultracentrifugation. Bovine serum albumin encapsulation efficiency increased with protein concentration up to 1mg/ml. Further increase promoted aggregation. The nanoparticles were spherical with a fluffy surface. Their zeta potential was 10 mV. Their size increased with the concentration of quaternized chitosan from 280.5 to 1152nm, and further increase led to aggregation. This concentration increase also promoted the encapsulation efficiency up to more than 40%, while decreasing the loading efficiency from more than 15% to ~10%. Particle size, encapsulation efficiency and protein release rate in HCl (up to 22% for 100 hours) also increased with chitosan molecular weight. Increasing degree of substitution of chitosan did not influence the particle size, but increased the encapsulation efficiency and slowed down protein release, although burst release was still observed. Therefore, the release of the encapsulated protein can be tailored by choosing the molecular weight and degree of substitution of quaternized chitosan. The nanoparticles were intended for oral delivery of proteins [109]. A close technique was used for salbutamol sulfate encapsulation. The pH of the solutions of chitosan (despite initial preparation in 1% CH₃COOH) and sodium alginate was adjusted to 5. The characteristics of the particles were as follows: size 288nm, zeta potential 23mV, encapsulation efficiency 21%, stomach retention time 11 hours (in vitro). Less than 75% of the encapsulated drug was released to fasted state simulating gastric fluid for 12 hours, but fast release at the initial phase was observed. The size of the particles prepared without pH adjustment or when chitosan was added before CaCl₂ was more than 800nm. In the terms of drug release profile or size this formulation excelled 24 others containing other polymers. The nanoparticles could be used for drug absorption enhancement [71]. For encapsulation of plasmid DNA encoding green fluorescent protein a different technique was used. The aim of the study was the development of several chitosan-containing nanoparticulate carriers of DNA and their comparison. In this case, the solutions of CaCl₂-DNA and chitosan were dripped one after another from a syringe to that of sodium alginate containing dextran sulfate. After incubation in the reaction mixture under stirring the nanoparticles were separated by centrifugation. Their characteristics were as follows: average size 196nm, polydispersity index 0.15, zeta potential 27mV, encapsulation efficiency 95%. The nanoparticles excelled those without dextran sulfate and those of pure chitosan in transfection efficiency of HEK 293 cells at 1µg of DNA for 48 hours. Unlike Lipofectamine, all the preparations of nanoparticles were not cytotoxic for these cells but had less transfection efficiency. The nanoparticles could be used as gene carriers in vitro and in vivo due to their biocompatibility [110].

As can be seen, dripping method is very popular because of its convenience. It does not require special skills and equipment, for pipette (or syringe) and stirrer (or ultrasonic bath) are enough. If the cells to be encapsulated are admixed to CaCl₂ solution, they never come into contact with sodium alginate that may be toxic to animal cells [111]. Moreover, the modern dripping techniques allow to generate nanoparticles.

**Dripping sodium alginate into CaCl₂**

Dropwise addition of sodium alginate solution to CaCl₂ solution is also widely employed. In this case the drug being encapsulated is always admixed to sodium alginate solution, because this drop becomes a particle. Drop size and shape determine those of the particle.

Dropwise addition of sodium alginate solution containing cyclophosphamide to CaCl₂ solution yielded beads having average diameter of 1mm. Bead drying retarded the release of 90-95% of the encapsulated drug into physiological saline from 80-100 minutes to 140-160 minutes. The beads could be suitable as a component of anticancer pharmaceutics because of their ability to be internalized by tumor cells [112]. Similarly produced microparticles with encapsulated crude turnip extract (applied in phosphate buffer) contained 0.295mg of protein per 1ml of matrix. Their average diameter was 390µm, and the encapsulation efficiency was 70%. They were used for treatment of wastewater containing phenol and chlorophenols, but the reaction products remained mainly in solution. The immobilization did not cause significant change of the enzyme catalytic constants. Polyethylene glycol protected the enzyme from inactivation, although some enzyme release to the solution still occurred. After each cycle the beads were washed to remove brown particles appearing inside them, and treated with CaCl₂. This step was essential for maintaining enzyme activity. Optimization of cycle time allowed to extend bead reusability to 15 cycles. Encapsulated peroxidase displayed higher efficiency of phenol removal but lower thermal stability than covalently immobilized enzyme. Development of a large-scale economically feasible process for industrial effluent treatment was expected [113]. In the case of beads produced using the same technique but in the presence of 0.9% NaCl strong swelling for the lowest molecular weight alginate tested was shown. Strong weight loss during bead preparation for the highest molecular weight alginate [2] and for those having low guluronic acid content and taken at low concentrations [2,114] was observed. Without NaCl the weight loss in all cases was less than 4% [114]. For the medium molecular weight alginate small weight loss was reported. The aim of the respective study [2] was the preparation of hydrogels with the applicability for immobilization of biological species. Indeed, cells can also be encapsulated using this technique. The pellet of yeast Candida tropicalis SSK01 cells was aseptically mixed into sodium alginate solution in Tris-acetate buffer (pH 7.5). The resulting solution was dripped into CaCl₂ solution using a micropipette. After incubation under stirring the beads were washed with the same buffer and stored in it at 4°C. The applicability of the immobilized cells for phenol degradation was demonstrated. The most significant operating factors and optimum levels of phenol degradation were found by means of statistical optimization.
approach. Functional relationship between independent variables like initial phenol concentration, temperature, pH and maximum percentage of phenol degradation was found out [115]. Almost the same technique was used for coencapsulation of human bone marrow mesenchymal stem cells at 2×10^7 cells per 1 ml of sodium alginate solution and an antibody to bone marrow protein 2 at 25µg/ml. Sodium alginate was coupled with RGD sequence (cell attachment peptide) and partially oxidized to increase its degradability. An in vitro experiment showed that the beads (average diameter 1 mm) released the encapsulated antibody in phosphate buffered saline for 10 days but still retained more than 30% of it after 14 days. The images acquired by means of bright field microscopy revealed uniform cell distribution inside the beads. The beads were transplanted into critical size calvarial defects in immunocompromised mice. The antibodies captured endogenous bone morphogenetic protein 2 and presented it to the stem cells, promoting their osteogenic differentiation and therefore bone repair and regeneration. The bone had a lamellar pattern, contained viable osteocytes, and was rimmed with osteoblasts participating in its formation. Dystrophic calcifications were not observed. Osteocytes and osteoblasts were observed even if the beads contained only the antibody or only the stem cells. But if the antibody was nonspecific and there were no encapsulated stem cells (i.e. negative control), fibrous and vascular connective tissue components consisting of interwoven bundles of collagenous fibers and calcium alginate were observed instead of regenerated bone. The bone regeneration strategy is a promising modality for repair and regeneration of craniofacial, axial and appendicular bone defects [116]. The co-immobilization of microalgae Chlorella vulgaris with auxin-producing bacterium Azospirillum brasilense or with chemically synthesized indole-3-acetic acid in calcium alginate beads by the same technique promoted microalgae multiplication and chlorophyll a, chlorophyll b, β-carotene, lutein and violoxanthin production. During solidification of the alginate into beads, numerous internal cavities of various sizes are formed randomly, and they provide the microenvironment for cells. Increasing biomass production of immobilized microalgae has important implications for environmental cleanup and some industrial processes [117]. The promoting effect of alginate oligosaccharides on root formation and growth in rice was also mediated by endogenous indole-3-acetic acid [30]. Using the same technique coupled with chitosan coating and fluid bed drying, beads having irregular structure and size of about 1 mm with encapsulated Bifidobacterium breve were prepared. After initial binding to alginate, the penetration of chitosan into the gel was seen only after 1080 min without any changes before this time point. Swelling and disintegration of the beads were pH-dependent. They released encapsulated bacteria into simulated intestinal solution (by bead disintegration) for 120 min after incubation in simulated gastric solution for 1 hour. In contrast to this, the beads without chitosan coating began to release bacteria at the stage of incubation in gastric solution, therefore they released much lower amount of viable cells to the simulated intestinal solution. The beads were produced in a scalable matter as a uniform granular product that could be used as a dry enteric delivery formulation for probiotic bacteria [118]. A close technique was used for the immobilization of bacteria. The main difference was the use of CaCl_2 solution in milk. After 8 -10 hours of milk fermentation the concentration of the released bacteria was less than 10^8 cfu/ml. (The ratio of beads to milk was adjusted to ~17%). Lactobacillus helveticus and Streptococcus thermophilus grew in the beads, while Lactobacillus delbrueckii ssp. bulgaricus did not [43]. Rifampicin and fluconazole were separately encapsulated into chitosan-coated alginate beads with a diameter of 1 mm in a similar way. Air drying retarded drug release into physiological saline. Further drug release rate decrease by 30% for up to 12 hours was observed with the increase of chitosan shell thickness from 25µm to 125µm that was, in turn, determined by the employed chitosan concentration. The use of these beads as antibacterial medicaments with prolonged action was suggested [119]. A slightly different technique was employed for the encapsulation of sunflower oil extract from tunic of ascidia. Sonicated emulsion of this extract in aqueous sodium alginate solution containing Tween 80 and activated charcoal powder at 50 °C was dripped into CaCl_2 solution through a syringe nozzle. The resulting beads were incubated in this solution followed by separation, incomplete drying, and transfer into chitosan solution. The characteristics of the beads prepared using the nozzles having diameter of 0.2 and 0.7mm were as follows: irregular shape, smooth surface, size 1.4 and 2.2mm, bead mass 2.5 and 3.1mg, carotenoid content 0.9 and 0.12µg/bead, respectively. They were stable in pure water, ethanol, and in HCl at pH 2.5 for at least 240 hours. Storage stability of dry beads for 5 months was shown. No visible degradation in water at 20-100 °C for 20 min was observed. The degradation rate increased with pH; however, the pH increase from 9.2 to 11 had no effect. The beads were suggested as a component of functional, i.e. healthy food products [120]. In another report multilayer coating was employed. Sodium alginate containing bovine serum albumin was extruded into stirred CaCl_2 solution through a commercially available encapsulator nozzle. The particles were than covered with poly-L-lysine, pectin, again with poly-L-lysine and finally with a new sodium alginate layer. Their diameter was ~400µm and depended on the nozzle used. The microcapsules were stable during in cubation in simulated gastric fluid for 3 hours followed by incubation in simulated intestinal fluid for up to 24 hours. They were stable for 24 hours at pH 1-9 as well, and had good mechanical stability. All the tests above were carried out using empty microcapsules. The loaded ones released half of the protein amount compared with the microcapsules without pectin and second poly-L-lysine layers. The microcapsules are expected to allow safe and efficient oral delivery of live bacterial cells for various clinical applications [121]. The encapsulator without subsequent coating was also used in the case of αTCP C2 cells. These cells synthesize prohormone convertase 1/3-derived proglucagon products. Injection of 1.5-2ml of the microcapsules with a size of 500-700µm in a total volume less than 3 ml to the intraperitoneal cavity improved glucose handling in mouse models of type 2 diabetes, providing a model for evaluating long-term delivery of proglucagon-derived peptides via cell therapy. Each mouse received about 4×10^7 encapsulated cells. In the case of mice with genetic defects in the leptin receptor signaling pathway the animals had significantly higher body temperature during acute housing at 4°C. If left untreated, these mice have defective cold-induced
termogogenesis, and their glucose-stimulated insulin secretion was also defective. The treatment did not change epidermal white adipose tissue and intrascapular brown adipose tissue of these defective mice [122]. Alginate microparticles coated with oligochitosan and containing encapsulated MCF-7 breast cancer cells were converted to alginate-chitosan microcapsules by treatment with EDTA dissolving the calcium alginate core. Sodium alginate solution containing the cells was dripped into CaCl$_2$ solution by means of a device with coaxial air flow or using an electrostatic generator of microparticles. The possibility of producing microcapsules with predetermined size by the means of the latter method was underlined but not demonstrated. Physiological saline was used for washing between the incubation steps. The microcapsules produced by two methods had average size of 750µm and 300µm, respectively, and were prone to swelling. Chitosan membrane thickness was about 82-89µm. Tumor spheroids were generated inside the microcapsules. The cells in the spheroids can better mimic cell behavior in a small size solid tumor than the cells in a monolayer culture. This new in vitro model of anticancer drug screening has advantages of spheroids produced using classic techniques [123]. In another report, sodium alginate solution containing drug (salbutamol sulfate), guar gum and/or pectin was dropped through a syringe into CaCl$_2$ solution and incubated under stirring. The beads formed were separated by filtration, washed and dried. Their mean diameter was 0.7-2.1mm depending on concentration of polymers. Encapsulation efficiency was 65-91%. The encapsulated drug was completely released during 3-7 hours of incubation in simulated intestinal fluid following 2 hours in simulated gastric fluid, where less than 20% was released. Sodium alginate concentration increase made the drug release more sustained. Guar gum and pectin improved it further, and the best formulation contained both polymers. The beads are promising pharmaceutical dosage forms, and better patient compliance can be expected [124]. In the same salbutamol sulfate encapsulation technique BaCl$_2$, Al$_2$(SO$_4$)$_3$ and magnesium aluminum silicate could be used instead of CaCl$_2$. Encapsulation efficiency was more than 77%. The size of oven dried nanoparticles produced using 2% sodium alginate and the respective salts was 921, 863, 899 and 782nm, and increased with sodium alginate concentration. 90% or more of the encapsulated drug was gradually released into phosphate buffer for 9 hours. Air-dried nanoparticles were larger in size due to incomplete dehydroxylation and showed slower drug release, but the encapsulation efficiency was comparable with that of oven dried ones. Particle characteristics did not change after 3 months of storage at 40 °C. Nanoparticles are expected to spread out in the gastrointestinal tract thus improving drug absorption and reducing local irritation [88]. Dried beads with encapsulated salbutamol sulfate (CaCl$_2$ was used) were dispersed in Eudragit S-100 solution in acetone for coating. The dispersion was poured into liquid paraffin, incubated under stirring to evaporate acetone, and finally the beads were washed with hexane and dried. They were spherical with a smooth surface, their size was 640-690µm with narrow size range, and encapsulation efficiency was 68-86%. It decreased with crosslinking time increase because of drug leakage and increased with sodium alginate and CaCl$_2$ concentration. As these concentrations and coating level increased, drug release rate decreased. Coating prevented drug release at pH 1.2 (HCl) and 6.8 (phosphate buffer). At pH 7.4 (phosphate buffer) more than 70% of the encapsulated drug was released for 21 hour. The preparation was intended for chronotherapeutic drug delivery for nocturnal asthma treatment and claimed as colon specific [125]. The same technique was used to produce microspheres with a diameter ~700µm containing encapsulated bovine serum albumin conjugated with tetramethyl rhodamine. They were incubated in CaCl$_2$ solution for 72 hours, then removed by filtration, washed and resuspended in 70% ethanol. After that they were shaken in silk fibroin solution, ultrasonicated to break down the aggregates and shaken again. They were washed with either water or methanol, dried and coated again. There were 3 fibroin layers in total. The shell was ~10µm thick. Loading efficiency was ~15% and was not changed by the coating. The coating retarded the conjugate release in phosphate buffered saline from 10 to more than 16 days. Incubation in EDTA solution led to alginate core dissolution, but the fibroin shell remained. Coating with fibroin has potential for biosensor and diverse array of controlled drug delivery applications [126].

For large-scale production of alginate beads having diameter of several millimeters a special device has been proposed. In this device, CaCl$_2$ solution is consistently fed to the center of the rotating disk, and above the disk there is a bowl fed with sodium alginate solution. The bowl is also rotating and spraying alginate onto the rotating disk where it contacts calcium chloride. The average bead diameter is determined by the disk diameter, its angular velocity, alginate solution viscosity and feed rate. It should be noted that alginate solution sterilization causes its depolymerization thus decreasing viscosity. From the ‘parent’ beads smaller ‘satellite’ beads are being detached making the bead size distribution bimodal. Improper selection of the parameters may result in producing undesirable alginate filaments instead of beads. The system could be applied to the immobilization of microorganisms, as well as plant or animal cell cultures. The device can be inserted in a fermentor for the immediate use of the beads. If the fermentation medium contains calcium, it can be used for bead production instead of CaCl$_2$ solution [127].

This variant of dripping method is also very popular and convenient. However, the method is generally limited to beads and microparticles because of difficulties with generating small alginate droplets due to its high viscosity.

**Calcium alginate nanoparticles as stabilizing agents for other nanoparticles**

The use of calcium alginate particles is not limited to encapsulation of various substances in molecular form. They can also be used as carriers for nanoparticles. The nanoparticles can be synthesized either in advance, as discussed above for the case of gold nanoparticles [89], or simultaneously with alginate particles, as described below. The synthesis of alginate particles using other ones as cores is not included here because it has been analyzed separately (see above).

Drop-wise addition of sodium alginate solution containing nifedipine to CaCl$_2$ solution in 1:1 alcohol/water mixture followed
by chitosan solution addition led to spherical nanoparticles having a size about 20-50nm and a fluffy surface with nifedipine nanocrystals inside. The average drug content was 6.7µg/mg, but less than 70% of the encapsulated nifedipine could be released. The particles could release nifedipine for more than 10 hours in a pH-sensitive way with an initial burst release for 2 hours. Although two of the three release media used were phosphate buffers, diffusion controlled release was revealed. This nanosystem offers an interesting potential for the delivery of hydrophobic compounds. If patients take it before sleep, drug concentration in plasma can reach peak value for initial burst, and then constant therapeutic concentration may be maintained by continuous and controlled release [128].

In another report dealing with nanoformulation of curcumin for delivery to cancer cells, ethanolic curcumin solution was added to CaCl₂ solution in water. Under these conditions some fraction of curcumin formed nanocrystals, and excess curcumin was removed by centrifugation. If CaCl₂ solution contained Pluronic F127, no precipitation of free curcumin was observed. The resulting solution was added dropwise to sodium alginate solution under stirring, followed by dropwise chitosan addition, and spherical particles with solid dense structure, uniform size distribution, and a mean size of 100nm were separated by centrifugation after incubation in the original solution. The pH of the final solution was made 4.7 by adjusting the values of alginate and chitosan solutions. Encapsulation efficiency in the presence of pluronic was more than 10%, and about 75% of encapsulated curcumin was released from lyophilized nanoparticles for 72 hours in phosphate buffered saline. Comparable internalization by HeLa cells was observed for free and encapsulated curcumin at 2 µM. The inhibition of cell proliferation of the DNA and its adducts is necessary for molecular biological assay to determine the level and nature of adduct formation and the extent of DNA damage [114,131].

However, in another report it was shown that the beads produced by internal gelation had lower mean sizes (2.2 mm at pH 4 and 3mm, respectively), higher shrinking (0.88% at pH 4 and 0.6%, respectively) and more homogeneous structure than those produced by external gelation. The shrinking increased with gelation pH decrease. The beads were produced by dripping alginate-CaCO₃ mixture in canola oil containing Span 80 into acetate buffer (for internal gelation) or into CaCl₂ solution (for external gelation). The internal pH of the bead was higher than that in the gelling solution. The empty beads were suspended in blue dextran solution and transferred to water. Blue dextran release for 15 days was 60% and 11%, respectively [68].

Aerosolization method

In aerosolization method, the solution of sodium alginate is converted to aerosol by a special device called nebulizer or atomizer (Figure 5). These terms are synonymous. In such devices pressurized carrier gas is needed for aerosol formation. Correspondingly, aerosolization can be called nebulization or atomization. The aerosol formed is carried by a flow of the carrier gas into CaCl₂ solution. The aerosol droplets become enclosed in gas bubbles inside CaCl₂ solution, therefore gas flow must not be too high in order to provide them enough time to diffuse to the solution. The method resembles dripping because aerosol also consists of
drops. However, here the only possibility is to aerosolize sodium alginate solution into CaCl$_2$ solution because in the reverse case bulk plain hydrogel is formed [2]. Another significant difference is nozzle placement. In dripping method, the nozzle is usually placed above CaCl$_2$ solution for drop formation. In aerosolization method the nozzle is usually placed inside CaCl$_2$ solution, because in the opposite case the yield of particles is very low. The carrier gas is usually air supplied by a pump. The process is accompanied by formation of secondary aerosol having other drop size distribution. Secondary aerosol leads to loss of solution and therefore to lower yields, but it can be prevented using a vigreux [63]. Aerosolization is usually considered as a method limited to production of alginate microparticles [77]. However, it has a potential of generating nanoparticles as well [63,132,133] because the aerosol drops can be made very small up to nanosize. It should be underlined that aerosolization is usually time-consuming, therefore, a fraction of nanoparticles stays in CaCl$_2$ solution for a prolonged time while other nanoparticles are being formed. If the concentration of CaCl$_2$ is too high, the nanoparticles will aggregate and even deposit [63]. It also limits the possibilities of scaling up the technique without changing the CaCl$_2$ solution.

The process of liquid atomization by a co-flowing gas stream is rather complex because of its multistage nature (Figure 6). At first, instabilities develop in the liquid layer, causing the formation, growth and detachment of liquid tongues. The initially formed large non-spherical parcels continuously break up into droplets. At a certain distance the break-up process finishes, and then the droplet size increases slowly due to coalescence [134]. The description above refers to the most common nebulizer type. There are, however, other types as well, those with hydraulic/airless spray nozzles being an example. The hydraulic nozzles utilize the hydraulic pressures to disintegrate the liquid jet into fine droplets [135].

The items 14 to 16 are used only with optional vacuum pump.

**Figure 5:** Aerosolization method. 1 – air pump; 2 – air filter (optional); 3 – air dryer (optional); 4 – sodium alginate solution containing drug; 5 – aerosol; 6 – nozzle; 7 – heater (optional); 8 – aerodynamic classifier (optional); 9 – magnetic stirrer (optional); 10 – CaCl$_2$ solution; 11 – air bubbles; 12 – capillary; 13 – vigreux (optional); 14 – liquid nitrogen; 15 – cold trap; 16 – exhaust tube.

The process of liquid atomization by a co-flowing gas stream is rather complex because of its multistage nature (Figure 6). At first, instabilities develop in the liquid layer, causing the formation, growth and detachment of liquid tongues. The initially formed large non-spherical parcels continuously break up into droplets. At a certain distance the break-up process finishes, and then the droplet size increases slowly due to coalescence [134]. The description above refers to the most common nebulizer type. There are, however, other types as well, those with hydraulic/airless spray nozzles being an example. The hydraulic nozzles utilize the hydraulic pressures to disintegrate the liquid jet into fine droplets [135].

**Figure 6:** Atomization regions of the spray from the hollow cone nozzle [136].
In the simplest variant of the aerosolization method, sodium alginate solution containing horseradish peroxidase and a small amount of sodium phosphate buffer was aerosolized above slowly stirred CaCl$_2$ solution using a manual disperser. After 2 hours of incubation the beads were separated by filtration and washed with phosphate buffer solution. The droplets produced by the disperser had rather big size, therefore the size of the beads was about 0.5mm. The beads contained 1.18µg of enzyme per 1mg of dry bead mass, and the encapsulation yield was 85%. The activity of immobilized horseradish peroxidase was significantly higher than that of the free enzyme. The immobilization also prevented enzyme activity loss at 60 °C for 1 hour, and the activity of the encapsulated enzyme was still measurable after storage of the beads for 7 months in phosphate buffer solution at 4 °C [137]. This was not the only example of enzyme activation after its encapsulation in calcium alginate. It was shown that at pH 5.5-6 or in alkaline medium the activity of horseradish peroxidase encapsulated in calcium alginate beads was higher than that of free enzyme. The reaction studied was phenol oxidation, and the beads were claimed as promising for continuous phenol removal from wastewater [41].

The encapsulation technique used was dropping CaCl$_2$ solution containing dextran and the enzyme into rapidly stirred sodium alginate solution followed by capsule transfer to a new CaCl$_2$ solution. The technique was also used for the encapsulation of monkey kidney cells, and their growing inside the capsules was observed. Capsule membrane thickness was found to be dependent on incubation time in sodium alginate solution, the capsule size and the concentrations of sodium alginate and CaCl$_2$. The possible fields of application of cell encapsulation techniques include bioreactors, artificial organs and others [111]. Enzyme activation caused by immobilization in alginate is not limited to external gelation techniques; it was also observed for glucose oxidase immobilized using emulsification-internal gelation technique [67], as discussed above.

For pre-industrial scale production of alginate microparticles an aerosolization device has been suggested. Compressed air after drying and filtration is directed to a piston pump fed with sodium alginate solution containing hydroxypropyl methyl cellulose and lysozyme being encapsulated. The solution is directed to airless spray-gun. The atomized droplets are collected into a tank filled with CaCl$_2$ solution (containing chitosan and acetic acid) and placed on a rotating platform. The particles are collected by filtration, washed and dried. Volume mean diameter of the produced particles depended on experimental conditions (sodium alginate concentration, nozzle size, swelling in water or phosphate buffered saline) and could be in the range from 69 to 274µm (without precise control). The particles larger than 400µm were also present. All the particles seemed to be aggregates. The particles released 40-90% of encapsulated lysozyme during the incubation in glycine buffer for 8 hours and then in phosphate buffered saline for 16 hours. The device appears to be suitable for production of microparticles showing gastro-resistance and suitability for oral applications or animal husbandry by processing feeding solutions of sodium alginate in a wide range of viscosities [138].

Aerosolization of sodium alginate solution containing drug diltiazem hydrochloride into CaCl$_2$ solution led to microparticles. In order to prevent their agglomeration, 20% (v/v) ethanol was added to CaCl$_2$ solution. The microparticles formed were dried and than rehydrated. Air flow rate increased their size and influenced their polydispersity. The buffers in the CaCl$_2$ and rehydration solutions (because of possible solubilization of alginate) and the presence of chitosan in the CaCl$_2$ solution were other factors influencing particle size. The particles having various sizes from 200 to 1100µm were produced. The drug release profiles were influenced by pH of the release medium, molecular weight of chitosan and amount of the drug loaded into the particles, but the equilibrium concentration was always reached after 30 min because of the high water solubility of diltiazem hydrochloride. Drug load, in turn, could be increased by addition of NaCl to sodium alginate solution because it decreased the electrostatic interaction between the drug and alginate. The surface tension, viscosity and pH of the CaCl$_2$ solution were identified as important factors for controlling particle agglomeration and stability as well as drug encapsulation [139].

The experiment with diffusion of 4 and 70 kDa FITC-dextran from calcium alginate microparticles having sizes 20 and 841 (or 948)µm revealed that for the larger particles there was a lower number of crosslinks per surface area. For 500 kDa FITC-dextran the particles were impermeable. The particles were produced by aerosolization method and by inkjet printing. The inkjet engine fires sodium alginate solution through a print head into CaCl$_2$ solution. The particles were used as artificial cells in the study of modeling diffusivity [140].

By aerosolization of alginate solution containing cells (islets isolated from mouse [141], rat or human [142] pancreas) and human serum albumin used as a stabilizer [141] into BaCl$_2$ solution [141,142] containing histidine as buffer and NaCl [139] using an air-driven droplet generator microparticles were produced. Their diameter was 900-1100µm [141] or 700-800µm [142]. After their transplantation diabetic mice became normoglycemic, but hyperglycemia immediately reappeared after explantation of the microparticles. No rejection was observed [141,142]. Each mouse received 900-1000 [141] or 1800 [142] microencapsulated islet equivalents. The peak C-peptide response in these mice was delayed compared with that of healthy mice. No overgrowth of the explanted microparticles was observed. Cytokines were demonstrated to damage encapsulated islet cells in vitro, but there was no cellular reaction [141] or weak cellular reaction at the particle surface. Slight reduction of insulin secretion capacity of rat islets occurred after transplantation [142]. The use of BaCl$_2$ as crosslinker provoked less fibrotic reaction than did CaCl$_2$ [141]. The same technique was used for porcine neonatal pancreatic cell cluster encapsulation, and the same positive effect was shown after their transplantation through a small abdominal incision into the peritoneal cavity of anesthetized diabetic mice. Each mouse received 10000 islet equivalents contained in 2.25ml of the microparticles. Cell differentiation occurred within the particles. Most capsules were free of overgrowth 2 and 20 weeks after
transplantation, although at 6 weeks 43% of the capsules were overgrown [143]. The proliferaton index of the β-cells before and 6 weeks after transplantation was almost the same, but it dramatically decreased at 20 weeks. IgG was deposited on the cells inside the microparticles, i.e. it could diffuse through the capsule, but without causing adverse effects. Another problem was batch-to-batch variation of encapsulated cells [143]. It should be noted, however, that the human clinical trial with barium alginate microparticles having diameter in the range 255-750µm with median average diameter 340µm failed because of necrosis of all the transplanted encapsulated islets of cadaver origin. The capsules were found intact and surrounded by fibrous tissue containing thin-walled capillaries with a mild histiocytic response. Each person received 98200-227900 islet equivalents. The median number was 178200. One person received three islet infusions for the 1st month and one more 6 months later. Another recipient received two infusions 10 months apart. The other two patients received one infusion each. Cytotoxic antibodies were detected in two persons. The loss of the encapsulated cells might occur due to either ischemic necrosis or inflammation [144]. The respective studies were aimed at the development of so-called bioartificial pancreas, i.e. pancreatic islets enclosed within immunobarrier membranes intended for transplantation to patients with type 1 diabetes [138-144]. In another report, Lactobacillus rhamnosus cell suspension in sodium alginate solution was atomized into CaCl₂ solution followed by additional incubation in this solution. The resulting microparticles were recovered by sedimentation, incubated in chitosan solution under shaking, washed with CaCl₂ solution and again recovered by sedimentation. They had size of ~390-450µm, loading efficiency of 38% and were roughly spherical in shape. The possibility of formation of biofilm-like globules inside the particles was shown. Such particles possessed higher resistance (in the terms of cell viability) to freeze-drying and to heat exposure at intermediate temperatures compared to those with encapsulated single cells. However, the resistance to refrigerated storage, simulated gastric juice and high temperatures was similar. Alginate particles were found to be overall superior to carrageenan ones. Chitosan-coated alginate capsules were characterized as the most suitable delivery system for the biofilm-like probiotics. Cell encapsulation represents the more plausible and widely accepted approach than development of heat-resistant probiotic strains via genetic modifications [145].

The synthesis of calcium alginate particles with predefined size

Aerosolization of sodium alginate solution containing mesoporous silica nanoparticles with a diameter around 500nm above calcium-containing solution using an air-dynamical nebulizer led to microspheres with a diameter around 20-30µm and silica nanoparticles inside. If the nanoparticles contained rhodamine 6G, its release from microparticles in phosphate buffer increased with buffer concentration increase (maybe because of alginate solubilization, as discussed below) and decreased with alginate concentration increase, but only up to 1% alginate. Sustained release could be maintained for 40 days. Doxorubicin encapsulation into the nanoparticles suspended in MEM medium enhanced the efficiency of its delivery into cultured HepG2 cells for 4 hours about 1.85-fold. Drug concentrations used for the experiment were 1, 5 and 10µM. Alginate functionalization with RGD-based targeting peptide using carbodiimide chemistry prior to particle synthesis enhanced drug delivery further, viz. about 1.6-fold, as shown for the case of fluorescein isothiocyanate. Functionalized microparticles specifically targeted αβ receptor of HepG2 cells and were taken up by the receptor-mediated endocytosis pathway. At 2-10µg/ml the microparticles without encapsulated doxorubicin were not cytotoxic, as revealed by incubation at 37 °C for 24 hours. The microparticles show significant potential as effective and visually observable transmembrane delivery carriers with high biocompatibility and targeting features for the intracellular controlled release of cell membrane-impermeable drugs. Further development of this nanomaterial could lead to a new generation of nanodevices for biomedical applications [146].
nanoparticles [63]. Porosity and mean pore diameter of calcium alginate beads as well as bead size (2.2mm for control beads) were shown to increase with the concentration of carbon dioxide-forming agent (CaCO₃ or NaHCO₃) incorporated into hydroxypropylmethylcellulose-containing sodium alginate solution being dripped into CaCl₂ solution containing acetic acid. In the case of NaHCO₃, pore size reached 39.7μm exceeding that of control (0.4μm) or that resulting from the use of CaCO₃ (6.1μm). At NaHCO₃: alginate ratio of 1:1 the released CO₂ burst the beads, thus making them nonspherical. Despite the increased porosity and pore size, the release of encapsulated riboflavin to phosphate buffered saline became more prolonged (up to ~2 hours), and the bead surface became smoother with the increasing concentration of CaCO₃, because of the participation of the released Ca²⁺ ions in alginate gelation. In the case of NaHCO₃, the opposite effect on the drug release was observed. The presence of CO₂ also improved bead buoyancy. CaCO₃-containing beads are an excellent candidate for floating drug dosage systems. These systems remain buoyant in the stomach without affecting the gastric emptying rate for a prolonged period [130]. In contrast, our experiments displayed nanoparticle size reduction in the case of aerosolization of sodium alginate solution containing NH₄HCO₃ into CaCl₂ solution containing acetic acid compared to the experiment without CO₂ generation. We ascribed this effect to burst of the nanoparticles at the initial stage of their formation [63].

It should be noted that electrospay in solution is another technique that has a potential of producing calcium alginate particles with predefined size. This technique is based on collection of charged droplets of polymer solution in a glass with crosslinking solution (usually CaCl₂). The control variables influencing particle size and size distribution are applied electric potential, collector distance, flow rate of polymer solution, its viscosity, conductivity, surface tension, concentration and polymer molecular weight. The possibility of variation of the size of the particles in wide limits has been demonstrated for alginate; however, the universal formula for size prediction still does not exist. Electrospay technique is reviewed in [147]; here we will discuss the possibilities of using aerosolization method for the synthesis of calcium alginate particles with predefined size. The discussion related to viscosity is valid for dripping method in general. Forced dissolution is also not limited to the particles produced using aerosolization method.

**Changing the viscosity of sodium alginate solution**

If alginate beads are produced by dripping sodium alginate into CaCl₂ solution, their size depends on the drop size, which is influenced by diameter of a nozzle and viscosity of alginate solution. The size of air-dried beads was also shown to decrease with hardening time increase, possibly because of leak of the encapsulated drug (theophylline), and some increase with hardening time increase, possibly because of leak of the solution. The size of air-dried beads was also shown to decrease with hardening time increase, possibly because of leak of the solution. The size of air-dried beads was also shown to decrease with hardening time increase, possibly because of leak of the solution. The size of air-dried beads was also shown to decrease with hardening time increase, possibly because of leak of the solution.

Fluidized-bed-dried as well as air-dried beads shrank during drying, but had much smoother surface and were less porous than freeze-dried ones. The two latter drying types did not cause the beads to become irregularly shaped. Theophylline, magnesium stearate and Kollidon K30 were admixed to sodium alginate solution before bead preparation. Bead size was in a millimeter range. The beads were intended for further incorporation into more complex drug delivery systems [148]. By means of dropwise addition of sodium alginate solution containing hemoglobin to CaCl₂ solution with the aid of a syringe beads with an average diameter of about 2mm were prepared, and at lower alginate concentration (and therefore lower viscosity) alginate microparticles with an average diameter of about 110μm were produced. CaCl₂ concentration was simultaneously lowered (in order to avoid aggregation of the microparticles). In order to avoid hemoglobin oxidation observed at the time of encapsulation, its concentration was increased 10 times, and it led to particle size increase to 156μm. Encapsulation and loading efficiencies for the latter case were 38% and 50%, respectively. A tight interaction between alginate and hemoglobin affecting the release of oxygen was observed. The particles could possibly be applied for therapeutic approaches aimed at achieving the remediation of particular pathologic conditions [149]. For effervescent atomization of glyceral aqueous solutions it was shown that with increasing solution concentration (and therefore viscosity) the number of droplets of larger diameter increased and the droplet size histogram was wider [150]. In no case drops smaller than 0.25mm were observed, although this might be due to the limitations of their measurement method employing a camera. In another study that also dealt with atomization of glyceral aqueous solutions but employed more advanced drop sizing technique it was similarly shown that the effect of viscosity acted to hinder the growth of instabilities in the liquid sheet, therefore resulting in longer break-up times and larger droplets [134]. A study of tapioca starch atomization also revealed that the reduction in viscosity not only eased the flow through the nozzle orifice but also boosted the surface wave oscillations and instabilities helping to break the solution stream into fine droplet spray patterns. Using chemically modified starch with opened ring the increase of variations in Sauter mean diameter across the spray centreline with an increase in solution viscosity and surface tension was demonstrated [135].

The viscosity of sodium alginate solution can be decreased in following ways:

- **concentration decrease.** The dependence of alginate viscosity on its concentration and shear rate is nonlinear [138].
- **Addition of a supporting polymer that can form interpolymer complexes with sodium alginate, such as polyethylene oxide.**
- **Addition of a surfactant (Triton X-100 can also form a complex with alginate leading to double effect); it may also reduce the surface tension.**
- **Addition of a cosolvent being able to interact with sodium alginate, such as DMSO [151].**
- **Addition of electrolytes [45,139].** We have observed calcium alginate nanoparticle size decrease if sodium alginate used for their preparation was dissolved in physiological saline instead of water [63], but it may affect gel strength because the
amount of calcium bound to alginate after addition of certain amount of Ca(NO$_3$)$_2$ solution to sodium alginate solution depends on calcium and sodium concentrations in solution. The volume fraction of hydrogel depends on the same parameters [152].

- Increase in temperature and/or in pressure, as shown for starch. The solution density and surface tension also decreased monotonically and reached their constant values above 90 °C [135]. We have observed calcium alginate nanoparticle size decrease if the nebulizer containing sodium alginate solution was heated [63].

- Using sodium alginate of lower molecular weight [151].

- Acid hydrolysis [153,154]. This is suitable only for alginic acid with subsequent neutralization to convert it to alginate.

- Alkaline degradation [45]. The viscosity of sodium alginate solution in water was shown to decrease during the first two weeks of storage because of hydrolysis, but this approach was not recommended [151]. Alginate hydrolysis may also occur at the time of its isolation from algae [152].

- Oxidation with H$_2$O$_2$ added [154,155] or generated by interaction of reduced glutathione [155], cysteine [153,155], cysteamine, ascorbic acid [153,155], sodium ascorbate [155], phenylhydrazine, dihydroxymaleic acid, thioglycolic acid, sulfite, dithionite, phloroglucinol, sodium hydrogen sulfide, 'fucosan' (endogenous phenolic compounds of brown algae) [153], etc. with atmospheric oxygen. Further reaction of hydrogen peroxide with the reducing compounds generates free radicals oxidizing alginate more rapidly. Ferrous ions can also be used for the same purpose because they decompose hydrogen peroxide leading to the same radicals, but at high concentrations they form gel with alginate [153]. When polynanunorote blocks having ~40 sugar units produced by sodium alginate treatment with 0.5 M HCl at 100 °C for 8 hours were further treated with 5% H$_2$O$_2$ for 30 min, oligomers having 1-15 sugar residues were formed [154]. But small fragments of alginate are not suitable for gelation.

- Chemical modification of alginate can both increase and decrease its viscosity, but it is usually accompanied by chain destruction and at high substitution degrees makes gelation impossible [45].

**Drop evaporation in preformed aerosol**

The total output from a nebulizer is made of aerosol and a significant component of water vapor. As the evaporative component increases, the cumulative solute output drops. The concentration of solute in the nebulizer solution increases with operating time in a nonlinear manner. At the same time the temperature of nebulizer solution can drop up to 12 °C lower during operation, but then equilibrium is reached. As the driving air pressure is increased, the equilibrium temperature is lowered, but the time to reach equilibrium remains constant. Increase in the ambient humidity of the supply air and decrease in the volume flow rate reduce solvent evaporation [156]. Heating of preformed diluted aerosol in order to induce water evaporation and therefore decrease drop size was proposed as a method to convert aerosols with micrometer-sized drops produced by common medical nebulizers to those with nanosized drops. The best heater type was air jacketed tube, and preheated air was flowing through the outer line in the opposite direction of the aerosol stream. In the case of aerosol of 0.1% albuterol sulfate it was possible to convert aerosol having mass mean aerodynamic diameter of 4.69µm to this having mass mean geometric diameter of 0.43µm. Dry nanoparticles of predetermined sizes can be formed by reducing drug concentration. The possibility to predict final drop size by means of computational fluid dynamics was demonstrated [157].

**The attempts to predict aerosol drop size distribution**

For several atomizer types there are equations allowing to predict Sauter mean diameter of the aerosol droplet. This, in turn, allows one to predict drop size distribution. The most important input parameter is viscosity because it has a cohesive effect and impedes the disintegration mechanisms leading to droplet formation. Other parameters for pressure-jet atomizers are liquid surface tension, air density, liquid density, discharge orifice diameter, injection pressure differential across the nozzle, half spray angle, and liquid mass flux. For pre-filming airblast atomizers the first three parameters are needed along with air velocity and air/liquid ratio; for plain-jet airblast atomizers discharge orifice diameter is needed as well [158]. Another model allows predicting number-based dimensionless drop size distribution near the region of their formation, i.e. not that of the drops actually reaching CaCl$_2$ solution. The input parameters are gas velocity, liquid velocity at the atomizer exit, gas density, liquid density, surface tension, drop mass mean diameter, linear growth rate of the sinusous or varicose waves, origin shift parameter, dimensionless critical diameter, and at least two more parameters [159]. A more advanced model based on maximum entropy principle, as were the ones mentioned before, is aimed at prediction of drop size distribution at the end of the primary breakup region, i.e. still not of that of final drops. At least the following input parameters are needed: mass flow rate, mean velocity of jet in nozzle outlet, gas density, liquid density, surface tension, drop mass mean diameter, shape factor for velocity profile, Reynolds number, breakup length and thickness of the liquid sheet [136]. Although some of these parameters can be substituted for other ones, unfortunately in real experiment of alginate aerosolization they are still very difficult to measure, and it makes the prediction impossible. In general, the predictive power of maximum entropy method is rather poor; it produces good agreement with the experimental data only after adjustment of parameters or if the representative drop diameter is known from direct measurements [160]. Obviously, in these two cases the method relies completely on the data it must predict, i.e. it becomes just a way of fitting the existing data and is no longer predictive. Moreover, the equations do not take into account the possibility of drop size change resulting from water evaporation discussed above. There is also a discrete probability function approach aimed at predicting drop size distribution from the fluctuating initial conditions, but it is again limited to primary atomization and requires prior knowledge of probability density function of the fluctuating initial conditions. These functions cannot be currently
the presence of alginate lyase was reported. Before the dissolution with an average diameter of about 2mm during 30 minutes of 1 hour of incubation [118]. Complete dissolution of alginate beads chitosan-covered dried calcium alginate beads with a size of ~1mm Phosphate buffered saline accelerated protein release from alginate thus accelerating the release of encapsulated rhodamine 6G [146]. was also demonstrated to dissolve calcium alginate [138,145]. proliferating and non-differentiating state [161]. Phosphate ion engineering. The printed cells could not degrade the surrounding hydrogel. Treatment with sodium citrate may help to improve the was proven for human corneal epithelial cells encapsulated in this sodium citrate was greater than that of alginate, the degradation was shown the possibility of using this reaction for adjustment of the size of preformed calcium alginate nanoparticles produced from commercially available low viscosity sodium alginate [63]. Sodium citrate can also be used for the same purpose, and the relation curve between the gelatin/alginate/collagen hydrogel degradation time and mole ratio citrate: alginate was obtained. If the amount of sodium citrate was greater than that of alginate, the degradation was too quickly to be controllable. Biocompatibility of sodium citrate was proven for human corneal epithelial cells encapsulated in this hydrogel. Treatment with sodium citrate may help to improve the alginate bioink system for the application of 3D bioprinting in tissue engineering. The printed cells could not degrade the surrounding alginate matrix themselves, causing them to remain in a poorly proliferating and non-differentiating state [161]. Phosphate ion was also demonstrated to dissolve calcium alginate [138,145] thus accelerating the release of encapsulated rhodamine 6G [146]. Phosphate buffered saline accelerated protein release from alginate particles by their degradation [112,126,138]. The dissolution of chitosan-covered dried calcium alginate beads with a size of ~1mm in simulated intestinal fluid (NaOH + KH₂PO₄) was complete after 1 hour of incubation [118]. Complete dissolution of alginate beads with an average diameter of about 2mm during 30 minutes of incubation in phosphate buffered saline at pH 7 under stirring in the presence of alginate lyase was reported. Before the dissolution study, the beads had been washed with phosphate buffered saline 3 times in order to remove excess Ca²⁺ [149]. It should be pointed out that some authors claim their alginate particles as demonstrating pH-dependent drug release while using phosphate buffers for alkaline pH, but in fact the presence of phosphate is the main cause of drug release because of particle dissolution. Phosphate buffers are often used as ‘simulated intestinal fluid’, sometimes even without providing their composition.

Non-crosslinking cations compete with calcium ions for carboxyl groups of alginate and therefore can disrupt electrostatic interactions between alginate chains. The affinity of the cations to alginate grows in the following order: K⁺ < Na⁺ < Mg²⁺. But this does not lead to bead disintegration, except at high salt concentration (50mM for NaCl or KCl and 10mM for MgCl₂). At lower salt concentrations bead swelling occurs if the beads were previously dried. This happens because at low concentrations the salts disrupt only dimer-dimer associations, but individual egg-box dimers remain intact. These dimer-dimer associations appear during bead drying and do not exist in fresh beads. Bead swelling in pure water did not occur. The beads used were prepared by drying sodium alginate solution into CaCl₂ solution under stirring with subsequent incubation in CaCl₂ solution. Bead diameter was ~3.2mm, the bead weight was 17.9mg before drying and 0.6mg after drying [162]. The applicability of this method to nanoparticles was also shown [73,95]. Alginate beads were chosen as a model system in the study of the changes in the food microstructure during drying in order to improve the quality of dried foods [162].

To our opinion, aerosolization method is preferable if calcium alginate nanoparticles with adjustable size and narrow size distribution, without the admixture of microparticles and without surfactants are needed in moderate quantities. (For beads and microparticles it may be easier to use dripping method). Although theoretical prediction of nanoparticle size has not been achieved yet, it is possible to obtain calibration data for a particular nebulizer. Moreover, particle size can be decreased in a controllable way by forced dissolution, as discussed above. This method allows fine adjustment of particle size after proper calibration. Another possible method of producing nanoparticles with partially predictable size is electrospray, but it is less safe because of high voltage.

**Encapsulation of cyclodextrin inclusion complexes in alginate particles**

Cyclodextrins are torus-like macro-rings built up from glycopyranose units (Figure 7). They have hydrophilic sites on the outer surface and a hydrophobic cavity inside. The diameter of the one end of the cavity is larger than that of the opposite end. There are numerous cyclodextrins, the most important ones being α-, β- and γ-cyclodextrins, comprising 6, 7, and 8 glucose units and having cavity diameter of 4.7-5.3, 6.0-6.5 and 7.5-8.3 Å, respectively. In an aqueous solution, the water molecules inside the cavity can be readily substituted by appropriate ‘guest molecules’ which are less polar than water. Upon dissolving the inclusion complexes formed, an equilibrium is established between dissociated and associated species [163].
Cyclodextrins can even form inclusion complexes with various volatile substances, although most of these complexes are rather unstable in the presence of water. The results of spray-drying complex mixtures of numerous flavors briefly incubated with cyclodextrins showed that γ-cyclodextrin was generally the most effective in retaining the flavor compounds during spray-drying, but the losses of volatilities on storage were greatest for γ-cyclodextrin and least in the case of α-cyclodextrin [164]. The synthesis of an inclusion complex of eugenol and β-cyclodextrin and its antifungal activity helping to preserve fruits has been reported [165]. Allyl isothiocyanate was shown to be released continuously from inclusion complexes with α- and β-cyclodextrins for 120 hours even at 98% relative humidity, but initial burst release was observed [166]. Both complexes had fungidal activity against Penicillium expansum, and the complex with β-cyclodextrin displayed bactericidal activity against E.coli and Listeria monocytogenes as well [167]. Inclusion complexes of 2-nonenone and β-cyclodextrin were active against fungal pathogen Botrytis cinerea, which led to the grey rot on strawberries, blueberries, raspberries and table grapes [168]: an inclusion complex β-cyclodextrin-hexanal was active against Colletotrichum acutatum, a fungus that is among the most common cases of berry decay [169]. Three techniques of the synthesis of hydroxypropyl-β-cyclodextrin-capsaicin inclusion complex were reported [170-172], although the aqueous solutions of the complex still contained free capsaicin [172]. Even acetaldehyde, a rather small molecule, was successfully encapsulated in β-cyclodextrin [173]. A more advanced technique proposed was acetaldehyde-diethylacetal encapsulation by hydroxypropyl-β-cyclodextrin. The inclusion complex was stable under normal shelf-storage conditions for one year, but immediately decomposed on contact with water [174]. However, smaller molecules show weaker binding. Similarly modified α- and β-cyclodextrins were shown to entrap aniline vapor in higher amounts than benzene [175]. Benzene binds much weaker than tert-butylbenzene with both native and carboxymethyl-β-cyclodextrin, as shown by molecular dynamics simulations [176]. The extracts of cinnamon bark and clave bud without isolation of the essential substances were used for encapsulation, and the formation of microparticles of the inclusion complexes by means of freeze-drying was reported. The antibacterial activity of the complexes against Salmonella enterica serovar Typhimurium LT2 and Listeria innocua was higher than that of the nonencapsulated extracts [177]. Numerous other examples of complexes of cyclodextrins with volatiles, the methods of preparing them, the methods by which the complex formation can be evidenced, the applications of the complexes and some other aspects are reviewed in [178].

It can be expected that alginate encapsulation of the inclusion complexes instead of free molecules may give an opportunity for additional retardation of their release.

The published examples of the immobilization of cyclodextrin inclusion complexes in alginate

The cationic polymer produced from β-cyclodextrin, epichlorohydrin and choline chloride was shown to form complexes with insulin, despite insulin molecules being too bulky to be wholly included into the cavity of cyclodextrin. Dropwise addition of sodium alginate solution containing the complex to CaCl₂ solution followed by chitosan addition led to beads [179]: the addition of CaCl₂ and then chitosan [180] or trimethylchitosan [181] to sodium alginate solution containing the complex led to nanoparticles [180,181]. The beads had irregular shape and relatively smooth surface with some wrinkles due to lyophilization. The encapsulation efficiency was up to 72%, being influenced by cationic polymer composition and amount, complexation time and CaCl₂ concentration. The beads released around 20% of insulin in simulated gastrointestinal fluid [179]. Their size was in the millimeter range, as stated later [180]. They were initially claimed as a promising system for improving insulin oral delivery efficiency [179], but they did not show significant improvement of insulin release efficiency in the following trials [180]. Most nanoparticles coated with chitosan had irregular shape and size in the range of 100-200nm. Encapsulation and loading efficiencies were up to 87% and 9.5%, respectively [180]. After freeze-drying in the presence of sucrose the nanoparticles coated with trimethylchitosan were spherical or subspherical. Aggregation was not observed. Their size was influenced by β-cyclodextrin, sodium alginate and CaCl₂ concentrations, and could
be varied from 187±20nm to 2393±1219nm. The same parameters influenced polydispersity index. For optimized formulation it was 0.362. Encapsulation efficiency was more than 93%. Higher encapsulation efficiency was associated with higher concentrations of trimethylchitosan and β-cyclodextrin as well as with lower pH of β-cyclodextrin. Loading efficiency was 5% [181]. The nanoparticles coated with chitosan released 48% of encapsulated insulin during 2 hours of incubation in simulated gastrointestinal fluid, and 40% of insulin was released during subsequent incubation in simulated intestinal fluid (containing phosphate) for 4 hours. In the case of uncomplexed insulin encapsulation these values were 60% and 18%, respectively, and the mean particle size exceeded 700nm. The release profile could be controlled by varying the molecular structure of the cationic polymer besides the alginate/chitosan mass ratio. The latter parameter along with cationic polymer content also affected the mean particle size [180]. The nanoparticles coated with trimethylchitosan released 5% and 94% of insulin in these media, respectively. Burst release of most of encapsulated insulin occurred in simulated intestinal fluid. Insulin permeation across Caco-2 cell monolayer from apical to basolateral side was significantly increased by these nanoparticles if compared with the nanoparticles prepared with unmodified β-cyclodextrin and with free insulin. 1ml of medium containing encapsulated insulin at 0.4mg/ml was used for the permeability studies, and almost 9% of insulin was transported for 240 min. The monolayer was grown on polyethylene terephthalate membrane filters with a pore size of 0.4μm [181]. The system provided enhanced insulin stabilization in the presence of pepsin [180]. The nanoparticles can be considered as a good candidate for oral insulin delivery [180,181].

Covalent binding of β-cyclodextrin to alginate via linkers was also reported. In the case of ethylenediamine linker the binding was achieved by means of carbodiimide chemistry. In contrast, binding through 1,6-hexanediame linker was carried out in a more complex way, beginning with turning sodium alginate into alginic acid with its subsequent neutralization with tetrabutylammonium hydroxide. Cyclodextrin was activated with tosyl group prior to the synthesis. Addition of Pluronic F108 to the modified alginate at 4% and 6% (w/v) for 24 hours with primary human mesenchymal stem cells seeded at 20000cells/well in 500μl/well of the growth medium showed no cytotoxicity. Biologically relevant mechanical properties of the hydrogels would be advantageous for biomedical applications such as drug delivery and cell transplantation [182].

It should be noted, however, that in the case of invertebrate immobilized in calcium alginate beads addition of β-cyclodextrin led to bigger pores and higher rate of release. The encapsulation efficiency dropped from 92% to 82%. The beads were prepared by dripping sodium alginate solution containing the enzyme into CaCl2 solution in acetate buffer (pH 3.8) containing β-cyclodextrin and trehalose. Trehalose protected the encapsulated enzyme during freezing, drying and thermal treatment, and β-cyclodextrin protected it during bead formation. The size of the beads depended on the drying technique and the composition, but was in the millimeter range. The enzyme release depended on pH and drying technique, but it took several hours in all the cases. The study was aimed at providing new alternatives for enzyme stabilization and release management in order to reduce costs and increase the productivity of the technological process employing invertase [183]. For spherical microparticles produced by spray drying the solutions of raffinose or trehalose containing hydroxypolypropyl-β-cyclodextrin in methanol: n-butyl acetate containing the mixture of porosity with cyclodextrin weight fraction up to 40% with subsequent decrease was revealed. The addition of cyclodextrin resulted in 2-times decrease of mass median aerodynamic diameters to about 2μm. This effect was ascribed to high porosity diminishing the contact areas between particles and therefore hindering aggregation. The preparations of microparticles had improved physical characteristics, physicochemical stability and aerosolization, suggesting their potential as carrier system for pulmonary delivery of biomolecules [184].

The unrealized possibilities of further improvement of small molecule encapsulation and release retardation

In theory, there is a way of encapsulation of small molecules in cyclodextrins by means of cavity size tuning, but there has been no attempt to combine these techniques with the encapsulation of the resulting inclusion complexes in alginate particles. Enzymatic synthesis of cyclothioamalts, viz. cyclo-(α(1 → 4)-4-thiomaltotetraoside, -thiomaltopentaoside and -thiomaltotriose has been reported [185]. They must have smaller cavities than that of α-cyclodextrin. Peracetates of cyclic (1 → 6)-β-D-glucan trimer and tetramer were chemically synthesized from derivatized linear tri- and tetrasaccharides, and their cavity hole size was calculated as 1 and 3.3 Å, respectively [186]. It should be noted, however, that the cavity may have dynamic nature. For example, a glycophane containing two benzoic acid spacers displayed great conformational flexibility [187].

Several other possibilities of cyclodextrin cavity tuning without decreasing its size exist. A single oxygen atom in the ring skeleton of β-cyclodextrin can be replaced by a disulfide bond in two different configurations making the cavity asymmetric [188]. In permethylated α- and β-cyclodextrins β-(1,4)-glucosidic bond can be introduced instead of α-(1,4)-glucosidic one to alter the cavity shape [189,190]. This technique was extended to 2,6-di-O-methylated α- and β-cyclodextrins [190]. The α-cyclodextrin analogues having a thiophene or triazole fragment inserted in the ring as well as α-, β- and γ-cyclodextrin analogues carrying fragments with triple bonds were reported [191].
A theoretically possible way to retard the release of encapsulated molecules from the inclusion complexes is steric hindrance introduction. An interesting example is the synthesis of hyperbranched polyglycerol grafted from β-cyclodextrin. This molecule can be further modified with amine groups [192]. But this technique was not intended for the combination with alginate.

In conclusion, the encapsulation of drugs in the form of inclusion complexes with cyclodextrins appears rather promising and underexplored. This is especially important for unstable or small molecules provided they can form the inclusion complexes. Cavity size tuning and steric hindrance introduction will probably be of limited applicability because they make the synthesis expensive, and biocompatibility of the resulting complexes is questionable. Another problem is that the stability of many inclusion complexes in water may lead to their decomposition at the time of alginate particle formation. Nevertheless, the use of cyclodextrins will surely provide additional power to the encapsulation methods.

**The Decrease of Alginate Pore Size**

Calcium alginate pore size is usually assumed to be in the range of 5-200nm. It is a so common assumption that reviews [193] may give it citing other reviews, and sometimes [42] it is given even without reference. Nevertheless, it is not always the case. The pores of calcium alginate hydrogel produced by treatment of spin-coated sodium alginate with CaCl₂ were found to have typical diameters of 50-300nm. The study dealt with calcium alginate hydrogel as a model of extracellular polymeric substances because of similar physicochemical properties and architecture [194]. In another study, alginate was spin-coated onto a poly-L-lysine-coated glass slide and then treated with CaCl₂. Pore size range of 4-18nm was observed. Moreover, their review of previously published literature indicated that pore sizes from 4nm up to 10.9μm had been reported [195]. In any case, there are some pores having much larger size than that of protein molecules. This is the main cause of burst release of encapsulated substances. Obviously, the situation with low molecular weight substances is even much worse than with proteins. The diffusion coefficients of glucose and ethanol in 2% calcium alginate are only slightly lower than in water. Their diffusion is mainly controlled by the rate at which they diffuse through water occupying the pores. Glucose and ethanol concentrations have no effect on the respective diffusion coefficients. Their decrease with increasing alginate concentration is probably due to a decrease in the number and length of the pores rather than a decrease in the pore diameters. The diffusion coefficients were needed to model reaction and diffusion in immobilized cell reactors [196].

The size of the pore openings can be effectively reduced by chitosan coating, as revealed by the study dealing with hydrogel film acting as a model of an immuno-isolation membrane [195]. Besides this effect, chitosan layer provides an additional diffusional barrier. Another option is the use of additional crosslinking agents.

In one example, alginate beads containing immobilized xylanase were produced by extruding sodium alginate solution containing the enzyme through a needle to CaCl₂ solution, and then additional overnight incubation in CaCl₂ solution was carried out. The beads were then incubated with 1% glutaraldehyde at pH 6.0 for 90 minutes under stirring and washed with acetate buffer. The immobilization shifted the enzyme pH optimum from pH 5.0 to 5.5, and treatment with glutaraldehyde shifted it further to pH 6.0. Encapsulation efficiency before and after glutaraldehyde application was 41% and 91%, respectively. Treatment with glutaraldehyde almost did not provide any additional enzyme stabilization at acidic or alkaline pH. Enzyme thermostability, stability during storage and in the presence of chemicals (HgCl₂, etc.) was increased, but the enzyme was operating more slowly. No additional shift of temperature optimum was recorded, but there was a shift to higher temperatures if compared with the free enzyme. After 4 reaction cycles the enzyme activity started to decrease, and after 12 cycles most of the enzyme leaked out. It was also washed out by bead treatment with sodium dodecyl sulfate [42]. Therefore, there was no ovalent linkage of the enzyme with alginate. The effect of glutaraldehyde was explained by enzyme crosslinking. Alginate crosslinking was not considered, although it was likely to happen. The beads could be efficiently used for fruit juice clarification and yield improvement [42].

In another example, sodium alginate solution containing diclofenac sodium [197] or neem seed oil [198] was added dropwise into methanol containing 1% glutaraldehyde and 1% of 10 M HCl [197] or 1% of 1 M HCl [198]. This resulted in formation of almost spherical beads with nonporous smooth surface and size 1.2-1.6mm [197,198]. Crosslinking time did not influence the particle size but decreased the release rate and bead swelling in water. Diclofenac release was studied at pH 1 for 3 hours and then at pH 7.4 [197], and that of neem seed oil in 0.1% Tween 80 [198]. The beads formed at 40 °C showed lower release rate than those formed at 25 °C, but this effect was less prominent [198]. Both parameters caused encapsulation efficiency decrease because of drug leaching during crosslinking. Increasing concentration of the substance being encapsulated in sodium alginate solution decreased encapsulation efficiency and increased release rate [197,198].

The longest diclofenac release time achieved for the beads with encapsulation efficiency of >55% was 300 min. Controlled release system is needed for diclofenac because of its short biological half-life and side effects such as ulcer bleeding or perforation of intestinal wall [197]. For the beads loaded with neem seed oil these parameters were >72% and 25 hours, respectively. There was no chemical interaction between neem seed oil and glutaraldehyde. The aim of the study was to create a controlled release formulation of a pesticide with a minimum residual toxicity [198].

Other techniques of crosslinking alginate with glutaraldehyde exist, but they were reported for alginate fibers or membranes, not for particles:

- 2ml of glutaraldehyde and 2ml of concentrated HCl in 180ml of 1:2 (v/v) water-acetone mixture for ~10 hours, then wash with water and drying [199]
- 2.5ml of glutaraldehyde and 2.5ml of concentrated HCl in 30:70 water:acetone mixture for ~12-14 hours, than wash with water and drying at 40 °C [200,201]
• 5% (v/v) glutaraldehyde and 1% (v/v) HCl in 90:10 methanol: water for 4 hours, then wash with water [202]
• 250mM glutaraldehyde and 0.05 M HCl in 50% ethanol for 48 hours at room temperature, then wash with water and drying under reduced pressure [151]

• 5-20% (v/v) glutaraldehyde (optimized concentration was 10-15%) and 0.05% (v/v) HCl in acetone at 40 °C for 48 hours, then wash with methanol, immersion in methanol for 24 hours at 40 °C, and vacuum drying. Some aldehyde residues were pendent from alginate chains instead of actual crosslinking [203]

• ≤3% glutaraldehyde and 0.1% of 35% HCl in dioxane at 50 °C for 4 hours with subsequent neutralization in 80:20 (v/v) methanol/0.4M NaOH mixture at room temperature for 20min, then wash with 80% methanol. In dioxane better crosslinking efficiency than in methanol or acetone was achieved, but <10% of alginate still remained water soluble [204].

Dropwise extrusion of sodium alginate solution into 0.05 M HCl with subsequent incubation for 4 hours resulted in beads having mean diameter of 2.9mm. The beads were crosslinked in a solution containing 1.58 M glutaraldehyde (3.16 M inside the beads) and 0.38 M HCl [205,206]. Control of reaction kinetics allowed to adjust the mean pore size from 720nm to less than 35nm. However, microscopy studies revealed pore size distribution of 25-200nm in the latter case [205]. In one example, pore size changed from less than 0.1µm in acid solution to almost 0.5µm in alkaline solution because of reversible gel swelling [206]. The time required to crosslink available hydroxyls could range from 1 to over 10 hours depending on the reaction conditions. The activity of guluronic and mannuronic acid residues was shown to be equal. The reaction had zero, second and first order with respect to alginate concentration, glutaraldehyde concentration, and acid concentration, respectively [205]. The addition of aprotic solvent (acetone, dioxane, dimethyl sulfoxide) to the crosslinking medium enhanced the reaction rate, while the use of protic solvents (alcohols) decreased it [206]. The reaction rate also increased with alginate molecular mass and with temperature. At room temperature the expected maximum crosslinking density (~5 monomeric units between crosslink points) was not reached because of glutaraldehyde loss in a parallel oxidative degradation reaction, but at 40-50 °C it was reached. At initial glutaraldehyde concentration less than 0.9 M the reaction was more diffusional controlled [205]. Glutaraldehyde can also turn into polyyclic or α,β-unsaturated polyglutaraldehyde. These products may react further with alginate to form complex network structures. Some fraction of glutaraldehyde reacted only with one hexuronic acid residue, i.e. it was not involved in crosslinking [206]. Detailed mechanism of the crosslinking reaction was also elucidated [205]. Crosslinked alginate can potentially be used as a pulsatile oral delivery vehicle for protein therapeutics, as a superabsorbent for environmental cleanup application, or as a scaffold for tissue engineered implants [205,206].

A technique of crosslinking calcium alginate nanoparticles using epichlorohydrin was reported. In this case, dried nanofibers were soaked in 50% ethanol containing 5M epichlorohydrin and 0.1M NaOH for 24 hours. The reaction was stopped by the addition of 5M HCl. The nanofibers were washed with 75% ethanol, then with water and dried under reduced pressure. The approach may be beneficial to the production of nanofibrous matrices mimicking the native extracellular matrix for a wide range of tissue engineering applications [151].

Tannic acid can be used as an additional noncovalent crosslinker. It was demonstrated for the nanoparticles produced by dropwise addition of CaCl₂ to the solution containing sodium alginate, tannic acid and diltiazem hydrochloride under stirring. The resultant suspension was aged for 10min followed by sonication and centrifugation, and the nanoparticles were resuspended in water and sonicated again. Drug encapsulation efficiency was found to increase with tannic acid concentration from 15% to 66%. Blank calcium alginate nanoparticles had the diameter of 475nm with polydispersity index of 0.56 and zeta potential of -23mV. Introduction of drug and tannic acid caused only a slight change of these parameters. All the nanoparticles were discrete and spherical. Tannic acid reduced the drug burst release within 30min in 0.1 M HCl from 44% to 11%. After 2 hours in this acidic medium the nanoparticles were transferred to phosphate buffered saline, and tannic acid reduced the cumulative drug release from 100% to 5% and prevented disintegration of the nanoparticles. Diltiazem hydrochloride is used in the treatment of hypertension, asthma and arrhythmia, but in the case of non encapsulated drug frequent administration is required [207]. We tried to produce crosslinked calcium alginate nanoparticles using aerosolization of sodium alginate solution containing glutathione (a drug being encapsulated) and phytic acid into CaCl₂ solution, but this technique did not prevent burst release if sodium alginate solution pH was not readjusted. Nevertheless, the attempt of using sodium cromoglicate instead of phytic acid led to more prolonged glutathione release even without pH adjustment, but overnight incubation of the nanoparticles in 1.11% glutaraldehyde in 0.4% HCl gave better results [63].

Several other methods of alginate crosslinking exist. One of them is based on activation of carboxyl groups of tetrabutylammonium alginate with subsequent crosslinking them using a diamine in the presence of triethylamine taken as catalyst. Another one is based on linking protonated carboxyl groups to hydroxyl groups using the carbodiimide coupling reagent. There is also a method in which partial oxidation of hydroxyl groups to aldehyde ones with subsequent crosslinking with adipic dihydrazide is used [45]. Methacrylated alginate can form hydrogels by free radical polymerization. If alginate with immobilized ligands is used for cell encapsulation, the cells can act as crosslinkers themselves, but the network exhibits low strength and toughness, as reviewed in [208].

Therefore, coating and additional crosslinking do lead to pore diameter reduction. However, no universal solution of the burst release problem exists. Crosslinking effectively reduces mean pore size, but some portion of big pores remains or appears during particle swelling. Moreover, drugs of low molecular weight may leak out during crosslinking. Coating is a powerful method. Besides
porosity decrease, it allows surface modification and charge adjustment. However, it ultimately increases particle size.

Conclusion

The characteristics of alginate such as its natural origin, abundance in nature, biocompatibility, biodegradability and hydrogel formation upon crosslinking with calcium ions make it a perfect base for micro- and nanoparticles for drug delivery and other applications. This conclusion is supported by numerous publications dealing with them, indicating significant interest of the scientific community to this versatile biopolymer. Moreover, alginate can be combined with additional biocompatible drug release retardants, e.g. cyclodextrins, creating additional largely unexplored possibilities for its clinical, agricultural and other use.

However, a question arises: why alginate nanoparticles are still not in use in real clinics, despite their great potential? Bulk alginate is already in use as an excipient in tablets, component of wound healing tapes and even as a dietary supplement, but its nanoparticulate form is not.

A very important or even the main reason may be the burst release of the encapsulated drugs. This is the main problem in creating alginate-based drug formulations, and it is caused by inherent properties of calcium alginate hydrogel - porosity and reversibility of gelation. Both problems may be overcome by covalent crosslinking and coating. However, chemically modified alginate is no longer natural and biodegradable, and the currently available coating methods either do not provide sufficient drug release retardation or lead to a so huge increase of the particle size that it becomes dangerous because of the possible embolism. Moreover, there is no method of producing small (i.e. not submicron-sized) calcium alginate nanoparticles with predictable size and guaranteed narrow size distribution without employing chemical surfactants, the safety of which is questionable. One should also remember that initially 'small' dry nanoparticle can become 4 times larger after swelling. Another problem is that hydrogel strength increases with alginate polymerization degree, but alginate of high molecular weight is far above the renal clearance threshold.

Nevertheless, the above-mentioned concerns do not override the wide perspectives of clinical use of alginate particles. New investigations must be directed to the use of alginate particles in the wide perspectives of clinical use of alginate nanoparticles with predictable size and guaranteed narrow size distribution without employing chemical surfactants, the safety of which is questionable. One should also remember that initially 'small' dry nanoparticle can become 4 times larger after swelling. Another problem is that hydrogel strength increases with alginate polymerization degree, but alginate of high molecular weight is far above the renal clearance threshold.

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