Accelerated Achilles tendon healing by PDGF gene delivery with mesoporous silica nanoparticles

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Abstract

We report the ability of amino- and carboxyl-modified MCM-41 mesoporous silica nanoparticles (MSN) to deliver gene in vivo in rat Achilles tendons, despite their inefficiency to transfect primary tenocytes in culture. We show that luciferase activity lasted for at least 2 weeks in tendons injected with these MSN and a plasmid DNA (pDNA) encoding the luciferase reporter gene. By contrast, in tendons injected with naked plasmid, the luciferase expression decreased as a function of time and became hardly detectable after 2 weeks. Interestingly, there were neither signs of inflammation nor necrosis in tendon, kidney, heart and liver of rat weekly injected with pDNA/MSN formulation during 1.5 months. Our main data concern the acceleration of Achilles tendons healing by PDGF-B gene transfer using MSN. Biomechanical properties and histological analyses clearly indicate that tendons treated with MSN and PDGF gene healed significantly faster than untreated tendons and those treated with pPDGF alone.

1. Introduction

Tendons have a crucial role in maintaining the body mechanics, as their main function is to connect muscles to bones. They are axial connective tissues that ensure transmission of the traction exerted by muscle to bone resulting in joint movements. Tendons are composed of a few cells and parallel arrays of collagen fibers. Their important mechanical function renders them liable to injury and rupture. In human, Achilles tendon is submitted to forces greater than 6-fold the bodyweight. Consequently, Achilles tendon injuries are one of the most common tendon disorders and concern a great part of population. They can be caused by different origins such as overuse that often occurs in sports, consequence of metabolic disease (gut) or age-related loss of elasticity. Tendon heals spontaneously but its regeneration is very difficult and requires a very long period - several months, even a year - which is physically disabling. Moreover, the return at the initial state of the biomechanical parameters is never reached.

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A B S T R A C T

Tendon regeneration proceeds in three stages: an inflammatory phase, a cell proliferation phase and a remodelling phase [1,2]. The inflammatory phase that occurs during the first days after the injury is characterized by fibrin clot formation followed by a mobilization of polymorphonuclear leucocytes as well as other cells involved in the inflammation process. These cells play a part in the cellular debris elimination and secretion of growth factors capable of promoting angiogenesis and tenocytes proliferation. During the proliferation phase, the revascularisation is set up and fibroblasts proliferation makes possible the production of extracellular matrix necessary to fill holes induced by lesions. The remodelling phase can last several months up to years. It is characterized by the reduction in the number of cells and a realignment of the matrix to allow a better response to tensions that the tissue can undergo. Unfortunately, healed tendons have altered biomechanical properties because of an increase of the water content and a decrease of the quantity and quality of collagen. So later, ruptures can occur, and sometimes a reduction in traction force (loading capacity) is observed due to the tendon tendency to form adherences [3,4]. These last years, surgery which consists in performing autografts has been improved. However, non-invasive strategies that allow an acceleration of tendon tissues healing are still of interest and remain to be explored.
The mechanism of tendon remodelling and especially the molecular actors that govern its induction and its regulation are known [5]. The process is strongly controlled by growth factors such as bFGF, PDGF (Platelet Derived Growth Factor) and IGF-1 acting at distinct phases of regeneration [6–8]. Indeed, PDGF-BB has a mitogenic activity on fibroblasts and stimulates the collagen I synthesis making it a factor of choice to enhance tendon healing. For instance, a treatment of injured Achilles tendons with PDGF-BB recombinant protein had induced a clear improvement of their regeneration [9]. However, the chronic character of these pathologies requires repeated injections of a large quantity of expensive recombinant. Moreover, the short lifespan of these molecules makes that only a small quantity is delivered to the target and a large part is disseminated in an uncontrolled way causing adverse effects. Thus, the development of alternative methods that will permit a high in situ production of these active molecules, such as gene delivery, is strongly desirable.

Different systems including viral or synthetic vectors can be used to deliver genes into the cells. Viral vectors are the most efficient but still remain less safe than synthetic ones, immunogenic and expensive to produce [10]. These last years, cationic lipids [11], polymers [12], carbon nanotubes [13] and silica nanospheres [14] have been extensively exploited as chemical vectors for delivering genes. Several studies have reported the use of nanoparticles as drug or gene delivery systems with a low toxicity, improved body distribution, high biocompatibility and improved efficacy [14–18].

MCM-41 mesoporous silica are inorganic materials that have a porous structure with hundreds of empty channels (mesopores) able to adsorb or encapsulate relatively large amounts of bioactive molecules. They have high surface area (>900 m²/g), a large pore volume (>0.9 cm³/g) and pore size around 2–6 nm. Mesoporous silica nanoparticles (MSN) have been used as in vitro DNA vectors [14].

Here, we have used MSN to transfect in vivo rat Achilles tendon. For this purpose, MSN surface was modified with amino or carboxyl groups.

2. Experimental procedures

2.1. Materials

3-aminopropyltriethoxysilane (APTES, 99%, Sigma–Aldrich, Saint Quentin Fallavier, France), toluene (for HPLC, 99.8%, Carlo Erba, Val de reuil, France), succinic anhydride (99%, Aldrich), anhydrous dimethylsulfoxide (DMSO, Sigma–Aldrich) and absolute ethanol (99.7%, Carlo Erba) were used as received. p-formaldehyde, glutaraldehyde, potassium hexacyanoferrate (III) and lead citrate were obtained from Electron Microscopy Sciences (Hatfield, USA).

2.2. Plasmids

pNFCMV-luc was a homemade plasmid DNA of 7.5 kb that encodes the NF κB motifs (termed NF) that recognize the NF κB binding sites. Note that human and rat PDGF-B share more than 90% of identity. Super coiled pDNA was isolated from bacteria by the standard alkaline lysis method, and purification was carried out with the Qiagen Mega Kit (Qiagen, Courtaboeuf, France).

![Functionalization scheme of MCM-41. APTES: 3-aminopropyltriethoxysilane.](image)

### Table 1

<table>
<thead>
<tr>
<th>Zeta potential (mV)</th>
<th>Without pDNA</th>
<th>With pDNA</th>
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</thead>
<tbody>
<tr>
<td>MSN</td>
<td>−38.5</td>
<td>−</td>
</tr>
<tr>
<td>MSN-NH₃</td>
<td>+17.9</td>
<td>−33.4</td>
</tr>
<tr>
<td>MSN-COOH</td>
<td>−49.6</td>
<td>−35.2</td>
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Zeta potential measurements of amino- or carboxyl-modified MCM-41 silica particles before and after mixing with pDNA. All measurements were performed in 10 mM Hepes buffer, pH 7.4.

2.3. Surface modification of mesoporous silica nanoparticles

MCM-41 type MSN was synthesized as described by Lelong et al. [19]. The silica surface was substituted with amino groups by reaction at room temperature for 12 h of 73 mg of MSN with 2.5 mL of 3-aminopropyltriethoxysilane (APTES) in anhydrous toluene. Then, amine-functionalized mesoporous silica materials (MSN-NH₂) were filtered, washed with toluene and dried at 60°C for 12 h under vacuum. MSN-COOH particles were prepared as follows. Firstly, 40 mg of MSN-NH₂ were added to 20 mL of toluene and the solution was kept at 50°C until solubilisation. Then, 150 mg of succinic anhydride was allowed to react overnight at room temperature. MSN-COOH particles were separated from the reaction medium by filtration, washed several times with ethanol and then vacuum-dried. The different MSN were characterized by Fourier transform infrared (FTIR) and zeta potential measurements.

2.4. FTIR experiment

FTIR spectra were collected on a Nicolet 710 FTIR spectrophotometer with a resolution of 8 cm⁻¹ by using the KBr method.

2.5. Zeta potential measurement

Measurements were carried out by electrophoretic mobility using a Zeta Sizer 3000 (Malvern Instruments, Orsay, France). The system was calibrated with the DTS 5050 standard beads from Malvern. Measurements were done 5 times with the zero field correction and performed in 10 mM Hepes buffer, pH 7.4, with the following parameters: viscosity, 0.891 cP; dielectric constant, 79, temperature, 25°C, F(Ka), 1.50 (Smoluchoffsky), maximum voltage was 15 V. The zeta potential was calculated using the Smoluchoffsky approximation; model Z (Malvern Instrument, Worcestershire, England).

2.6. pDNA/MSN formulations

MSN nanoparticles at 10 μg/ml in distilled water were sonicated for 30 min before use. They were then added to 150 μl NaCl aqueous solution containing pDNA, and the mixture was kept for 30 min at room temperature before transfection. pDNA/MSN formulations (50 μl final volume) were prepared at different weight ratios (μg/μg) from 2:0.6 to 2:5. Formulations were freshly prepared before use.

2.7. Transmission electron microscopy analysis

pDNA/MSN formulations were dropted onto the 400 mesh carbon surface of electron microscopy copper grids. After 2 min, the grids were stained for 1 min with 5% uranyl acetate and 0.4% lead citrate and blotted dry. Samples were analysed using a transmission microscope CM20 (Philips) at 200 kV operating voltage.

2.8. Atomic force microscopy (AFM)

The morphology of the pDNA/MSN was analysed by atomic force microscopy (Molecular Imaging Pico +). The samples were prepared by mixing 0.1 μg of pDNA with various amounts of MSN. After 30 min incubation, samples were dropped on freshly cleaved mica surface and allowed to interact for 5 min and then blotted dry.
2.9. Agarose gel electrophoresis

MSN were mixed with pDNA at indicated weight ratio in 50 μl of 10 mM Hepes, pH 7.4. pDNA/MSN or pDNA alone were run on 0.8% agarose gel containing 0.6X ethidium bromide for 3 h at 80 V. Images were taken using a UV transilluminator (Geneflash, Syngene Bio Imaging).

2.10. Primary culture of tenocytes

Tenocytes were obtained from adult Achilles tendon explants. Adult Wistar rats (average weight of 250 g, from Etablissement Dupré (Saint-Doulchard, France) were sacrificed and their tendons were harvested and cut in small pieces in Hank’s Buffered Salt Solution pH 7.5 (HBSS). Then, small tendon pieces were placed in tenocytes’ medium (DMEM 1% glutaframe, 10% Fetal Calf Serum, 50 μg/ml ascorbic acid, 250 U/ml penicillin, 250 μ/mL streptomycin, 0.2% fungizone) and cultured at 37°C under humidified atmosphere containing 5% CO₂. After 10 days, first primary cells sprout out from the explants. They were harvested with trypsin, reseeded in 75 cm² flask and maintained in tenocytes’ medium. Half of the medium volume was replaced every two days.

2.11. In vitro transfection

Two days before experiment, tenocytes were seeded at 2 × 10⁴ cells on 2 cm² well. Then, tenocytes were transfected with 2.5 μg pNCFMV-luc mixed with MSN as described above. After 4 h, the medium was removed and cells were cultured for two days in tenocytes’ medium. The transfection efficiency was determined by measuring the luciferase activity in cell lysates as follows. The culture medium was removed and cells were lysed on ice during 10 min in lysis buffer (1% Triton X-100, 25 mM Tris-phosphate, 1 mM DTT, 1 mM EDTA, 15% glycerol, 1 mM MgCl₂, pH 7.8). In vitro luciferase substrate (beetle luciferin, Promega, Madison USA) was added to lysate supernatant in the presence of 1 mM ATP. The Relative Light Units (RLU) produced was measured with a luminometer (Lumat LB9507, Berthold, Thoiry France) and expressed as RLU per mg of proteins upon protein quantification using the BCA colorimetric assay.

2.12. In vivo transfection

Adult Wistar rats were stored in CBM’s animal house at 22°C and fed ad libitum for at least one week before experiments. Rats were anesthetized either with a mixture of ketamine (250 mg/kg) and xylazine (10 mg/kg) in 0.9% NaCl by intra-peritoneal injection or by isoflurane gas system (AErrane, Baxter, Maurepas, France). Rats’ posterior legs were washed with Vetedine® (Centravel, Allfort, France) and shaved. Skin was incised on few millimeters in Achilles tendon region with a stainless steel surgical blade n° 12 (Swann-Morton). Achilles tendon was then harwood along its total length three times using a carbon steel surgical blade n° 12 (Swann-Morton). Fifty microliters of pDNA/MSN at a weight ratio of 2:5 were slowly injected in the middle section of Achilles tendon using a Hamilton PBS600 repeater delivery system (5 μl dose per injection). Finally, skin was sutured and rats were closely observed until their wake up. After surgery, animals were treated with subcutaneous injection of lynadine (250 μL/kg) to relieve animals’ pain.

The transfection efficiency of pNCFMV-luc treated rats was determined by measuring the luciferase activity in tendon as follows. Rats were euthanized by lethal CO₂ inhalation at day 1, 4, 6, 10 and 15 after surgery. Achilles tendons were harvested and maintained in PBS on ice. Tendons were crushed in liquid nitrogen with mortar and pestle. The powder was added in ice cold lysis buffer (luciferase Cell Culture Lysis, Promega, Madison USA) and the mixture was kept for 3 h on ice before luminescence measurement. The Relative Light Units (RLU) was measured with a luminometer after adding luciferase substrate (Luciferase Assay System, Promega, Madison USA) to tendon lysates and expressed as RLU per mg of proteins.

2.13. In vivo toxicity assessment

Experiments were carried out with 5 rats per group. Each group received a weekly injection during 6 weeks. The first group was injected with saline solution (150 μl NaCl), the second one with 50 μg MSN and the third one with 20 μg pDNA mixed with 50 μg MSN. Transfected rats were euthanized by lethal CO₂ inhalation. Liver, spleen, kidneys, tendons, and mesenteric lymph nodes were harvested and fixed in 10% p-formaldehyde at room temperature. Then organs were embedded in paraffin wax (Tissue Tek, Sakura Bayer Diagnostics), sliced in serial sections and stained with HES (hematoxyline eosine safranin) (Tissue Tek) and toxicity analyses were performed by Novaxia (St Laurent Noun, France). Each slide was inspected microscopically by a pathologist to search for any evidence of damage such as adhesions, necrosis, cellular infiltration or hypervascularity.

2.14. Biomechanical tests

Control and transfected rats were euthanized by lethal CO₂ inhalation at indicated times. Achilles tendons were harvested and stored in PBS at -20°C before use.

They were carefully dissected and cleaned to remove muscle tissues. The tendon extremities were glued with cellulose polyanacetate and care was taken to prevent gel spreading over the tendon. Then the tendon was clamped between two metal jaws covered with rubber, and mounted vertically on a conventional mechanical test machine (MTS synerijge 400). To strengthen the interface between the sample and the jaw, the tendon was surrounded with a piece of rubber. The tendon was pre-loaded with a velocity of 0.6 mm/min until the applied force reaches 0.1 N. The length (L0) between both extremities of the tendon was measured and the tendon was preconditioned by a cyclic loading, stretching the tendon to 110% L0 with a velocity of 0.6 mm/min. After pre-conditioning, specimens were stretched to failure at the same rate and the force to failure was recorded (Max. load). The maximal stress (Max. stress) was measured by the force divided by the tendon surface. In addition the Young’s modulus (E) was also determined.

After mechanical tests, the tendons were fixed in fresh Bouin’s picroformol solution for up to 48 h. Thereafter, tendons were embedded in paraffin and cross sections of 5 μm thickness were cut and stained with Van Gieson’s trichrome. The surface of each tendon was manually evaluated from tendon sections mounted on glass slides and analysed by optical microscope using QWIN Leica software.

2.15. Histological analysis

To assess the healing effect of PDGF-B gene transfer, control and injured tendons were harvested at indicated times. They were fixed in 10% p-formaldehyde solution, paraffin embedded, sliced in serial sections (thickness: 4 μm), mounted on glass slides and counterstained with HE stain. Experiments were carried out and analysed by LeicaQ5 (Nantes, France).

Fig. 2. Fourier-transform IR (FTIR) absorbance spectra of MSN and modified MSN.

Fig. 3. pDNA/MSN interactions MSN-COOH and MSN-NH₂ were mixed with pDNA at a pDNA/MSN weight ratio (μg / μg) of 2/0.6 (line 1), 2/1.25 (line 2 & 5), 2/2.5 (lines 3 & 6) and 2/5 (lines 4 & 7). Line C corresponds to pDNA alone. Agarose gel electrophoresis was carried out and DNA was revealed by Ethidium Bromide fluorescence upon DNA intercalation. Arrow: retarded DNA.
2.16. Statistical analysis

Data were expressed by mean ± SD or SEM as indicated. Statistical differences were analysed by a unilateral Mann–Whitney U-test using XLStat 2007 software, and significance was defined as p-value <0.05.

3. Results

3.1. MSN surface modifications

MCM-41 mesoporous silica nanoparticles (MSN) were functionalised with amino or carboxyl groups as described in Fig. 1. The global charges of MSN and modified MSN were assessed by measuring their zeta potential at different steps of functionalization procedure (Table 1). Untreated MSN carried a net negative charge of −38.5 mV due to the ionisation of the surface hydroxyl groups. In contrast, the amine modified ones (MSN-NH₂) had a net positive surface charge of 18 mV as a consequence of the presence of amino groups on their surface. MSN-COOH exhibited a negative zeta potential of −49.6 mV resulting from the carboxyl groups created by reaction of amine groups of MSN-NH₂ with succinic acid molecules.

Silica surface modification was also monitored systematically through the different stages by FTIR spectroscopy (Fig. 2). The unmodified MSN showed the typical vibration bands of siliceous

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**Fig. 4.** TEM and AFM analyses. A. Representative Transmission Electron microscopy images of naked plasmid DNA (a), MSN-NH₂ (b), MSN-COOH (c), pDNA/MSN-NH₂ (d), pDNA/MSN-COOH (e). B. Intermittent mode AFM images of nanoplexes deposited on mica surface by blotting. Left: DNA/MSN-COOH 2:2.5 Right DNA/MSN-NH₂ 2:5. Images suggest that most nanoparticles are linked to DNA. When MSN were not mixed to DNA they did not bind to the mica surface, which confirms the attractive interaction between MSN and DNA.
materials, such as that of asymmetric stretching Si–O–Si at ca. 1085 cm⁻¹, symmetric stretching Si–O–Si at ca. 800 cm⁻¹, and stretching vibrations of Si–OH groups at ca. 960 cm⁻¹ (Bands located at 1630 and 3430 cm⁻¹ come from adsorbed water). The grafting of APTES in MSN-NH₂ samples was confirmed by the presence of adsorption bands at ca. 2900 cm⁻¹ attributed to C–H stretching vibrations from propyl groups, and a broad band near 3300 cm⁻¹ due to N–H stretching. After reaction with succinic anhydride, amide II band emerges near 1450 cm⁻¹, and amide I band increases intensity and width of 1650 cm⁻¹ peak.

3.2. Interactions between DNA and silica nanoparticles

MSN-NH₂ and MSN-COOH were mixed with a pDNA at various weight ratio and their interactions were investigated by zeta potential measurements and agarose gel electrophoresis. When mixed with pDNA, the global charge of MSN-NH₂ became negative (from +17.9 mV to −33.4 mV). This result suggests that electrostatic interactions occurred between the amino charges of MSN-NH₂ and the phosphate groups of pDNA (Table 1). In the case of MSN-COOH, the zeta potential in the presence of pDNA remained negative (−35.2 mV). Since, the number of amino groups of MSN-COOH was low compared to MSN-NH₂, pDNA was not expected to interact strongly with MSN via electrostatic interactions (Table 1). The interactions were also evidenced by agarose gel electrophoresis (Fig. 3). Whatever the amount of the two MSN types, the migration of pDNA was not completely retarded indicating that the interaction strength was weak. Only one part of pDNA was completely retarded with both MSN types and the amount increased with the pDNA/MSN weight ratio suggesting the presence of some pDNA/MSN complexes. Note that in the presence of MSN up to 100 μg, MSN/DNA interactions were not improved (not shown). TEM and AFM observations were performed to assess MSN/DNA association (Fig. 4). TEM micrographs of silica nanoplexes made with the two types of MSN showed that nanoparticles are surrounded by loose strands of DNA. However, TEM and AFM showed that a significant amount of DNA remains unbounded.

According to agarose gel electrophoresis, TEM and AFM observations, pDNA was found to interact weakly with two types of MSN. It seems that the nature of the charge or functional group carried by the MSN did not influence their capacity to interact with pDNA. The data suggest that pDNA may enwrap MSN via surface adsorption with some strand inserted inside pores. Indeed, MSN has a honeycomb-like structure with mesopores large enough to absorb a high amount of biomolecules [17]. Note that the 3 nm pore size of our MSN is compatible with the diameter of a double stranded DNA molecule.

3.3. In vitro transfection efficiency of tenocytes

In vitro transfection efficiency of primary tenocytes was evaluated with a plasmid DNA (pDNA) encoding luciferase reporter gene (pLuc) in the presence of various ratios of MSN-NH₂ or MSN-COOH nanoparticles and Jet-PEI™ as gold standard transfection reagent (Fig. 5). The transfection efficiency with MSN-NH₂ and MSN-COOH at pDNA/MSN weight ratio of 2:5 and 2:10 was very low and similar to that obtained with naked pLuc. Comparatively, the luciferase activity produced with Jet-PEI™ polyplexes was 100-fold higher than with bare pDNA. We found that these two types of MSN/pDNA formulations were also unable to transfet other cell types including human embryonic kidney (HEK293) cells, known to be easily transfected by most of transfection reagents (not shown). This absence of efficacy on cultured cells may be due to the weak interactions of pDNA with both MSN. It is known that pDNA compaction is usually required to avoid its degradation by serum nucleases in culture medium and to enhance its cellular uptake. Moreover, the negative zeta potential of pDNA/MSN probably did not favour their internalization by the cells. Indeed, when the tenocytes were incubated with MSN formulated with fluorescein-labelled pDNA, flow cytometry analysis showed that there was no difference between the cell-associated fluorescence intensities measured after 4 °C and 37 °C incubation (not shown). Accordingly with the above data, this result evidenced a very low pDNA uptake by tenocytes, and therefore it is not surprising that transfection was not efficient.

3.4. Achilles tendons transfection

In spite of the absence of transfection of primary tenocytes, we tested the efficiency of these pDNA/MSN upon direct injection in rat Achilles tendons. In the first set of experiments, tendons were transfected with pLuc, and 3 days post-transfection the luciferase activity was evaluated. Surprisingly, the two MSN types were able to transfect efficiently tendons at a pLuc/MSN weight ratio of 2:2.5
The efficacy at a pLuc/MSN weight ratio of 2:5 was not significantly different (not shown). It is worth to note that similar transfection efficacy was obtained on injured or healthy tendons (not shown). Although MSN-NH₂ was as efficient as MSN-COOH, we decided to use the latter because we thought that anionic silica nanoparticles will be less toxic than the amino ones and would lead to a better clearance. Indeed, it is known that positive charged particles could be stuck within negatively charged extracellular matrix components. Next, dose response experiments were carried out with 2.5, 5, 10 and 20 μg of pLuc formulated with MSN-COOH in tendons. Fig. 6 indicates that the gene expression was dependent on pLuc dose and a plateau was reached at 10 μg pDNA (1.6 × 10⁸ RLU/mg of proteins).

The kinetics of gene expression was performed with 5 μg pDNA (Fig. 7). One day post-transfection, the luciferase activity obtained from lysates of tendons transfected with pLuc/MSN-COOH was high (10⁷ RLU/mg of proteins) but not significantly different to that obtained with pLuc alone. Then, this activity decreased the following days but not with the same manner. At day 10 and day 15 post-injection, the luciferase activity in pLuc/MSN-COOH nanoplexes-injected tendons detected was 10 and 100-fold less than that measured at day 3, respectively. Comparatively, in tendons injected with pLuc alone, this activity was 114-fold and 1500-fold lower than that at day 3. Similar differences were also observed when 20 μg pDNA was injected in tendons.

### 3.5. MSN toxicity

Repeated injections (one per week during 6 weeks) of either free MSN-COOH or pDNA/MSN were done locally in rat Achilles tendons. Rats were checked up every day for close observations, and vital organs (liver, spleen, kidney, mesenteric lymph node and lungs) as well as tendons were harvested after 6 weeks for histopathological analyses. As shown in Fig. 8, no histopathology changes in terms of inflammation, necrosis or structural tissue organization were observed in vital organs of treated groups versus untreated ones. It is worth to note that no signs of inflammation were observed in tendons after 3 days post-injection.

### 3.6. Healing activity of PDGF gene transfer

The absence of toxicity upon pDNA/MSN transfection prompted us to evaluate their efficacy to transfer the PDGF-B gene for tendon healing. Rat Achilles tendons were injured and then treated or not with plasmid encoding PDGF gene (pPDGF). For that, 20 μg of pPDGF were used alone or mixed with MSN-COOH. In control rats, healthy tendons were injected with physiological serum or with a plasmid encoding luciferase gene. The beneficial effect of the treatment was assessed by biomechanical tests and tendon histology. In terms of biomechanical properties, parameters defined by maximal load to failure, maximum stress and Young’s modulus were significantly greater in pPDGF/MSN or in naked pPDGF-treated groups than in untreated one (Table 2). It is worth to note that there is a high variability of Young’s modulus values of...
injured tendons treated with pPDGF alone (88.79 ± 16.39 MPa) which is not the case of tendons treated with pPDGF/MSN. Remarkably, the values of maximal load, maximal stress and Young’s modulus of tendons treated with pPDGF/MSN are very close to those of healthy (control) tendons (96.57 ± 7.7 vs 96.25 ± 4.19 MPa).

To improve the organization of tendon structure, we treated injured tendon with two injections of pPDGF spaced by one week interval. Histological analyses at 21 days post-injury showed that a faster regenerating activity occurred in treated tendons than in the untreated groups (Fig. 9). This was supported by the following observations: i) a higher number of fibroblasts was observed in untreated tendons than in tendons treated with pPDGF or pPDGF/MSN indicating that untreated tendons were still in proliferative phase after 2 weeks of injury and ii) treated tissues were nicely organized by contrast to those that were mock treated. More importantly, tendons that have been treated with pPDGF/MSN exhibited the highest structural organization with more aligned fibers compared to tendons treated with pPDGF alone.

4. Discussion

Several reports have shown that silica nanoparticles can be used as carrier to deliver gene in vitro [14]. Silica nanoparticles are usually modified by various means to exhibit positive charges providing electrostatic binding with negatively charged DNA. In the present study, we provide evidences that mesoporous silica nanoparticles, substituted either with amino or carboxyl groups, can be used in vivo for gene delivery in rat Achilles tendons despite being inefficient in vitro on primary cultures of tenocytes.

Both pDNA/MSN-NH₂ and pDNA/MSN-COOH nanoplexes exhibited a negative zeta potential of about −35 mV at pH = 7.4. This suggests that direct electrostatic interaction is not the main driving force for association of pDNA on MSN. It is probable that hydrogen bonding between silanol and phosphate groups plays an important role in pDNA-MSN interaction [20,21]. Our results could also suggest that segments of pDNA were inserted inside MSN pores. In the case of MSN-NH₂, a propyl group was used as spacer between the surface of MSN and the amino group. This was also the spacer used to prepare ORMOSIL amino silica nanoparticles [22]. In this case, pDNA/ORMOSIL interactions were also weak. Indeed, pDNA was not completely retarded under agarose gel electrophoresis. The propyl length may be too short to allow strong electrostatic interactions with pDNA due to screening effects. Indeed, a complete pDNA retardation was obtained at a pDNA/MSN weight ratio of 1:5 (μg:μg) when a hexyl-amino-propyl spacer was used [23] as well as when MSN was capped with polyamidoamine dendrimer [24]. The absence of strong interactions between MSN and pDNA as well as the negative zeta potential of pDNA/MSN explain the absence of efficient transfection of cultured tenocytes.

### Table 2

<table>
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<tr>
<th>Populations</th>
<th>Maximal load (N)</th>
<th>Maximal stress (MPa)</th>
<th>Young’s modulus (MPa)</th>
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<td>pPDGF (n=8)</td>
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<td>Untreated (n=12)</td>
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<td>20.18 ± 2.62</td>
<td>68.15 ± 7.63</td>
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Maximal load, maximal stress and Young’s modulus obtained with the mechanical tests: control: healthy tendons, pPDGF: injured tendons injected with pPDGF alone, pPDGF/MSN: injured tendons treated with pPDGF/MSN—COOH, Untreated: injured tendons treated with saline. Values are mean ± standard error of the mean (SEM).

a Significant statistical difference (p < 0.05) compared to untreated group.

b No significant statistical difference compared to healthy group.

Fig. 9. Histological analyses of pDNA/MSN-treated Achilles tendons. Injured Achilles tendons were treated with pPDGF alone or formulated with MSN-COOH. Twenty days post-treatment healthy and treated tendons were harvested, fixed in 10% p-formaldehyde solution, paraffin-embedded, sliced in serial sections (4 μm), mounted on glass slides and HES counterstained. Images shown are representative sections (10 fold magnification) of at least 4 tendons from healthy tendon (A), non-treated tendon (B), pPDGF-treated tendon (C) and pPDGF/MSN-treated tendon (D). The delineated circle corresponds to injury area.
as well as other cell lines in vitro. This result is coherent with previous reports clearly indicating that for in vitro transfection, positive complexes are required to get enhanced interactions with plasma membrane and high enough protection against extracellular degradation. In the case of silica nanoparticles, Kneuer et al. [18] have shown that a high weight ratio of silica pDNA/nanoparticles (1:30) was needed to get positive charged nanoplexes efficient for in vitro transfection. The use of such large amount of silica nanoparticles could induce biological side effects (see below [25]). Of interest, our MSN are able to transfer gene efficiently in vivo in tendons. Such behavior has also been found for amphiphilic block copolymers such as Lutrol or Pluronic that are not efficient to transfect cells in vitro whereas they show high efficacy to transfect skeletal muscles upon intramuscular injection or lung upon intratracheal instillation [26–28]. In our case, no transfection was observed upon intramuscular injection of pDNA/MSN formulations (data not shown). The probable explanation is that pDNA/MSN were able to diffuse upon injection in tendons allowing the transfection of a large area whilst they stuck at the site of injection and did not diffuse at all upon intramuscular injection (Pichon et al., unpublished observations).

Data concerning the kinetics of gene expression reveal the gradual diminution of the luciferase activity that can likely be contributed to the time release behavior of pDNA adsorbed on or entrapped in MSN.

Our main data concerns the acceleration of the regeneration activity by PDGF gene transfer with MSN. Tendons are viscoelastic tissues and therefore their internal organization mainly based on collagen I is crucial [29,30]. In rat Achilles tendocytes, Wang et al. have shown that in vitro transfection of PDGF-B gene promotes a positive effect on collagen I production [31]. Note that, the level of luciferase produced in tendons upon transfection was close to the nanogram range). Our data clearly show a fast recovery of biomechanical property of pPDGF/MSN treated tendons which is sustained by the active dose of recombinant PDGF-BB (related to the nanogram of luciferase produced in tendons upon transfection was close to the nanogram of luciferase produced in tendons upon transfection). Our data also show an increased recovery of biomechanical property of pPDGF/MSN treated tendons which is sustained by an active dose of recombinant PDGF-BB (related to the nanogram of luciferase produced in tendons upon transfection). Our data also show an increased recovery of biomechanical property of pPDGF/MSN treated tendons which is sustained by an active dose of recombinant PDGF-BB (related to the nanogram of luciferase produced in tendons upon transfection).

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The adverse biological effects of nanomaterials such as silica nanoparticles raise safety concerns on their use. Several reports have assessed the biological side effects that could induce silica nanoparticles. They have shown that silica nanoparticles can induce inflammatory (NFκB activation) and oxidative stress responses both in vivo and in vitro [32–35] but these effects were only observed at high concentrations [25,36]. Recent studies have demonstrated by comet assay that silica nanoparticles with size varying from 20 to 400 nm do not exert significant genotoxicity [25,36]. The toxicity of our pDNA/MSN was addressed by investigating the chronic effect they could induce. Gross observations of tendon and histological analyses indicated that no local inflammation or necrosis was induced after several injections. Note that no acute inflammatory process was observed after one or 3 days post-injection (not shown). We are, however, conscious that before human application, a battery of standardized tests to quantify genetic aberrations must be performed to cover all potential forms of DNA damage that may be induced following the local injection of silica nanoparticles in Achilles tendons.

5. Conclusions

Results reported in this study clearly demonstrate that mesoporous silica nanoparticles can be exploited to efficiently deliver genes in Achilles tendons. The delivery of pDNA with MSN allows a sustained gene expression for at least 2 weeks. Our remarkable result concerns the improvement of healing of injured tendons by PDGF gene transfer. Nevertheless, co-transfection with other genes encoding active molecules involved in healing will be addressed to restore a tendon structure as aligned as in healthy tendons.

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Appendix

Figures with essential color discrimination. Figs. 2, 4, 8 and 9 in this article are difficult to interpret in black and white. The full color images can be found in the on-line version, at doi:10.1016/j.biomaterials.2010.02.077.

References
