Functional importance of coacervation to convert calcium polyphosphate nanoparticles into the physiologically active state

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1. Introduction

The first precise visualization and documentation of particles around 100 nm came from R. A. Zsigmondy [1] using dark field ultramicroscopy; he coined the term “nanoparticles” for particles in the 10–100 nm range. Soon later, Langmuir [2] and Blodgett [3] introduced the concept of monolayer surface chemistry [4]. Nanoparticles (NP) occur as fine colloidal suspensions in the atmosphere, originating from volcanic eruptions or produced by human activities [5]. NP also existphysiologically, e.g. in blood platelets, consisting of polyphosphate (polyP) with an average size of 100–200 nm [6,7]. The formation of the polyP NP proceeds through polyP precipitation in the presence of divalent metal ions, especially Ca2+. Prior to the present report, our group succeeded in preparing such polyP nanoparticles from Na-polyphosphate (Na-polyP)
together with Ca\(^{2+}\) in a bioinspired way.

In most previous studies on polyP, the physico-chemical mechanism by which NP become functionally active has remained open. Going back to Bütschi [8], who introduced studies on colloidal dispersions from water-in-water emulsions [9], it was recognized that solid phases (as a precursor stage) must undergo a transition into an aqueous intermediate product that is biologically and functionally active. The resulting intermediate associates, which consist of amorphous, non-crystalline particles, are also referred to as coacervates [10]. The process of coacervation is based on the formation of polymer droplets in solution, driven by the separation into two liquid phases, a highly concentrated colloidal phase, the coacervate, and a highly dilute colloidal phase [11]. The two colloidal aqueous phases are immiscible and are in thermodynamic equilibrium. The reversible transition between an immiscible and a miscible colloidal system is mandatory for the functional activity of a coacervate. While large positive or negative zeta potentials (e.g., around +30 and −30 mV) of particles dispersed in solution cause repulsion, leading to stabilization of (bioinert) NP, low zeta potentials promote coacervate formation. This property is exploited by the polyP/Ca\(^{2+}\)-NP when added to a physiological environment. When these particles are transferred to a proteinaceous milieu, their zeta potential drops and transformation into the coacervate phase occurs [12]. The coacervate phase represents the biologically active form of polyP and similar potentially active molecules [12,13].

The relevance of coacervation as a physiological control mechanism between inert NP and functionally active coacervate is usually underestimated [12,14]. For the formation of the NP, a superstoichiometric ratio between Ca\(^{2+}\) and phosphate units ([PO\(_4\)\(^3-\)]\(^-\)) of polyP [12] was chosen to ensure that the polyanionic polyP chains in the NP are surrounded by a net-positive Ca\(^{2+}\) surface. Such particles have been shown to be more readily taken up by human Caco-2 cells than those with a negative surface charge [15].

PolyP is a distinguished polymer that underscores the physiological relevance of a controlled transition between the NP state and the coacervate phase. This polymer is accumulated in the acidocalcisomes with an acidic milieu [16]. At the pH ~5 present there, phosphoric acid occur as H\(_2\)PO\(_4\), which in turn forms a Ca\(^{2+}\) salt Ca(H\(_2\)PO\(_4\))\(_2\), which is readily soluble in water (18 g L\(^{-1}\)). At a pH around 7 (near the pK\(_{a2}\) of 7.09), ~50% of the phosphoric acid is present as H\(_2\)PO\(_4\) and ~50% as HPO\(_4^{2-}\), respectively. With increasing pH, the solubility of the Ca\(^{2+}\)-phosphate salts decreases rapidly and an almost insoluble salt Ca\(_6\)(HPO\(_4\))\(_2\) (0.02 g L\(^{-1}\)) is formed [17]. In the acidocalcisomes, where polyP is stored, the phosphorus content (141 μg [in ~200 insect eggs]) far exceeds the Ca\(^{2+}\) content (18 μg), suggesting that in this acidic environment the phosphate deposits largely exist as polyP/Ca\(^{2+}\)-NP [18]. From the acidocalcisomes of the blood platelets, polyP is released into the extracellular space. The phosphate concentration there (1 × 10\(^{-3}\) M) is comparatively low compared to the intracellular concentration of 200 × 10\(^{-3}\) M [19]. Even the comparatively high intracellular Ca\(^{2+}\) level of 1 mM [20] does not favor the precipitation of the calcium salts of the monomeric phosphate under the existing pH conditions. Later under “Results” it is shown that at a pH ~7 aggregates of the calcium salts of polymeric phosphate (Ca-polyP) are formed that are in an aqueous coacervate phase, or at pH ~10 in the particulate NP state.

In general, it is obvious that the degree of solubility of Ca-phosphate/polyP salts/NP determines both the direction and the intensity of the downstream reactions. Therefore, to get an evidence-based view of the functional potential of Ca-polyP-NP in silico, modeling and simulation studies were performed with Ca\(^{2+}\) and polyP in an aqueous milieu. Based on our data reported here, a fairly straightforward vision could be derived revealing a pH-dependent sorting and clustering of Ca\(^{2+}\) in the polyP environment. Furthermore, elucidating the pH dependence of the NP/coacervate mechanism also provided additional understanding of the accessibility of this inorganic polymer in a biological environment. In the coacervate phase, the release of polyP is faster and more extensive compared to the kinetics of polymer release from the NP phase.

In the final series of experiments, it is documented that cells cultivated on the coacervate phase show a more intense proliferation and cell layer formation. As a functional consequence, the cells growing on the coacervate release considerably more ATP into the extracellular matrix (ECM) than cells cultured on particulate polyP (Ca-polyP-NP). The property of polyP to function as a storage form for metabolic energy is dependent on the enzymatic cleavage of polyP via the enzyme alkaline phosphatase (ALP) and the subsequent phosphotransferase enzyme, adenylyl kinase (ADK), which catalyzes the reversible interconversion of the three adenosine phosphates (ATP, ADP, and AMP) [21].

The presented coacervate, a Newtonian fluid, with its biocompatible viscosity offers human epithelial A549 cells a suitable environment to spread and to develop their functional potential. These cells have been found to release ATP in a Ca\(^{2+}\)-dependent manner [22]. The data shown here also underscore the importance and necessity of the coacervate phase of Ca-polyP as a transition state of the biologically inert, stable storage form of polyP, the Ca-polyP nanoparticles, which enables the polymer to exhibit its energy-rich and growth-stimulatory properties.

2. Materials and Methods

2.1. Materials

Na-polyphosphate (Na-polyP) with an average chain length of 40 orthophosphate (P\(_i\)) units was purchased from Chemische Fabrik Budenheim (Budenheim; Germany).

2.2. Preparation of Ca-polyP-coacervate

The coacervate of Na-polyP was obtained by mixing an aqueous solution of Na-polyP (100 mL of a 0.48 M solution; based on P\(_i\)) with CaCl\(_2\)-\(\cdot\)2H\(_2\)O (C3306, Sigma; 100 mL of a 1.2 M solution). Unless otherwise noted, the CaCl\(_2\) solution was added to the polyP solution under vigorous mixing, using a magnetic stirrer at a rate of 5 mL min\(^{-1}\). The pH was kept constant at pH 7; the duration of the preparation was 30 min. The resulting coacervate was washed several times with cold distilled water and stored at 4 °C. "Ca-polyP-Coa". Where indicated, the sequence of mixing was reversed: the polyP solution was added to the CaCl\(_2\) solution.

The viscosity of the coacervate was determined using a Brookfield DV3T Rheometer (Brookfield, Middleboro; MA).

2.3. Preparation of Ca-polyP-NP

Amorphous nanoparticles of Ca-polyP, “Ca-polyP-NP”, were prepared as described [23]. If not mentioned otherwise, a 2.5-fold molar excess of CaCl\(_2\)-\(\cdot\)2H\(_2\)O over the concentration of Na-polyP was used to prepare the NP. The solution was kept constant at a pH of 10 with NaOH. The reaction was continued for 12 h under stirring conditions. The obtained NP were washed in water and in ethanol and then dried (50 °C).

In one series of experiments, the polyP solution was poured into the CaCl\(_2\).

2.4. Microscopic analysis

High resolution SEM (scanning electron microscopy) images were taken with a Zeiss Gemini 1530 (Zeiss Oberkochem; Germany). The samples were dried prior to mounting onto a microscope aluminum stub. The light microscopical images were taken with a VHX-600 Digital Microscope (Keyence, Neu-Iserbarg, Germany). The TEM (transmission electron microscopy) analysis was performed with a Tecnai 12 microscope (FEI Electron Optics, Eindhoven; Netherlands) as described [24].

2.5. Fourier Transformed Infrared Spectroscopy

The material, dried NP or coacervate, was ground to a composite
powder and then analyzed by FTIR (Fourier Transformed Infrared Spectroscopy). The analyses were performed with an attenuated total reflectance-FTIR spectroscope/Varian IR spectrometer (Agilent, Santa Clara, CA).

2.6. Simulation method

The molecular mechanics and molecular dynamics simulations and predictions of interatomic forces between Ca$^{2+}$ and polyP were performed using the Materials Studio-Discovery Studio packages and the Analysis Module in InsightII/Discover program (BIOVIA-Accelrys Inc., San Diego, CA, USA). The protocols are based on the data collected with the COMPASS force field algorithm [25,26]. After drawing the polyP chain with a length of 20 P$_1$ units and subsequent implementation of the additional option “clean geometry” of the program, 16 polyP molecules together with 350 Ca$^{2+}$ ions and an aqueous environment were computed, again in the “clean geometry” mode 1000-times until the data output reached a variation of <1%. Energy minimizing was performed in the presence of ~5000 water molecules at a density of 1.15 g cm$^{-3}$. The simulations were run at a temperature of 297 K.

2.7. PolyP release assay

The following samples were used. The respective polyP polymer either as free Na-polyP or in combination with particulate Ca-polyP-NP was embedded in a 2% aqueous hydrogel prepared from hydroxyethyl cellulose (HEC; Caelo, Hilden; Germany). After a heating step (121 °C under positive pressure), the HEC hydrogel was supplemented with 600 μg mL$^{-1}$ of Na-polyP or 60 μg mL$^{-1}$ of Ca-polyP-NP or the two components together (600 μg mL$^{-1}$ of Na-polyP and 60 μg mL$^{-1}$ of Ca-polyP-NP), as indicated under “Results”. Samples (1 mL) were dispensed into 24-well plates and incubated with 2 mL Ham’s medium/serum (FBS). Immediately after plating or after different time periods aliquots were taken and subjected to the enzymatic polyP determination.

The release of polyP from the polyP-hydrogel was quantitated using the “Phosfinity” total polyphosphate quantification kit (Aminovirne B.V., 6361HK Nuth; Netherlands). This assay is based on the enzymatic degradation of polyP with a highly specific enzyme releasing orthophosphate, which is quantified using a colorimetric phosphate detection reaction. The absorbance is read at 882 nm in a spectrophotometric plate reader. First, the concentration of orthophosphate in the reaction mixture was determined at the beginning of the enzymatic reaction. The concentration values obtained were subtracted from the values in the assays at the end of the enzymatic reaction. The polyP material detected by the “Phosfinity” assay thus only quantifies polymeric polyP. A polymer-specific enzyme is used to quantify the polyP. The amounts of polyP (with all chain lengths) were calculated based on a calibration curve using standard phosphates between 2 and 200 μM.

2.8. Preparation of the PVA matrices

A 4% PVA solution (polyvinyl alcohol); $M_w$ 146,000–186,000; #363065 Sigma-Aldrich) was used to plate the A549 cells as described before [27]. The polymer was supplemented with 200 μg mL$^{-1}$ of “Ca-polyP-NP” and 1 mL of the PVA hydrogel was pipetted into 24-well plates; “PVA/polyP:NP”. After two washing cycles, the layer-covered wells were used to overlay A549 cells; 2 mL cell aliquots (20 0.103 cells mL$^{-1}$). To process the polyP phase of the “Ca-polyP-NP” caged in PVA to the coacervate state, the “PVA/polyP:NP”-coated 24-well plates were incubated for 24 h with 2 mL of Ham’s F-12K medium with 10% FBS. During this treatment, the nanoparticles in the wells were converted into the “PVA/polyP:Coa” coacervate layers.

2.9. Cultivation of A549 cells

To determine the ATP release, A549 cells (#86012804 Sigma, Taufkirchen, Germany), a human lung (carcinoma) cell line, were cultured in Ham’s F-12K (Kaighn’s) medium (#21127022; Gibco/Thermo Fisher Scientific, Dreieich; Germany) supplemented with 10% FBS, 1% penicillin-streptomycin, and 4 mM glutamine as described [28,29]. The cells were grown in 24-well plates in a humidified atmosphere of 5% CO$_2$ in air (37 °C).

Spreading experiments of A549 cells were performed in 24-well plates coated with 1 mL of the respective PVA matrix. The cells at a density of 20 0.103 cells mL$^{-1}$ were suspended on the matrices in 100 μL. After incubation for 48 h, the cells were fixed (3.7% formaldehyde; 20 min), permeabilized, and subsequently immune-stained for the distribution of the cell surface associated mucin 1 with anti-mucin 1 (#PAS-61524; Thermo Fisher Scientific; dilution 1:1000) and then reacted with Alexa Fluor 488-conjugated goat anti-rabbit IgG secondary antibody (failing in green) as described [29]. The nuclei were stained red with propidium iodide (#P4864; Sigma-Aldrich). The specimens were inspected with the fluorescence microscope (Olympus, Hamburg; Germany).

2.10. Uptake studies of the NP

The A549 cells at a density of 100 0.103 cells mL$^{-1}$ were suspended on the “PVA/polyP:NP” layer and incubated for 3 h in medium/serum. Then the cells were removed from the substrate by gentle shaking and transferred to a non-supplemented PVA layer. After additional 3 h, the cells were fixed, embedded (LR-white resin [# 62661; Sigma-Aldrich]), sliced and contrasted [24], and then inspected by TEM.

2.11. Determination of the extracellular ATP level on polyP containing matrices

The ATP released to the extracellular space was quantified using the luciferin-lucerase-based Enlighten assay (Promega, Madison; WI, USA) as described [24]. The A549 cells were grown in 24-well plates until they reached 90–100% confluency. Then the cells were suspended in Ham’s medium without serum and incubated for a further 2 h at a cell density of 100 0.103 cells mL$^{-1}$. The wells were coated with three different PVA matrices: PVA control, “PVA/polyP:NP”, and “PVA/polyP:Coa”. To stop the reaction, 0.5 mL aliquots were collected and transferred to chilled polypropylene tubes (#Z334006; Sigma) and centrifuged (12,000 g; 5 min; EBA 200; Hettich GmbH, Tuttingen; Germany). Aliquots of 100 μL were taken from the supernatants and transferred to the luciferin-lucerase assay. The values obtained were correlated with a standard curve for ATP to obtain the level of ATP released from the cells. ATP concentrations are given as pmol 0.106 cells$^{-1}$. As additional controls, the cells were pre-incubated with 40 μM ApA ($P^1P^2P^3P^4$adenosine-5’-pentaphosphate pentasodium salt; #D4022, Sigma), an inhibitor of ADK [30], or 1 mM LEV (levamisole hydrochloride; #L9756; Sigma), which inhibits ALP [31], for 10 min before cell seeding.

2.12. Staining of A549 cells for ALP

The cells were cultured on either a plain PVA surface or a polyP-supplemented PVA hydrogel (either “PVA/polyP:NP” or “PVA/polyP:Coa”) at a cell density of 10 0.103 cells mL$^{-1}$ and incubated in medium/serum for two days. When mentioned, the cultures were supplemented with 1 mM LEV. Then samples were taken, washed with phosphate-buffered saline containing 0.1% Tween-20 and fixed with 70% (v/v) ethanol. After a further washing cycle with PBS, the cells were reacted with the ALP staining solution (#ab242287, ALP staining kit; Abcam, Cambridge; UK) and washed again with PBS and inspected by light microscopy.
2.13. Statistical analysis

The analyses were carried out with the SigmaPlot 13.0 (Systat Software, Erkrath; Germany) and the resulting graphs were plotted with the help of GraphPad Prism 7.02 (GraphPad Software, San Diego; CA). At least five or ten (as indicated) replicates were performed for each series. The results are given as mean ± standard deviation. The paired t-test was applied to assess the significance of differences. The p-values < 0.01 are considered as statistically significant (*). When comparing members from different groups, these significant differences were also marked (#).

3. Results

3.1. Driving forces of “Ca-polyP-Coa” and “Ca-polyP-NP” formation

Both the polyP coacervate and the polyP NP form after the interaction between soluble anionic polyP (Na-polyP) and cationic Ca\(^{2+}\) ions (CaCl\(_2\)) in an aqueous environment. Broadly, the coacervate is formed at neutral pH, while NP form at pH ~10. During this process electrostatic attractions between the oppositely charged ions take place. However, since ion-dipole attraction forces must also be taken into account in ionic solutions, it is important to estimate not only the strength of the interionic bonds between the ionic compounds but also the strength of the ion-dipole forces [32]. To approach a solution, docking and molecular dynamics simulations were performed as described under “Materials and Methods”. Using these algorithms, topographic interactions of the three partners (polyP [16 polyP molecules], Ca\(^{2+}\) [120 Ca\(^{2+}\) ions] and water [~5000 water molecules]) were predicted.

At pH 7, the three components form an almost random arrangement pattern right from the start. The Ca\(^{2+}\) ions are arranged in a mixing mode around the polyP molecules throughout the aggregate (Fig. 1 A and B). Likewise, the water molecules are evenly distributed in the assemblies. After masking the water, the random, stochastic organization of polyP and Ca\(^{2+}\) is evident (Fig. 1 B). After hiding the polyP, the arrangement of the water molecules in the center of the aggregates can be seen (Fig. 1 C).

The simulation pattern of polyP, Ca\(^{2+}\) and water at pH 10 shows a different pattern (Fig. 1 D–F). A clustering of the polyP molecules and partial separation of the Ca\(^{2+}\) ions is seen. It is obvious that under the setting input into the algorithm of the modeling system, the cationic Ca\(^{2+}\) becomes at least partially separated from the anionic polyP aggregations, indicating a shift between short- and long-range electrostatic interactions. Blinding out the polyP molecules highlights the presence of water within the polyP cluster (Fig. 1 F).

3.2. Kinetics of “Ca-polyP-Coa” and “Ca-polyP-NP” formation

Compared to the NP formation, the process of “Ca-polyP-Coa” formation (at pH 7) is a slow process. Starting with a solution of 0.48 M Na-polyP into which 1.2 M CaCl\(_2\) is introduced, the first increase in turbidity is seen after ~9.5 mL of CaCl\(_2\) (Fig. 2 I-A). Later in the course of aggregate formation (as mentioned later, coacervates are formed at pH 7), the turbidity decreases again and levels off at 20% after ~25 mL. The resulting aqueous aggregate has a viscosity of 10–15 cP. The kinetics of coacervate formation, measured by dynamic viscosity, is independent within the temperature range 14 °C and 55 °C.

Very much faster is the aggregate formation at pH 10, which reflects the formation of the nanoparticles “Ca-polyP-NP” (Fig. 2 II-B). Under these higher pH conditions, the turbidity increases almost immediately. The high turbidity remains constant during the subsequent preparation period.

By eye inspection, the difference in the kinetics of aggregate formation at pH 7 and pH 10 becomes obvious (Fig. 2 II-A and B). At pH 7, the first aggregate can be seen after addition of 6 mL of CaCl\(_2\) and the fluffy, bulky coacervate deposits appear after ~10 mL (Fig. 2 II-A). Again in contrast to the formation at pH 7, the first solid flocs appear already after 0.5–1.5 mL of CaCl\(_2\) (Fig. 2 II-B) and the turbid appearance forms after the addition of ~10 mL. Even after mixing is stopped the material does not sediment during a 30-min incubation period.

Fig. 1. PolyP (chain length of 20 P units) interacting with Ca\(^{2+}\) and water at (A–C) pH 7 and (D–F) pH 10. At neutral pH, the three components of the aggregates, polyP, Ca\(^{2+}\) and water, are arranged stochastically, while at pH 10 a clustering of the polyP molecules and a distinct separation of the Ca\(^{2+}\) ions are found.
Following the initial procedure [23], adding CaCl2 (in a
2.5-fold molar excess) to the polyP solution, the dense aqueous coacervate forms at a pH of 7 (Fig. 2 III-A to C). The appearance of the dense/aqueous electrostatic complex after drying the sample on the stub is a continuous, slightly granular layer. The diameter of the granules is ~200
nm (Fig. 2 III-B). They are porous with holes of ~10 nm (Fig. 2 III-C).

If – under the same concentration conditions – the reaction is run at pH 10, only nanoparticles with a size range of 80–120 nm appear (Fig. 2 III-D to F).

FTIR: Differential infrared spectroscopy was performed on the material obtained by pouring the CaCl2 solution into the polyP solution. The spectrum of the material at pH 7 (coacervate) differs from that collected at pH 10 (NP). The characteristic signatures for polyP are the signals at characteristic wavenumbers given here shift from those of Na-polyP. The change of the turbidity was followed by measuring the changes in the absorbance at 700 nm. (Fig. 2III-B). This change is most likely due to the binding of Ca2+ to the internal phosphate units.

Indicative is the change of the νas(PO2)3 2+ signal if the CaCl2 solution is not dropped into the polyP solution, but Na-polyP is added to the CaCl2 solution in the reverse order (Fig. 2 III-C). Under this condition, the νas(PO2)3 2+ signal is seen around ~1250 cm
–1 not only at pH 7 but also at a pH 10. From this finding it was deduced that the νas(PO2)3 2+ units undergo immediate binding to the fully available Ca2+ ions. The morphology - sequence dependence (CaCl2 – Na-polyP): The morphology of the Ca2+: polyP salts is dependent on the sequence in which the two starting solutions are mixed together. This fact becomes visible by SEM analysis. According to the initially described sequence of adding CaCl2 solution to Na-polyP solution [23], the coacervate is formed at pH 7, while nanoparticles develop at pH 10 (Fig. 3 I-A versus –C). If Na-polyP is added to the CaCl2 solution at pH 7, Ca2+/phosphate deposits are formed, which are identified as coacervates by SEM (Fig. 3 II-B). Surprisingly, bulky coacervate depositions are also formed at pH 10, as identified by SEM (Fig. 3 II-D) and confirmed by FTIR (Fig. 3 I-C). Hence, under these pH 10 conditions and when 100 mL of a 0.48 M Na-polyP solution is added to 100 mL of a 1.2 M CaCl2 solution, coacervate depositions are formed over the complete reaction process. The additional SEM image in Fig. 3 II-E (pipetting scheme: 100 mL of 1.2 M CaCl2 added to 100 mL of a 0.48 M Na-polyP solution) shows the pronounced porous morphology in the developed NP.

The coacervate-like polyP phase is probably caused by a strong and rapid binding of the Ca2+ ions to the polyP polyanions as soon as the Na-polyP solution is added to the CaCl2 solution. In turn, no sorting out of Ca2+ from polyP ions takes place, which has been predicted for the interaction of the two ions during the addition of CaCl2 to Na-polyP (Fig. 1 E). This assumption is also consistent with the published data [37].
Interatomic forces predictions: The calculation of the interacting forces of Ca\(^{2+}\) and polyP revealed a relocation movement of the two partners as a function of time (Fig. 3 III). At pH 7, the two components Ca\(^{2+}\) and polyP (Fig. 3 III-A) organize themselves rather randomly, while in the following steps the two interacting members sort apart and finally form two concentric rings with the Ca\(^{2+}\) on the outside and polyP as the core (Fig. 3 III-B to D). It can be predicted that these assemblies form the liquid droplets (Fig. 3 III-E) with the polyP inside (core) and the Ca\(^{2+}\) ions around it [38]. The ions are surrounded with hydrate shells [39,40], allowing the droplets to merge and form the water-rich coacervate. At pH 10, the components Ca\(^{2+}\) and Na-polyP also form a concentric ring pattern, but this is more dense (Fig. 3 III-F). This dynamic is based on the stronger affinity of Ca\(^{2+}\) for the more ionized polyP, especially since at pH 10 the second OH of the phosphate end-groups of the polymer (pKa-2 of 7.09) is also almost completely dissociated. Consequently, the liquid droplets formed at pH 7 are larger than the particles at pH 10 (Fig. 3 III-E).
versus Fig. 3 III-F to H). As the “Ca-polyP-NP” nanoparticles are dried, the water in the pores is removed.

3.4. Differential polyP release kinetics from nanoparticles versus coacervate

The release of polyP was determined from a HEC hydrogel embedded either with Na-polyP alone (600 µg mL⁻¹) or in combination with “Ca-polyP-NP” (60 µg mL⁻¹). These concentrations and concentration ratios were chosen because the optimal effect on wound healing could previously be achieved with them. When the release of polyP from the hydrogel was determined with Na-polyP alone (600 µg mL⁻¹), ~40% of the polymer was released during the first 2 days. An additional ~20% was released at day 4 (Fig. 4 I-A). After 9 days, 72% (434 µg mL⁻¹) of the Na-polyP polymer was found in the medium/serum surrounding the sample. The kinetics show that most of the polymer is set free during the first 4 days. The comparison of the determined released polyP from the hydrogel supplemented with Na-polyP (600 µg mL⁻¹) alone with that together with “Ca-polyP-NP” (60 µg mL⁻¹) showed values that do not differ significantly (Fig. 4 I-A). In principle, this result was to be expected, since Na-polyP was present in the hydrogel in a 10-fold excess over “Ca-polyP-NP”.

Therefore, the release kinetics from the hydrogel was determined with “Ca-polyP-NP” as the sole component. In this series of experiments, a clear difference in the kinetics was evident compared to the results obtained with the gel containing both Na-polyP nanoparticles (Fig. 4 I-B). During the first 4 days of incubation, only a non-significant increase in polyP was measured in the assays with “Ca-polyP-NP” alone. The increase becomes significant after an incubation period of 6 days with ~37%. Extending the incubation period to 9 days resulted in a release of 69% (41 µg mL⁻¹) of the polymer from the particulate “Ca-polyP-NP”. From the data in Fig. 4 I, it is concluded that the release of polyP from the gel containing the Na-polyP is almost twice as fast compared to the hydrogel supplemented with particulate polyP.

3.5. Transformation of nanoparticles into the coacervate phase

In an earlier report, we showed that the high zeta potential of the nanoparticles is reduced in the presence of peptides, leading to the transformation of the nanoparticles into the coacervate phase [12]. This process is now supported by FTIR and also SEM analyses.

As shown above (Fig. 3 I-B), the νas(P=O) signal in the FTIR spectrum of the Ca-polyP coacervate, which is present as a distinct peak at a wavenumber of 1255 cm⁻¹, changes into a flat shoulder in “Ca-polyP-NP”. This signal is also not present in the polyP-free PVA matrix (Fig. 4 II); again, this νas(P=O) band, which is ascribed to the stretching vibrations of (PO2) ν, is absent in the PVA layer with “Ca-polyP-NP”, the “PVA/polyP-NP” hydrogel. However, when this layer is incubated for 24 h with Ham’s medium/FBS, the νas(P=O) signal at 1256 cm⁻¹ can be clearly seen, indicating that the nanoparticles in the PVA hydrogel undergo a conversion to the coacervate phase, “PVA/polyP:Coa” (Fig. 4 IV B).

This conversion is also seen in microscopic images (Fig. 4 III). The surfaces of the PVA hydrogel are flat (Fig. 4 III A) in contrast to the matrix supplemented with “Ca-polyP-NP” (Fig. 4 III B). After incubation with medium/serum, polyP undergoes a transformation into a flat coacervate gel (Fig. 4 III-C and D).

3.6. Increased spreading of A549 cells on polyP-enriched PVA matrices

The A549 cells were plated in 24-well plates coated either with the nanoparticles “PVA/polyP-NP” or with the coacervate obtained from the NP phase by incubation with medium/serum “PVA/polyP:Coa” (Fig. 5 I). The cells attached to the particulate polyP phase have a large size (~50 µm) and show a scattered distribution pattern and the nuclei are flat (Fig. 5 I A and C). The cytoplasm was stained with mucin 1 antibodies and the nuclei with propidium iodide. The increase in the growth number during the 24 h incubation period was determined to be 47.1 ± 7.3%. In contrast, the cells grown on the coacervate phase of polyP are densely packed and their nuclei are arranged in condensed, massy cell bodies.
Fig. 5. Increased biological activity of A549 cells after cultivation on PVA polyP coacervate, “PVA/polyP:Coa”. (I) Different cell morphology of A549 cells cultured for 48 h on (A and C) particulate “Ca-polyP-NP” (“PVA/polyP:NP”) versus those contacted with (B and D) “PVA/polyP:Coa”. On the coacervate layer, the density of the cells is higher and the cell bodies are more compact compared to the particulate polyP in PVA. (II) Response of A549 cells to different polyP environments. The cells were suspended on PVA layers, either on PVA control (black bars), particulate “PVA/polyP:NP” (red bars) or on “PVA/polyP:Coa” (blue bars) and incubated for 1 h or 2 h. Then the supernatants were collected and the ATP content determined using the luciferin-luciferase-based Enlighten assay. Significant pairs are marked (* p < 0.01). In parallel series, the effect of the ADK inhibitor Ap5A (hatched) and the ALP inhibitor LEV (cross-hatched) on the ATP production of A549 cells cultured onto the three different matrices was determined as well. Significant differences are marked (p < 0.01).

Fig. 6. NP in the cytoplasm of A549 cells and distribution of ALP in A549 cells, the enzyme that hydrolyzes the polyP in NP. (I) Uptake kinetics of polyP NP by A549 cells; TEM. (A) At time zero, no nanoparticles (NP) can be seen. (B and C) After incubation onto the “PVA/polyP:NP” layer for 2 h or 3 h, (D) the cells were transferred on a non-supplemented PVA substratum and incubated for a further 3 h. On the “PVA/polyP:Coa” layer, the cells take up NP. The NP are absent on the plain PVA surface; the darker shaded areas could represent coacervate areas (Coa?). (II) Increased expression of ALP in A549 cells cultured on “PVA/polyP:Coa”; light microscopy. (A) The cells growing on the hydrogel without polyP (“PVA”) for 2 d show a low staining reaction and only a scattered distribution pattern compared to those cultured on (B) “PVA/polyP:NP”. (C) In cultures on “PVA/polyP:Coa”, however, the staining is significantly stronger and the density of the cells is higher. (D) Suppression of the staining reaction after addition of the ALP inhibitor LEV to cultures on “PVA/polyP:Coa”.

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(Fig. 5 I-B and D). During the 24 h incubation, the cell number increased by 73.0 ± 5.6%.

3.7. ATP release from A549 cells on coacervate versus nanoparticles

The A549 cells react differently when cultivated on either the PVA-control matrix, the PVA-based matrix supplemented with “Ca-polyP-NP” (200 μg mL⁻¹), “PVA/polyP:NP”, or on the coacervate phase of polyP. The A549 cells grown on the “PVA/polyP:NP” responded after a 2-h incubation period with a small but significant release of ATP compared to PVA without polyP. In contrast, the cells attached to “PVA/polyP:Coa” layers produce a strong ATP burst already after 1 h, which is significant and ~150% higher compared to the PVA control and the “PVA/polyP:NP”-supplemented layer (Fig. 5 II). After a further 1 h, the ATP release on the coacervate layer increases by about two-fold.

The ATP production by the A549 cells is significantly inhibited by the ADK inhibitor Ap5A and also by the ALP inhibitor LEV with all different PVA matrices (Fig. 5 II).

3.8. Uptake of NP into A549 cells

The cells were first incubated in medium/serum on a “PVA/polyP:NP” layer. Samples were taken at time 0 and after incubation for 2 and 3 h. Then the cells were removed from the polyP-supplemented matrix and transferred to a non-supplemented PVA layer and the incubation was continued for 3 h. Slices from embedded cells were inspected by TEM (Fig. 6 I). At time 0, no NP are visible in the cut cells (Fig. 6 I). After a 2–3 h, NP accumulate in the cytoplasm (Fig. 6 I-A). After a further 3 h, no distinct, electron-dense particles can be seen in the cytoplasm; only clusters of diffuse darker shaded areas are visible, presumably reminiscences of coacervate droplets.

Fig. 7. Site of polyP formation and its distribution pattern in humans. Left panel: Every cell contains polyP with the highest concentration in the large bone marrow cells, the megakaryocytes (size between 50 and 150 μm). From these precursor cells 2–3 μm small platelets are formed and released into the blood-stream; one megakaryocyte produce 1000 to 3000 platelets. In these blood cells the chain-shaped polyP undergo to Ca⁺⁺ the transition to Ca-polyP-NP of a 100 nm size. Right panels: The phases of polyP processing: polyP chain to nanoparticle, and finally and in particular in the extracellular matrix system (ECM) to coacervate. Bottom: Always, when a damaged organ is under regeneration, blood platelets deliver the polyP components, like in bone, cartilage, viral tissue damage (e.g. Covid) or wounds, which contribute to the restoration.

3.9. Effect of polyP on the expression of ALP

To assess the effect of the different polyP phases on ALP expression, A549 cells were seeded on following surfaces: plain PVA, “PVA/polyP-NP” or “PVA/polyP:Coa”. After an incubation period of 2 d, the cells were subjected to a specific ALP staining. In the case of the PVA cultures, only a very low cell density is visible on the PVA matrix (Fig. 6 II-A). The degree of staining of cells cultivated on “PVA/polyP-NP” is higher (Fig. 6 II-B). However, an incomparably high staining level is seen for those cells grown on “PVA/polyP:Coa” (Fig. 6 II-C); when these cultures with “PVA/polyP:Coa” were additionally exposed to 1 mM LEV, the red/purple staining of the cells is almost completely reduced (Fig. 6 II-D).

4. Discussion

First of all, inorganic NP made from native or processed metals are stable in a biological environment, possibly also inert, such as gold, silver, copper, and aluminum NP. This property stems from their inherent antibacterial activity and/or in vitro cytotoxicity. They obtain their biomedical impact due to their antibacterial function [41]. Among the inorganic nonmetallic materials constituting NP, those based on phosphate or silica gained bio-medical impact [42–45] because they are soluble (can form a dispersion) under physiological conditions. A new area started when it was possible to prepare polyP-NP that elicit regenerative activity [46] in a bioinspired way [23]. The special feature of these polyP-NP is that they undergo a transition from an “inert” and consistently stable NP state into a physiologically active coacervate phase [12]. In the extracellular space, both unprocessed polyP and Ca-polyP-NP are transported via the blood platelets through the blood vessels [6,7,47]; Fig. 7. Both short-chain polyP (~50 P₁ units) and longer-chain polyP (~150 P₁ units) are transported via these cell fragments and released from them upon activation. In the circulating blood, the polyP...
incorporated into NP remains attached to the platelet surface while the soluble polymer chains are set free. When the platelets are exposed to ECM, they are activated by interaction with collagen and laminin, and also with fibrinogen in vascular events [48]. The activation of the platelets upon contact with ECM proteins occurs via their adhesion receptors, especially when the endothelial barrier is disrupted [49]. PolyP in the circulating blood or plasma has a half-life of ~1.5–2 h [50], compared to platelets, which persist for 7–10 d and constitute a density of 150–400 · 10^9 platelets per mL. In addition to the platelets, macrophages contain large amounts of polyP [51]. These migration competent cells, which are important for the inflammatory status, produce migrasomes, which are loaded onto retraction fibers that trail behind these migrating cells, the podocytes [52]. The migrasomes are vesicular structures filled with cellular contents and small vesicles. Platelets also produce migrasomes, which they release after activation [53]. It is to be expected that Ca-polyP-NP are also involved in this recently published cell migration pathway.

PolyP is an essential buffering molecule in eukaryotic cells [54]. More specifically, it has been reported that polyP is a crucial molecule especially in the mitochondria, where this polymer controls the Ca^{2+} exchange across the mitochondrial inner membrane and calcium-buffering system within the matrix [55].

PolyP-NP undergo coacervation in the presence of peptides especially in the extracellular space. This process is supported by the aqueous environment of the ECM which is also rich in Ca^{2+} [12]; Fig. 7. Through the blood vessels and the ECM, polyP is distributed to the different organs, bone, cartilage and to wounds and, if present, also to viruses such as the coronavirus 2 [56,57]; Fig. 7. In the liquid gel-like phases (intra- and extracellular sites), polyP undergoes enzymatic hydrolysis in the presence of ALP, a process that, together with ADR, initiates ATP generation [21]. Both enzymes occur in the intra- and extracellular space as well as cell surface-associated [12,58]. Furthermore, Ca-polyP-NP are taken up intracellularly via an endocytosis mechanism [59]. There, the NP remain closely associated with the cell and form sites where intracellular coacervation also takes place [60].

The aim of the present study was to elucidate and model the initial events of the coacervation process of polyP, its pH dependence and – importantly – its phase-dependent effect on the physiological generation of extracellular ATP [21]. Again, as defined, coacervates are formed as condensed liquid-like droplets between oppositely charged polymers. However, the polyP coacervates form in the presence of polyP together with divalent, “monomeric” cations. A series of coacervation studies of Na-polyP with divalent (e.g., Ca^{2+} and Mg^{2+}) but also trivalent (e.g., Al^{3+}) cations have been performed and published [61,62], mainly focusing on material science. Here we focus on a biomimetically engineered NP fabrication process that is also used in nature to deposit polyP in cell organelles, specifically in the acidocalcisomes [16].

High concentrations of polyP with a chain length between 40 and 150 Pi units are present in the acidocalcisomes (reviewed in: [21]). In the protozoan Trypanosoma cruzi, these organelles contain not only polyP (~1400 mmol kg^-1) but also high levels of Mg^{2+} (650), Ca^{2+} (170), Zn^{2+} (150), and Na^{+} (160) [63]. Consequently, the polymer is present in these organelles as free chains (Na-polyP) or particulate Ca-polyP-NP. The fabricated Ca-polyP-NP [23] or Mg-polyP-NP [64] have a size of around 100 nm, are porous [46], and have a high zeta potential with the consequence that the particles have a high stability [12]. Despite this physical property that causes repulsive electrostatic forces of the particles resisting aggregation, the Ca-polyP-NP are distinctly biocompatible and evoke a pronounced regenerative activity in vivo [65]. When applied to bone defects, the particles are resorbed and induce new hydroxyapatite mineral deposition at this site. The unprecedented feature of these polyP-NP is their characteristic phase transition from the NP state into an aqueous coacervate in the physiological environment. This process is driven by peptides in the body fluids that lower the zeta potential and cause the transition into the coacervate state [12]. In this conversion, the size of the polyP chains (~35 Pi units) and the Ca:P atomic ratio 1.78 remain unchanged (starting from a NP preparation at a CaCl2 to Na-polyP weight ratio >2.5) [23].

So far, there are no data that indicate differences between the physiologically occurring polyP fractions, the medium-chain polyP molecules (around 40 Pi units) compared to the longer-chain polyP molecules (~400 Pi units). Since the water solubility of both polymer fractions is not dramatically different at room temperature, there is presently no reason to assume that these polyP fractions differ to a greater extent in their tendency to form coacervates.

Under conventional conditions, adding the CaCl2 solution to the polyP solution, the coacervate is obtained at pH of 7 while the NP are formed at pH 10. Applying the simulation approach using molecular mechanics and molecular dynamics algorithms, a drastic difference in the distribution patterns of the polyP chains and the Ca^{2+} ions at pH 7 and pH 10 is found. At pH 7, the condition under which the coacervate is formed, there is a random distribution of the polymers and Ca^{2+} (Fig. 1), allowing liquid droplets to develop along the Ca^{2+} shell surrounding the polyP core (Fig. 3 III). In contrast, at pH 10, the condition favoring NP formation, the Ca-polyP-NP are formed, there is a random distribution of the polyP and Ca^{2+} molecules, the NP remain unchanged (starting from a NP preparation at a CaCl2 to Na-polyP weight ratio >2.5) [23].

The remarkable biochemical feature of polyP is the presence of the high-energy phosphoanhydride bonds that link the Pi units within the polymer chain [67–69]. During enzymatic hydrolysis by ALP, starting from the termini of the polyP chain, the released free energy is partially stored in ADP, which is used for driving next steps such as ATP synthesis (reviewed in: [21]). Subsequently, the phosphotransferase ADK catalyzes the interconversion of ADP to ATP and AMP. The generation of ATP from polyP is a fast process and takes place within 1–2 h [70]. In the present study, A549 cells were plated on a PVA hydrogel supplemented without or with polyP. After an incubation period of 48 h, the cells seeded on PVA with particulate Ca-polyP-NP show a lower cell density compared to cells on the polyP coacervate “Ca-polyP-Coa”. Using this setting, the ATP generation and release from the cells on PVA without polyP was
significantly lower compared to the cells growing on particulate or even more on coacervated polyP. The key roles of the two enzymes ALP and ADK have previously been proven by inhibition studies with their inhibitors LEV and Ap5A [71]. These results underscore the importance of the coacervate phase in achieving the optimal utilization of polyP. Furthermore, after exposure to the polyP coacervate, A549 cells express a higher level of ALP, a finding that is strengthened by a higher growth rate of the cells.

The cells growing on the PVA hydrogel were also used to gain insight in the fate of the polyP-NP after endocytotic uptake, which is inhibitable by trifluoperazine [59]. For this series, the cells were grown on the “PVA/polyP:NP” layer. After 2–3 h, the NP can be clearly seen in the cytoplasm by TEM visualization. This duration for particle uptake by endocytosis is within the range published for other systems [72]. After transfer to a polyP-lacking matrix, the NP disappear in the cytoplasm, leaving – apparently – space for coacervate droplets.

By definition, as mentioned, a coacervate, especially a coacervate of the complex type, is built from colloidal droplets formed by liquid–liquid phase separation in a solution containing both polycations and polyanions [73]. In contrast, proteinaceous gelatin-alginate coacervates are formed according to an “egg-box” model in the presence of Ca2+ [74]. In this design pattern, it is proposed that in a statistical ion-binding process, two polymers are linked in a cooperative manner via Ca2+. However, when focusing on the polyP-driven coacervation process, Ca2+ is a vital member of a two-reactant coacervation process alongside polyP [61]. Thereby, Ca2+ is not a simple linker but an essential partner pretending a polymer.

5. Conclusion

Due to their similarity to human hard tissue, calcium phosphate nanoparticles have been fabricated, which have qualified as a suitable filler and implant to restore bone defects due to their excellent biocompatibility together with their favorable biodegradability (reviewed in: [45]). Calcium phosphate as a multi-phase scaffold has the tendency to crystallize from the amorphous phase into the different crystalline phases. This property is advantageous for curing bone defects, especially if applied as multiphase calcium phosphate mixtures. Although this property is desirable for this application, it has the disadvantage that the material is only metabolized slowly or not at all in the body [75]. Two results prompted us to investigate the effect of polyP on the stabilization of the amorphous calcium phosphate; first, that β-glycerophosphate can be omitted from the mineralization cocktail used in mineralization studies in vitro, and second, that polyP is a potent inducer of ALP [76]. This enzyme, ALP, is a regulator of bone mineralization and has been implicated in the supply of inorganic phosphate for hydroxyapatite formation [77]. Furthermore, ALP is an enzyme that cleaves polyP [78]. The result is monomeric phosphate, which is an essential and tunable operator for hard tissue/bone formation and resistance of cells to growth stress, and protects against virulence of pathogens [79].

Aside from being an accelerator for the formation of growth promoting and organismic defense metabolites, polyP has two additional and singular properties, first, conversion into the coacervate state and second, the fact that it is the extracellular generator for the supply of ATP to cells. As reported here, oppositely charged (poly)electrolytes such as the polyanionic polyP and Ca2+ ions, when mixed in aqueous milieu at neutral pH (pH 7), undergo liquid–liquid phase-separation, coacervation, via cooperative electrostatic interactions. This associative phase separation forms a dense, fluid, polymer-rich phase (the coacervate) and a very electrolytic, aqueous phase. Likewise, and shown here, when the polyanion is added to the Ca2+ environment, the coacervate is also obtained at pH 10. Coacervation of polyP and Ca2+ is also a process occurring physiologically if the NP synthesized in the cellular acidic vacuoles are transferred to a physiological peptide-containing medium. Since the process of coacervation, as outlined, is not restricted to the extracellular matrix but also occurs intracellularly, it is very likely that also there, the polyP-NP that have been taken up by endocytosis undergo coacervation. As we have described earlier [12], it is the physiologically active form of polyP that enables the cells to embed, protect and, above all, exploit the regeneratively active biomaterial polyP.

The third distinguished feature of polyP is the harvesting of the Gibbs free energy (AG) liberated in parallel with the enzymatic cleavage of the energy-rich phosphoanhydride bonds within the polyP chains, leading to ATP formation [21]. From there, this high-energy phosphate metabolite generates ATP during the ADK reaction. The generator-associated enzymes ALP and ADK are present both extra- and intracellularly and enable ATP generation in both compartments. In the extracellular space, ATP is required for extracellular chaperone [80] and kinase activities and intracellularly for feeding anabolic cellular processes and signal transmitters (reviewed in: [21]).

The present study focuses on a modelling approach to shed light on the molecular events during polyP coacervation. The data gathered contribute to a better understanding of the biphasic nature of polyP and the formation of these phases, as well as their physiological roles, the coacervate as the functionally active state and the NP as the extra- and intracellular metabolic energy storage form of this ubiquitous inorganic, but physiological, in all cells present polymer.

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Author contributions

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Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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