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Nanoparticles as transfection agents: a comprehensive study with ten different cell lines

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The performance of transfection agents to deliver nucleic acids into cells strongly depends on the cell type. In a comprehensive study, nine different cell lines and primary human mesenchymal stem cells were transfected with DNA encoding for enhanced green fluorescent protein (eGFP). As transfection agents, two kinds of cationic multi-shell calcium phosphate nanoparticles and the commercially available transfection agent Lipofectamine were used. The transfection efficiency was measured by fluorescence microscopy by counting the percentage of green fluorescent cells which expressed eGFP as well as qPCR. Furthermore, the uptake of fluorescent calcium phosphate nanoparticles was measured by fluorescence microscopy. The cell viability was measured by the MTT test after incubation with nanoparticles and Lipofectamine. All cell types took up nanoparticles (with different efficiency), but the expression of eGFP was strongly different, demonstrating that the uptake not necessarily leads to processing of a gene. A clear correlation was found between the transfection efficiency and the cell viability that was independent on the transfection agent: a high transfection efficiency was clearly correlated with a low cell viability and vice versa.

Introduction

The non-viral introduction of DNA into living cells is called transfection. The DNA has to cross the cell membrane and to enter the nucleus where its genetic code is transcribed and the corresponding protein is produced.¹⁻⁵ Together with gene silencing where siRNA is suppressing the transcription of a specific protein,⁶⁻⁹ this approach is called gene therapy, an exciting area of biomedical research.^{8,10,11}

The efficiency of transfection is notoriously poorly reproducible and difficult to compare between different studies. The main factors are the target cell type and the kind of transfection agent. Given the fact that for a successful transfection, the DNA has to cross the cell membrane, to escape the lysosome, to enter the nucleus and to avoid degradation on its pathway, it is clear that the cell type and the transfection agent will strongly influence this process at each step. 1,7,12-20 Therefore, many experiments reported in the literature are difficult to compare because different cell types and different transfection agents were used. Furthermore, it is not always clear whether a low transfection rate was due to a poor particle uptake or to a poor processing of the DNA by the cell.

We have therefore treated ten cell types of different origin and differentiation with nanoparticles to compare the efficiencies of uptake and transfection with eGFP-encoding DNA. As control reagent, the commercial transfection agent Lipofectamine was used. All cells were incubated in parallel with the same batches of nanoparticles and transfection agent. Of course, all cell lines have different properties with respect to particle uptake, ability to be transfected, internal protein synthesis, sensitivity towards nanoparticles. It was our goal to elucidate exactly these differences (without going too deep into the intracellular mechanisms and differences between cell types) by challenging the cells under identical conditions with identical nanoparticles.

Experimental

Materials

Poly(ethyleneimine) (PEI, $M_{\rm w}=25~000~{\rm g~mol}^{-1}$) was obtained from Sigma-Aldrich. Fluorescent FITC-labelled (fluorescein isothiocyanate) PEI ($M_{\rm w}=25~000~{\rm g~mol}^{-1}$) was obtained from Surflay (Berlin, Germany). For transfection, pcDNA3eGFP was used. Plasmid DNA (pcDNA3eGFP) coding for eGFP (enhanced green fluorescent protein) was isolated from *Escherichia coli* using a NucleoBond® PC 10000 EF endotoxinfree plasmid DNA purification kit (Machery-Nagel, Düren, Germany). This plasmid DNA had a molecular weight of $M_{\rm w}=1~871~704~{\rm g~mol}^{-1}$.

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All other reagents were of analytical grade and used without further purification.

Instruments

Scanning electron microscopy was performed with an ESEM Quanta 400 instrument with gold/palladium-sputtered samples. Dynamic light scattering and zeta potential determinations were performed with a Zetasizer nanoseries instrument (Malvern Nano-ZS, laser: $\lambda = 633$ nm) using the Smoluchowski approximation and taking the data from the Malvern software without further correction. The particle size data refer to scattering intensity distributions (z-average). Centrifugation was performed at 4 °C with a Heraeus Fresco 21 instrument for smaller volumes, and a Sorvall WX ultra series centrifuge for larger volumes.

The amount of calcium was determined by atomic absorption spectroscopy (AAS) with an M-Series AA spectrometer (ThermoElectron Corporation, Schwerte, Germany).

Transfection efficiencies were determined by transmission light microscopy and fluorescence microscopy with a Keyence Biorevo BZ-9000 instrument (Osaka, Japan). The microscope was equipped with Keyence GFP-B (excitation: 470/40) filters. Images were recorded with the BZ-II viewer software and processed with the BZ-II analyzer software. Real-time quantitative PCR (qPCR) was carried out with a 7500 Fast Real-time System (Applied Biosystems, Life Technologies, Carlsbad, USA). Data were recorded and analysed with the 7500 Fast Real-time System software. UV/vis measurements were performed with a Cary 300 BIO UV-visible spectrophotometer with 1 cm quartz microcuvettes. The cell viability was determined with the MTTtest by spectrophotometric analysis with a Multiscan FC instrument (ThermoFisher scientific, Vantaa, Finland) at $\lambda = 570 \text{ nm}.$

Freeze-drying of nanoparticle dispersions was performed with an Alpha 2-4 LSC freeze-dryer (Martin Christ, Osterode, Germany) as described in ref. 21.

Lipofectamine 2000® was obtained from Life Technologies (Invitrogen, Carlsbad, USA). This commercially available transfection agent consists of positively charged lipid molecules. By mixing DNA with Lipofectamine 2000®, positively charged liposomes are formed, which are attracted to the cellular membrane by electrostatic interactions. After a contact between the liposomes and the cellular membrane is formed, the liposomes are able to fuse with the cellular membrane to release the nucleic acids into the cell.22,23

All errors given represent standard deviations.

Synthesis of triple-shell calcium phosphate nanoparticles: CaP-DNA-CaP-PEI

Cationic triple shell calcium phosphate nanoparticles (CaP-DNA-CaP-PEI) were synthesized in analogy to earlier reported procedures.^{24,25} Equal volumes of solutions of calcium-L-lactate (6.25 mM, pH 9) and diammonium hydrogen phosphate (3.74 mM, pH 9) were pumped into a solution of DNA (pcDNA3eGFP; 1 mg mL⁻¹) to give a final DNA concentration of 167 $\mu g \text{ mL}^{-1}$ (e.g. 217 μL of calcium and phosphate solution

each and 87 µL DNA solution). Calcium and phosphate solutions were pumped through a Y-connector with a syringe pump, resulting in a nucleation time of 0.25 s for calcium phosphate formation. The primary nanoparticles were stabilized by coating with DNA. Excess DNA was removed by centrifugation at 20 000g for 15 min. The precipitated nanoparticles were then redispersed in the same volume of water by ultrasonication for 10 s (UP50H, Hielscher, Ultrasound Technology; sonotrode 2, cycle 0.8, amplitude 70%). After this procedure, single-shell calcium phosphate nanoparticles with a shell of DNA were obtained.

The second shell of calcium phosphate and the third shell of PEI were added as follows. Equal volumes of calcium-1-lactate (6.25 mM, pH 9) and diammonium hydrogen phosphate (3.74 mM, pH 9) solutions were subsequently added (e.g. 217 µL) to the colloidal dispersion, followed by addition of PEI solution (2 mg mL $^{-1}$ in water) to give a final concentration of 87 μ g mL $^{-1}$ of PEI (e.g. 45 µL). Excess PEI was removed by ultracentrifugation and redispersion in water.

The concentration of DNA in the nanoparticles was determined by measuring the absorbance at 260 nm in the supernatants of the first centrifugation step. After purification of the nanoparticle dispersion from free nucleic acids, excess ions and free PEI, about 7.6 $\mu g \text{ mL}^{-1}$ of DNA was left in the colloidal dispersion, which corresponds to about 9% of the DNA applied during the synthesis.

Synthesis of calcium phosphate nanoparticles with a silica shell: CaP-DNA-PEI-SiO2-SH

Calcium phosphate nanoparticles with a silica shell (CaP-DNA-PEI-SiO₂-SH) were synthesized as reported earlier.²⁶ 5 mL of calcium-1-lactate solution (18 mM, pH 10), 5 mL of diammonium hydrogen phosphate solution (10.8 mM, 10 pH) and 7 mL of PEI solution (2 mg mL⁻¹) were simultaneously pumped during 1 min into 20 mL of purified water. The dispersion was stirred for 20 min. To add DNA to the nanoparticles, 900 µL of this dispersion were mixed with 100 μ L of DNA (1 mg mL⁻¹).

To add a silica shell, 5 μL TEOS and 2.6 μL 30–33% aqueous ammonia solution were dissolved in 4 mL ethanol. The CaP-DNA-PEI nanoparticle dispersion (1 mL) was added and stirred for 16 h at room temperature. The particles were purified by centrifugation at 20 000g for 30 min and redispersed in 1 mL of water by ultrasonication for 10 s (UP50H, Hielscher, Ultrasound Technology; sonotrode 2, cycle 0.8, amplitude 70%). Subsequently, the silica shell was functionalized with thiol groups by silanization. 5 μL of 3-mercaptopropyltrimethoxysiloxan (MPS) were dissolved in 4 mL of ethanol and stirred for 8 h after the addition of the colloidal dispersion. Purification was performed as described above. The concentration of DNA in the dispersion was determined by dissolving the particles in hydrochloric acid and measuring the absorbance at 260 nm. The analysis showed that about 43 $\mu g \text{ mL}^{-1}$ of DNA were present in the colloid which corresponds to about 43% of the initial amount.

After the synthesis, both types of particles were freeze-dried in the presence of trehalose as a cryoprotectant.²¹ 10 mg mL⁻¹ of D-(+)-trehalose dihydrate (Sigma-Aldrich) were added to a 1 mL

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aliquot of the dispersion. The aliquots were shock-frozen in liquid nitrogen and subsequently freeze-dried at 0.31 mbar and -10 °C. The particles were redispersed in the same amount of water before use in cell culture experiments.

Synthesis of fluorescent single-shell calcium phosphate nanoparticles: CaP-PEI

Fluorescing single-shell CaP-PEI nanoparticles were prepared by functionalizing the calcium phosphate core with FITClabelled PEI. Equal volumes (0.5 mL) of calcium-L-lactate (6.25 mM, pH 9) and diammonium hydrogen phosphate (3.74 mM, pH 9) were pumped into a solution of PEI-FITC (2 mg mL⁻¹, 0.2 mL). The dispersion was purified by ultracentrifugation (20 000g, 20 min) and redispersed in the same amount of water by ultrasonication for 10 s (Hielscher UP400S; 20% amplitude). These particles were used in the cell culture experiments after the synthesis without prior to freeze-drying.

Incubation of cells with nanoparticles

The amount of calcium in the colloidal dispersions was determined by AAS. This was converted into the amount of calcium phosphate by assuming the stoichiometry of hydroxyapatite, Ca₅(PO₄)₃OH. Before the AAS measurement, the particles were dissolved in hydrochloric acid. Using the density of hydroxyapatite and the particle diameter by SEM, we computed the particle concentration, assuming a spherical shape of the particles (see ref. 27 and 28 for a typical calculation).

Nine cell lines and one primary cell culture of human mesenchymal stem cells (hMSCs) were cultivated in cell culture flasks (Sarstedt, USA) under 5% CO2 and humidified atmosphere and 37 °C according to standard cell culture protocols. All cell types and passage ranges as well as the corresponding cell media are given in Table 1.

For microscopic imaging and viability assays, all cells were passaged at the same day, except for the suspension cell lines (THP-1 and U937). In order to obtain adherent cells, THP-1 and U937 cells were previously seeded in 96-well plates (Sarstedt, USA) in a density of 5×10^3 cells per well in 100 μL of cell medium and differentiated into macrophages by the addition of 100 nM of PMA (4β-phorbol-12β-myristate-13α-acetate, Sigma-Aldrich, USA) solution per well and incubated for 3 days. Afterwards, the cell medium was changed and the cells were further treated exactly like the other cell lines.

One day before the incubation with nanoparticles or Lipofectamine, all adherent cell lines were trypsinized with trypsin/ EDTA solution (Gibco™, Life Technologies, USA) and seeded in 96-well plates in a density of 5×10^3 cells per well in 100 µL of cell medium (2×10^3) cells for CV-1 cells because of the high proliferation rate).

Approximately 24 h later, the cell culture medium was removed and 100 µL of fresh medium containing either DNAloaded nanoparticles or Lipofectamine was added to the cells. The final concentration of DNA was 0.3 µg per well. This corresponded to about 7.97×10^9 CaP-DNA-CaP-PEI particles per well or 1.31×10^9 CaP-PEI-DNA-SiO₂-SH particles per well. The transfection with Lipofectamine was carried out according to the manufacturer's recommendation, adjusted for 96-well plate volumes. The concentration of DNA in Lipofectaminetreated samples was 0.2 µg per well. Approximately 18 h after the incubation with transfection agents, the cell medium for all samples was changed and the cells were incubated for another 48 h. The transfection efficiency was obtained by imaging the cells by fluorescence and light microscopy and computed as follows:

Table 1 List of all cell types and cell culture media used. DMEM: Dulbecco's modified Eagle's medium; FCS: fetal calf serum; NEAA: non-essential amino acid cell culture supplement; MEM: minimum essential medium

#	Cell line	Passage	Description	Characteristics	Cell culture medium
1	HeLa	P21-24	Human epithelial cervical cancer cells	Adherent, epithelial	DMEM + 100 U mL ⁻¹ streptomycin/ penicillin + 10% FCS
2	MG-63	P37-39	Human osteosarcoma cells	Adherent, fibroblasts	
3	Huh 7	P7-9	Human hepato cellular carcinoma cells	Adherent, epithelial-like hepatoma cell line	DMEM + 100 U mL ⁻¹ streptomycin/ penicillin + 10% FCS + 1% NEAA
4	CV-1	P8-10	Monkey kidney fibroblast cells	Adherent, fibroblasts	•
5	Mode-K	P13-15	Mouse intestinal epithelial cells	Adherent, epithelial	DMEM + 100 U mL ⁻¹ streptomycin/ penicillin + 10% FCS + 2 mM _L -glutamine
6	MC3T3	P9-11	Osteoblastic cell line from mouse calvaria	Adherent, fibroblast-like	αMEM + 100 U mL ⁻¹ streptomycin/ penicillin + 10% FCS
7	U937	P8-10	Human histiocytic leukemia cell line	Suspension, monocytes	RPMI-1640 + 100 U mL ⁻¹ streptomycin/ penicillin + 10% FCS + 10 mM hepes + 2
8	THP-1	P47-49	Human acute monocytic leukemia cell line	Suspension, monocytes	mM ь-glutamine
9	4T1	P21-23	Mammary carcinoma cells from mouse mammary gland	Adherent, epithelial	RPMI-1640 + 100 U mL ⁻¹ streptomycin/ penicillin + 10% FCS
10	hMSC	P6-7	Human bone marrow mesenchymal stem cells	Adherent, stem cells	Mesenchymal stem cells (MSC) growth medium

(number of green fluorescent cells)/(number of all cells) × 100%

Two images were taken per well, and for each cell type two separate wells were prepared. The whole experiment was performed twice, giving a total of $2 \times 2 \times 2 = 8$ data points for each cell type.

qPCR analyses

qPCR probes the production of eGFP on the mRNA level. Again, an average is obtained by analysis of a whole cell culture sample. After trypsinization and centrifugation (900 rpm, 3 min), the cells were lysed in lysis buffer (Qiagen, Hilden, Germany) and frozen at -80 °C until total RNA isolation. Total RNA was purified from lysed cells with the RNeasy Kit from Qiagen (Hilden, Germany) according to the manufacturer's instructions. After isolation of the RNA, about 1 µg of RNA was used for cDNA synthesis. cDNA was synthesized with MMV-L (H-) point mutant reverse transcriptase (Promega, Fitchburg, USA). About 5 to 20 ng of cDNA were used for qPCR analysis. Each sample in qPCR was analysed in duplicate. SYBR green master mix (life technologies, Carlsbad, USA) was used for amplification and detection. Relative amounts of mRNA were calculated from standard curves and normalized to the housekeeping gene RPS9. The following primers were used:

RPS9:

5' CTGGACGAGGGCAAGATGAAGC

3' TGACGTTGGCGGATGAGCACA

eGFP:

5' CTACGGCGTGCAGTGCTTCAG

3' CTCGGCGCGGGTCTTGTAG

In all cases, cells cultivated under the same conditions but without any treatment, were used as control.

Cell viability

The cell viability was analysed by an MTT-assay 48 h after the transfection. MTT (3-(4,5-dimethylthiazol-2-yl)-2,5diphenyltetrazolium bromide; Molecular probesTM, Life Technologies, Carlsbad, USA) was dissolved in PBS (5 mg mL⁻¹) and then added to the required amount of cell culture medium to give a final MTT concentration of 1 mg mL⁻¹. The cell culture medium of the transfected cells was replaced with the MTTcontaining medium (150 µL) and incubated for 1 h at 37 °C under 5% CO2 in humidified atmosphere. Then the MTTcontaining medium was removed and 150 µL of dimethyl sulfoxide (DMSO) was added to each well and incubated for another for 30 min. Finally, two aliquots of 50 µL from each well were taken for the photometric analysis at $\lambda = 570$ nm. The absorption of the supernatant of the transfected cells was normalized to that of the control (untreated cells).

Uptake of nanoparticles by cells

For the uptake studies, cells were seeded into 8-well culture slides (Falcon, USA) with 5 \times 10^4 cells per well in 500 μL of cell medium as described previously for the transfection experiments. The cells were treated with 70 μL of PEI–FITC

functionalized single shell calcium phosphate nanoparticles $(1.32 \times 10^{10} \text{ particles})$ and left for 3 h incubation. Afterwards, the cells were washed 3 times with PBS and fixed with 3.7% solution of formaldehyde to remove any adhering or dispersed nanoparticles. Then, the formaldehyde solution was removed, the cells were washed twice with PBS and stained with Cell Mask® (Molecular ProbesTM, Life Technologies, USA) and DAPI (4',6-diamidino-2-phenylindole, Molecular ProbesTM, Life Technologies, USA) dyes to visualise the cell membrane and the nucleus, respectively.

The nanoparticle uptake was visualized by fluorescence microscopy with a Keyence Biorevo BZ-9000 microscope equipped with filters for TRITC (excitation: 540/25, emission: 605/55), GFP BP (excitation: 470/40, emission: 535/50) and DAPI (excitation: 360/40, emission: 460/50) channels under $40 \times$ magnification.

Results and discussion

We have studied the reaction of cell types of different origin and differentiation state to nanoparticles (see Fig. 1 and Table 1). We used a range of cells of animal origin as well as of human origin.

Two types of calcium phosphate nanoparticles were synthesized and loaded with eGFP-DNA for transfection of the different cell lines. The full characterization of the particles is given in Table 2.

SEM images of the particles show mainly spherical primary particles in the size range between 30 nm and 140 nm (Fig. 2). DLS analysis gave particle sizes between 400 nm and 750 nm, indicating some degree of agglomeration in the dispersion. Note that an excessive use of ultrasound to enhance the dispersion is not recommended because the biomolecules may be damaged.²⁹ The zeta potential of +20 to +25 mV is at the border for a purely electrostatic colloidal stabilization, but the presence of the polyelectrolyte polyethyleneimine (PEI) adds a steric component to the colloidal dispersion.²⁴

The first barrier for a successful transfection is the cell membrane. In order to verify whether all cells take up the calcium phosphate nanoparticles, fluorescing particles without DNA, but with an outer layer of FITC-PEI were prepared. The uptake of calcium phosphate nanoparticles by HeLa cells has

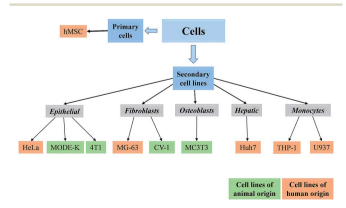
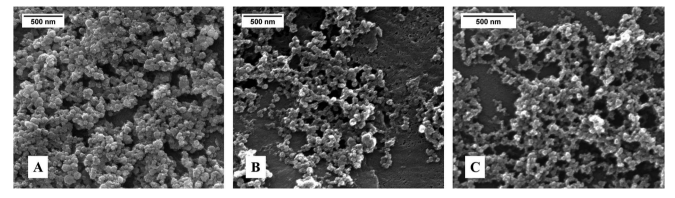


Fig. 1 All cell types used in the transfection experiments.

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Table 2 Properties of the nanoparticles used. All concentrations refer to the stock solutions that were then added to the cells. The particle concentration is based on a number of assumptions (see experimental part), therefore it is associated with a relative error of about $\pm 25\%$

Type of nanoparticles	Concentration of DNA/ $\mu g \ mL^{-1}$	Diameter by SEM/nm	Diameter by DLS/nm	PDI by DLS	Zeta potential by DLS/mV	Number of particles/mL ⁻¹	Number of DNA molecules per particle
CaP-PEI-DNA- SiO ₂ -SH	46 ± 10	40-140	739 ± 24	0.4-0.56	+24 ± 1	${\sim}1.25\times10^{11}$	~119
CaP-DNA-CaP-PEI CaP-PEI-FITC	8 ± 3 —	40-100 30-70	437 ± 180 209 ± 5	0.3-0.5 0.5-0.7	$^{+20}\pm 1$ $^{+21}\pm 1$	\sim 2.02 × 10 ¹¹ \sim 1.88 × 10 ¹¹	~12 —
Cai ILI IIIC		30 70	207 ± 3	0.5 0.7	121 1	-1.00 × 10	



SEM images of the calcium phosphate nanoparticles: (A) CaP-PEI-DNA-SiO₂-SH; (B) CaP-DNA-CaP-PEI; (C) CaP-PEI-FITC.

been previously investigated by confocal laser scanning microscopy (CLSM). The uptake of nanoparticles occurred already during 3 h of incubation.28

To study the uptake of nanoparticles, we used positively charged single-shell particles (CaP-PEI-FITC) with a very similar diameter of the DNA-loaded particles. Their zeta potential was identical to that of the DNA-loaded particles (Table 2). The ten cell types readily took up calcium phosphate nanoparticles during 3 h of incubation (Fig. 3). Although the extent of particle uptake was different for the cell lines, and appeared to vary between individual cells, nanoparticles and cells were clearly co-localized. This indicates that the particles are present inside the cell and not just adsorbed on the surface, as shown earlier for HeLa cells.28,30

This is a prerequisite for the subsequent transfection that is clearly fulfilled for all cell types. However, there are different possibilities for cells to take up nanoparticles (e.g. endocytosis, micropinocytosis, phagocytosis).16,31-34 The nanoparticles end up in different intracellular compartments (endosome, lysosome, phagosome), and the DNA might be degraded before it enters the lysosome and finally the nucleus.

For the transfection experiments, the amount of DNA per well was kept constant for all cell types in the case of the nanoparticles. As control, Lipofectamine was used with the DNA concentration as recommended by the manufacturer. Because the concentration of DNA was different in both nanoparticle formulations, the number of particles per well was also different. However, the number of particles per well was much larger than the number of cells (Table 2).

Fig. 4 shows the expression of eGFP by the different cell lines after the transfection. Brightfield and fluorescence images were

microscopically recorded. While transfected cells were visible only in the fluorescence images, all cells were visible in the brightfield images. The transfection efficiency was obtained from microscopic images by counting the fraction of green cells. The numerical transfection data are shown in Fig. 5 and Table 3.

The numbers of cells that were counted to determine the transfection efficiency are given in Table 3. The numbers denote the cells counted in the brightfield images, i.e. 100% of the cells of which a fraction was green fluorescent due to transfection with eGFP. For THP-1 and U937 cells, no transfection was observed (no green fluorescent cells).

Clear distinctions in the transfection efficiency between the ten cell types are seen. The most effective transfection results were obtained with HeLa, Huh7, Mode-K and MG-63 cell lines with both nanoparticles and Lipofectamine. Lipofectamine was the most efficient transfection agent, but the efficiency of the nanoparticles was typically close to Lipofectamine. There were no major differences between the two types of cationic calcium phosphate nanoparticles, indicating that their surface termination (thiol or PEI) did not play a major role.

A set of cell types that was transfected with much lower efficiency comprised 4T1, MC3T3, hMSC, and CV-1 cells. The only two cell lines that were transfected by any of three applied transfection agents were the monocyte cell lines U937 and THP-1, which were previously differentiated into macrophages. Although these cells took up the nanoparticles as shown in Fig. 3, a transfection of these cell lines did not occur. This indicates that the DNA was not processed inside the cells.

Fig. 5 quantitatively shows that some cell types are more easily transfectable than others. In fact, the cells that were

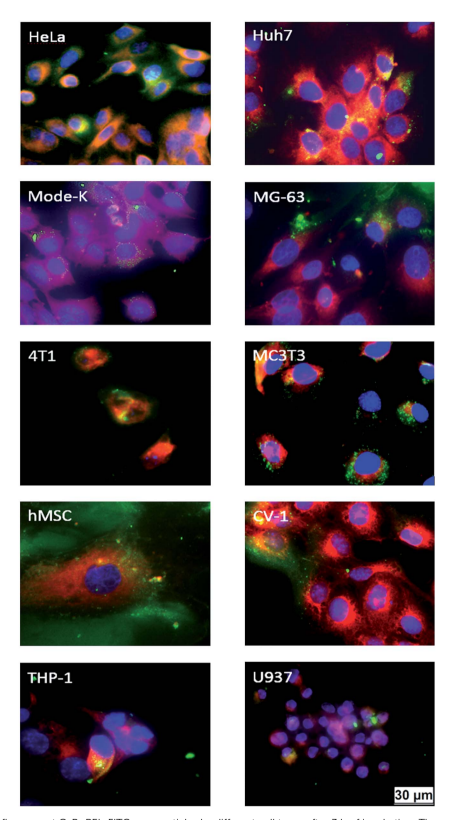


Fig. 3 Uptake of green fluorescent CaP-PEI-FITC nanoparticles by different cell types after 3 h of incubation. The cell nucleus was stained in blue by DAPI, and the cell membrane was stained red by CellMask. The cells were thoroughly washed before imaging to remove dispersed and weakly adhering nanoparticles. All cells took up the cationic nanoparticles, although to a different extent.

transfected with high efficiency were cells that divide more rapidly, namely HeLa, Huh7, Mode-K, and MG-63. It is noteworthy that for efficiently transfected cells, a higher cytotoxicity

was observed as well. These cells were also highly abundant on the images (Table 3), indicating their higher proliferation rate. The correlation of the transfection efficiency with the

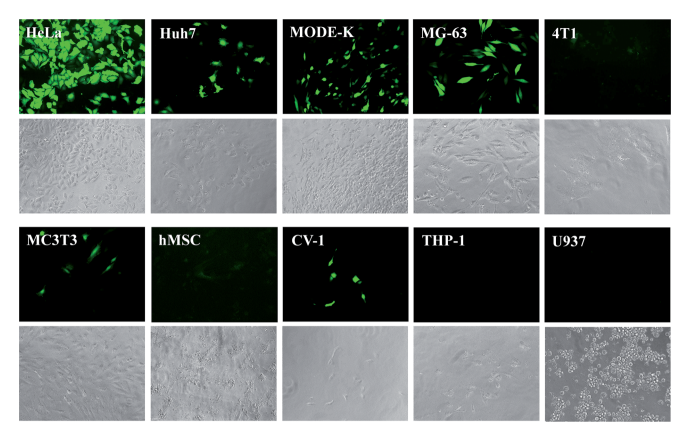


Fig. 4 Fluorescence and brightfield micrographs, showing the eGFP expression in different cell types after transfection with CaP-DNA-CaP-PEI nanoparticles. Brightfield images are shown below the corresponding fluorescence images. For CaP-PEI-DNA-SiO₂-SH-nanoparticles, very similar results were obtained (see Table 3).

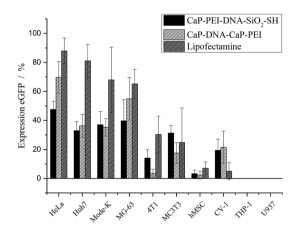


Fig. 5 Transfection efficiency of different cells after incubation with nanoparticles and Lipofectamine. In the case of nanoparticles, 0.3 μ g DNA per well were applied. In the case of Lipofectamine, 0.2 μ g DNA per well were applied (96 well format).

proliferation rate is in line with reports in the literature that showed a dependence of the transfection on cell division. In order to promote protein expression the plasmid DNA must enter the nucleus. This crucial step was shown to occur after mitosis in HeLa cells, independently from the transfection agent used, as observed by life cell imaging.^{35,36} The protein

Table 3 Number of living cells (both transfected and untransfected) counted in the individual experiments (brightfield image) to determine the transfection efficiency. In the case of the THP-1 and U937 cells, not a single fluorescing cell was detected by fluorescence microscopy (transfection efficiency zero)

	${\it CaP-PEI-DNA-SiO}_2{-\it SH}$	CaP-DNA-CaP-PEI	Lipofectamine
HeLa	878	2461	436
Huh7	1325	497	735
Mode-K	448	2262	896
MG-63	794	652	886
4T1	187	220	88
MC3T3	532	747	102
hMSC	85	111	56
CV-1	208	139	49
THP-1	>200	>200	>200
U937	>200	>200	>200

expression was observed 2–5 h after mitosis.³⁶ Furthermore, it was shown for hMSC that cells were transfected more efficiently under conditions that promote cell division.³⁷ On the other hand, cells that divide very slowly or not at all were transfected only poorly or even not at all (4T1, hMSC, THP-1, U937).

The low transfection efficiency in monocytes/macrophages may also be due to high enzymatic activities in phagosomes/phagolysosomes. It is likely that these cell lines take up the nanoparticles and Lipofectamine by phagocytosis and not by

endocytosis. Macrophages are scavenging cells with a high level of digestive enzymes which will degrade DNA in the phagosome, resulting in an inefficient transfection.38

To underline the uptake and transfection results, the transfection efficiency was also measured at the mRNA level of protein expression (Fig. 6). The two cell types Huh7 and hMSC were analysed by qPCR. Transfection was carried out with CaP-PEI-DNA-SiO₂-SH nanoparticles.

The different transfection efficiencies of the different cell types were even more pronounced by qPCR analysis. The relative expression of eGFP in untreated control cells was normalized to 1. In the case of the Huh7 cells, the mRNA signal of eGFP was 20 000 times elevated in comparison to the untreated control cells. In the case of hMSC, the signal was only 1000 times higher.

On the other hand, the cell viability showed a completely different trend (Fig. 7). The cell lines that were efficiently transfected also displayed a high toxicity, and poorly transfected cell lines were less sensitive, even though all cell lines took up

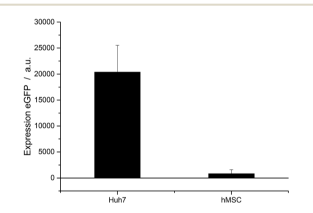


Fig. 6 Relative expression of eGFP by Huh7 cells and hMSC, determined by qPCR. The relative expression is normalized to the signal of untreated control cells. 1.5 µg of DNA per well were applied with CaP-PEI-DNA-SiO₂-SH nanoparticles (24 well format).

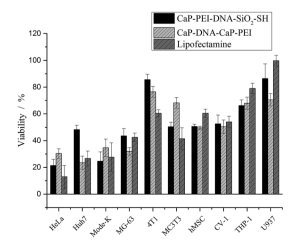


Fig. 7 Viability of cell lines after incubation with different transfection agents. In the case of nanoparticles, 0.3 µg DNA per well were applied. In the case of Lipofectamine, 0.2 µg DNA per well were applied (96 well format)

particles (Fig. 3). This correlation was independent from the transfection agent used, *i.e.* nanoparticles or Lipofectamine.

Fig. 8 shows the correlation between the cell viability and the transfection efficiency. It is important to note that a correlation never proves a causality, but may point to the fact that two variables (in this case, the transfection efficiency and the cell viability) depend on a third common parameter. This parameter is not yet known, but possibly the cationic compounds Lipofectamine and poly(ethyleneimine) are on the one hand promoting the uptake and the endosomal escape of DNA, and on the other hand have harmful side effects.39-42

Liu et al. showed that lysosomal rupture may cause necrosis.43 However, the transfection efficiency depends also on the lyso-/endosomal release of the DNA in the case of the transfection agents used (Fig. 9). Elevated intracellular calcium levels induced by nanoparticles should not play a major role causing the toxicity by apoptosis as shown earlier. 44,45

Fig. 9 shows the general mechanism which we propose for the uptake of calcium phosphate nanoparticles by eukaryotic cells. A nanoparticle loaded with pDNA (plasmid DNA) (I) must overcome a number of biological barriers to enter the cell nucleus and finally be processed inside. 46 After endocytosis (II), the nanoparticles end up in an endosome that then fuses with a lysosome (III). Unless the lysosome bursts before the full degradation of the DNA, the degraded DNA will not be effective anymore, and the transfection efficiency will be zero. 1,5,19,47 An endosomal escape is supported by an increased osmotic pressure inside the lysosome due to neutralization of acid by suitable basic compounds (IV). Poly(ethyleneimine) is well known for the so-called proton sponge effect,48 and the acid-soluble calcium phosphate will also lead to a considerable number of calcium and hydrogen phosphate ions after dissolution. 43,44,49,50 The lysosome ruptures (V) and released its cargo into the cytosol. Excess calcium is pumped out (VI) and the DNA is entering the nucleus, possibly together with PEI as polyplex.

Cell uptake studies performed with different synthetic vectors and on different cell types showed that endocytosis occurs efficiently either by the clathrin and/or by caveolae-

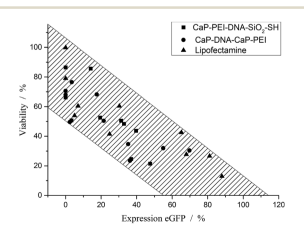
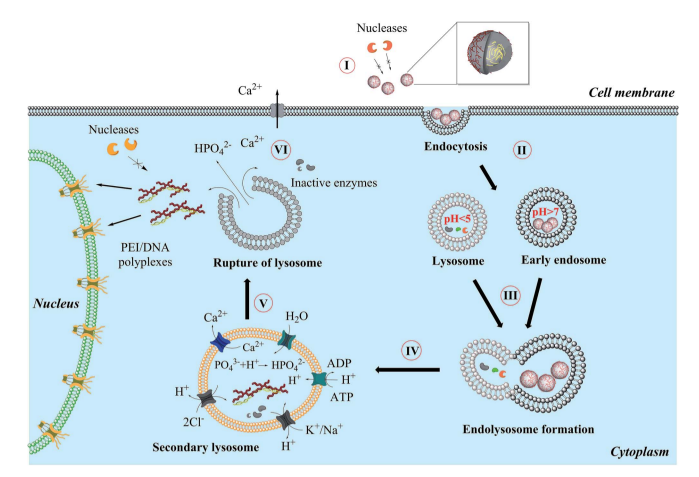


Fig. 8 Correlation between the transfection efficiency and the cell viability. All data points occur in the shaded region, independent of the transfection agent. The shaded area was chosen to include all data points in this inverse proportionality.



General mechanism proposed for the uptake and intracellular processing of calcium phosphate nanoparticles by cells.

dependent pathways. 47,51-57 Cationic calcium phosphate nanoparticles were shown to enter HeLa cells by micropinocytosis.²⁸

When the endocytosis of pDNA/cationic polymer polyplexes occurs by the clathrin-dependent pathway, these polyplexes can escape from the endosomal route and reach the cytosol, taking advantage of the proton sponge effect induced by the polymers that not only carry cationic groups for condensing pDNA, but also basic groups like secondary amines42,48,58-60 or imidazole rings.61 To enhance the nuclear targeting of pDNA, specific DNA sequences recognized by transcription factors such as NFkB can be inserted into pDNA.⁶²⁻⁶⁴ However, the sub-cellular trafficking of pDNA to the nucleus up to now remains unclear and poorly described in the literature.46

Conclusions

Different cell lines show a different susceptibility to transfection, despite the fact that they all take up nanoparticles. This is negatively correlated with the cell viability, irrespective of the transfection agent, pointing to a hitherto unknown common mechanism of transfection and cytotoxicity. Transfection systems that show a good transfection efficiency cause to a high cytotoxicity and vice versa. Altogether, we have shown that cells react very differently to transfection agents, including nanoparticles, and that one may not generalize results obtained for one cell line.

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References

- 1 F. Pérez-Martínez, J. Guerra, I. Posadas and V. Ceña, Pharm. Res., 2011, 28, 1843-1858.
- 2 P. Midoux, C. Pichon, J. J. Yaouanc and P. A. Jaffres, Br. J. Pharmacol., 2009, 157, 166-178.
- 3 S. Bhattacharya and A. Bajaj, Chem. Commun., 2009, 4632-
- 4 M. Morille, C. Passirani, A. Vonarbourg, A. Clavreul and J. P. Benoit, Biomaterials, 2008, 29, 3477-3496.
- 5 I. S. Zuhorn, J. B. F. N. Engberts and D. Hoekstra, Eur. Biophys. J., 2007, 36, 349-362.
- 6 J. Kurreck, Angew. Chem., Int. Ed., 2009, 48, 1378-1398.
- 7 J. Wang, Z. Lu, M. G. Wientjes and J. L. S. Au, AAPS J., 2010, 12, 492-503.

Paper

8 A. S. Abdolhamid, S. S. Hedayati, R. H. Poor, S. Madani, S. S. Poor and S. Panahi, *J. Genet.*, 2010, **89**, 527–537.

- 9 P. N. Pushparaj and A. J. Melendez, *Clin. Exp. Pharmacol. Physiol.*, 2006, 33, 504–510.
- 10 X. Guo and L. Huang, Acc. Chem. Res., 2012, 45, 971-979.
- 11 D. Reischl and A. Zimmer, Nanomedicine, 2009, 5, 8-20.
- 12 J. Zabner, A. J. Fasbender and T. Moninger, *J. Biol. Chem.*, 1995, 270, 18997–19007.
- 13 M. Nishikawa and L. Huang, *Hum. Gene Ther.*, 2001, **12**, 861–870.
- 14 C. M. Wiethoff and C. R. Middaugh, J. Pharm. Sci., 2003, 92, 203–217.
- 15 S. C. de Smedt, K. Remaut, B. Lucas, K. Braeckmans, N. N. Sanders and J. Demeester, Adv. Drug Delivery Rev., 2005, 57, 191–210.
- 16 H. Hillaireau and P. Couvreur, Cell. Mol. Life Sci., 2009, 66, 2873–2896.
- 17 M. S. Shim and Y. J. Kwon, FEBS J., 2010, 277, 4814–4827.
- 18 F. M. Kievit and M. Q. Zhang, *Adv. Mater.*, 2011, **23**, H217–H247.
- 19 P. Kesharwani, V. Gajbhiye and N. K. Jain, *Biomaterials*, 2012, 33, 7138–7150.
- 20 V. Sokolova and M. Epple, Angew. Chem., Int. Ed., 2008, 47, 1382–1395.
- 21 J. Klesing, S. Chernousova and M. Epple, *J. Mater. Chem.*, 2012, 22, 199–204.
- 22 X. Ming, K. Sato and R. L. Juliano, *J. Controlled Release*, 2011, **153**, 83–92.
- 23 B. Dalby, S. Cates, A. Harris, E. C. Ohki, M. L. Tilkins, P. J. Price and V. C. Ciccarone, *Methods*, 2004, 33, 95–103.
- 24 V. Sokolova, S. Neumann, A. Kovtun, S. Chernousova, R. Heumann and M. Epple, *J. Mater. Sci.*, 2010, 45, 4952– 4957.
- 25 V. V. Sokolova, I. Radtke, R. Heumann and M. Epple, *Biomaterials*, 2006, 27, 3147–3153.
- 26 D. Kozlova, S. Chernousova, T. Knuschke, J. Buer, A. M. Westendorf and M. Epple, *J. Mater. Chem.*, 2012, 22, 396–404.
- 27 G. Doerdelmann, D. Kozlova, S. Karczewski, R. Lizio, S. K. Knauer and M. Epple, J. Mater. Chem. B, 2014, 2, 7250–7259.
- 28 V. Sokolova, D. Kozlova, T. Knuschke, J. Buer, A. M. Westendorf and M. Epple, *Acta Biomater.*, 2013, 9, 7527–7535.
- 29 M. L. Wu, S. S. Freitas, G. A. Monteiro, D. M. Prazeres and J. A. Santos, *Biotechnol. Appl. Biochem.*, 2009, 53, 237–246.
- 30 V. Sokolova, O. Rotan, J. Klesing, P. Nalbant, J. Buer, T. Knuschke, A. M. Westendorf and M. Epple, *J. Nanopart. Res.*, 2012, 14, 910.
- 31 I. Canton and G. Battaglia, *Chem. Soc. Rev.*, 2012, **41**, 2718–2739.
- 32 T. G. Iversen, T. Skotland and K. Sandvig, *Nano Today*, 2011, **6**, 176–185.
- 33 G. Sahay, D. Y. Alakhova and A. V. Kabanov, *J. Controlled Release*, 2010, **145**, 182–195.
- 34 G. J. Doherty and H. T. McMahon, Annu. Rev. Biochem., 2009, 78, 857–902.

- 35 K. Hakamada and J. Miyake, *J. Biosci. Bioeng.*, 2012, **113**, 124–127.
- 36 I. Kirchenbuechler, D. Kirchenbuechler and M. Elbaum, *Exp. Cell Res.*, 2015, DOI: 10.1016/j.yexcr.2014.11.019.
- 37 W. J. King, N. A. Kouris, S. Choi, B. M. Ogle and W. L. Murphy, *Cell Tissue Res.*, 2012, **347**, 689–699.
- 38 S. Dokka, D. Toledo, X. Shi, J. Ye and Y. Rojanasakul, *Int. J. Pharm.*, 2000, **206**, 97–104.
- 39 H. T. Lv, S. B. Zhang, B. Wang, S. H. Cui and J. Yan, *J. Controlled Release*, 2006, **114**, 100–109.
- 40 A. Paul, C. J. Eun and J. M. Song, *Polymer*, 2014, 55, 5178-5188.
- 41 W. T. Godbey, K. K. Wu and A. G. Mikos, *Biomaterials*, 2001, 22, 471–480.
- 42 A. Akinc, M. Thomas, A. M. Klibanov and R. S. Langer, *J. Gene Med.*, 2005, 7, 657–663.
- 43 Z. Liu, Y. Xiao, W. Chen, Y. Wang, B. Wang, G. Wang, X. Xu and R. Tang, *J. Mater. Chem. B*, 2014, 2, 3480–3489.
- 44 S. Neumann, A. Kovtun, I. D. Dietzel, M. Epple and R. Heumann, *Biomaterials*, 2009, **30**, 6794–6802.
- 45 Y. C. Tseng, A. Yang and L. Huang, *Mol. Pharm.*, 2013, **10**, 4391–4395.
- 46 B. Maury, C. Goncalves, G. Tresset, M. Zeghal, H. Cheradame, P. Guegan, C. Pichon and P. Midoux, *Biomaterials*, 2014, 35, 5977–5985.
- 47 C. Pichon, L. Billiet and P. Midoux, Curr. Opin. Biotechnol., 2010, 21, 640-645.
- 48 G. Creusat, A. S. Rinaldi, E. Weiss, R. Elbaghdadi, J. S. Remy, R. Mulherkar and G. Zuber, *Bioconjugate Chem.*, 2010, 21, 994–1002.
- 49 A. E. Ewence, M. Bootman, H. L. Roderick, J. N. Skepper, G. McCarthy, M. Epple, M. Neumann, C. M. Shanahan and D. Proudfoot, *Circ. Res.*, 2008, **103**, e28–e32.
- 50 Y. Dautova, D. Kozlova, J. N. Skepper, M. Epple, M. D. Bootman and D. Proudfoot, *PLoS One*, 2014, 9, e97565.
- 51 L. Billiet, J. P. Gomez, M. Berchel, P. A. Jaffres, T. Le Gall, T. Montier, E. Bertrand, H. Cheradame, P. Guegan, M. Mevel, B. Pitard, T. Benvegnu, P. Lehn, C. Pichon and P. Midoux, *Biomaterials*, 2012, 33, 2980–2990.
- 52 C. Troiber and E. Wagner, *Bioconjugate Chem.*, 2011, 22, 1737–1752.
- 53 D. Vercauteren, J. Rejman, T. F. Martens, J. Demeester, S. C. De Smedt and K. Braeckmans, *J. Controlled Release*, 2012, **161**, 566–581.
- 54 T. E. Park, B. Kang, Y. K. Kim, Q. Zhang, W. S. Lee, M. A. Islam, S. K. Kang, M. H. Cho, Y. J. Choi and C. S. Cho, *Biomaterials*, 2012, 33, 7272–7281.
- 55 C. Y. Hsu and H. Uludag, *Biomaterials*, 2012, 33, 7834–7848.
- 56 Z. Hyvonen, V. Hamalainen, M. Ruponen, B. Lucas, J. Rejman, D. Vercauteren, J. Demeester, S. De Smedt and K. Braeckmans, J. Controlled Release, 2012, 162, 167–175.
- 57 S. Yamano, J. Dai, C. Yuvienco, S. Khapli, A. M. Moursi and J. K. Montclare, *J. Controlled Release*, 2011, 152, 278–285.
- 58 O. Boussif, F. Lezoualch, M. A. Zanta, M. D. Mergny, D. Scherman, B. Demeneix and J. P. Behr, *Proc. Calif. Acad. Sci.*, 1995, **92**, 7297–7301.

RSC Advances

- 59 R. K. Oskuee, A. Philipp, A. Dehshahri, E. Wagner and M. Ramezani, *J. Gene Med.*, 2010, 12, 729–738.
- 60 C. Lin and J. F. Engbersen, *J. Controlled Release*, 2008, **132**, 267–272.
- 61 C. Pichon, C. Gonçalves and P. Midoux, *Adv. Drug Delivery Rev.*, 2001, 53, 75–94.
- 62 A. Hayer, M. Stoeber, D. Ritz, S. Engel, H. H. Meyer and A. Helenius, *J. Cell Biol.*, 2010, **191**, 615–629.
- 63 C. Goncalves, M. Y. Ardourel, M. Decoville, G. Breuzard, P. Midoux, B. Hartmann and C. Pichon, *J. Gene Med.*, 2009, 11, 401–411.
- 64 A. Mesika, I. Grigoreva, M. Zohar and Z. Reich, *Mol. Ther.*, 2001, 3, 653–657.