E2F decoy oligodeoxynucleotide ameliorates cartilage invasion by infiltrating synovium derived from rheumatoid arthritis

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Abstract. This study examined the ability of E2F decoy oligodeoxynucleotides (ODN) to inhibit proliferation of synovial fibroblasts derived from patients with rheumatoid arthritis (RA). The effect of E2F decoy ODN on cartilage invasion by RA synovium in a murine model of human RA was also investigated. E2F decoy ODN were introduced into synovial tissue and synovial fibroblasts derived from patients with RA using hemagglutinating virus of Japan (HVJ)-liposomes. The effect of E2F decoy ODN on synovial fibroblast proliferation was evaluated by MTT assay and by RT-PCR for the cell cycle regulatory genes proliferating-cell nuclear antigen (PCNA) and cyclin-dependent kinase 2 (cdk2). Changes in production of inflammatory mediators by RA synovial tissue following transfection with E2F decoy ODN were assessed by ELISA. Human cartilage and RA synovial tissue transfected with E2F decoy ODN were co-transplanted in severe combined immunodeficient (SCID) mice. After 4 weeks, the mice were sacrificed and the implants histologically examined for inhibition of cartilage damage by E2F decoy ODN. E2F decoy ODN resulted in significant inhibition of synovial fibroblast proliferation, corresponding with reduced expression of PCNA and cdk2 mRNA in synovial fibroblasts. The production of interleukin-1β (IL-1β), IL-6 and matrix metalloproteinases (MMP) such as IL-1, IL-6, TNF-α, MMP-1 and MMP-3 (2), and expression is clearly apparent at cartilage-pannus junctions (3). The importance of proliferative activity in RA and its association with production of proinflammatory cytokines has been studied (4-6). Therefore, cell cycle regulators are attractive candidates for therapeutic targets to halt joint destruction in RA.

Introduction

Rheumatoid arthritis (RA) is characterized by synovial hyperplasia with infiltration of various inflammatory cells resulting in invasion of articular cartilage and bone (1). Synovial fibroblasts are thought to play an important role in the pathogenesis of joint destruction in the arthritic joints. Synovial cells are the major source of proinflammatory cytokines and matrix metalloproteinases (MMP) such as IL-1, IL-6, TNF-α, MMP-1 and MMP-3 (2), and expression is clearly apparent at cartilage-pannus junctions (3). The importance of proliferative activity in RA and its association with production of proinflammatory cytokines has been studied (4-6). Therefore, cell cycle regulators are attractive candidates for therapeutic targets to halt joint destruction in RA.

The transcription factor E2F regulates the expression of multiple cell-cycle regulatory genes that are critical to cell growth and proliferation. In G1/S phase, E2F forms an inactive complex with the hypophosphorylated retinoblastoma (RB) gene product, cyclin A and cdk2. In this condition, the transcriptional activity sequestered E2F is repressed. Once RB is phosphorylated, E2F is released and becomes free to bind to a specific cis element in the promoter region of cell cycle regulatory genes c-myc, c-myb, cdc2, and cdk2 and proliferating cell nuclear antigen (PCNA), thereby trans-activating the expression of these genes (7-10). Cell cycle biology involves a complex interaction of multiple growth factors, their receptors, secondary messengers, oncogenes and transcriptional factors. Therefore, the transcriptional factor E2F provides a good single target for cell cycle blockade.

It has been demonstrated that a synthetic double-stranded oligodeoxynucleotide (ODN) with high affinity for a target transcription factor may be introduced into target cells as a ‘decoy’ to bind the transcription factor, thereby altering gene transcription (11). We have previously demonstrated inhibition of synovial cell proliferation in vitro and amelioration
of joint damage in vivo using NFkB decoy ODN (12). To determine the utility of cell cycle inhibition in treating RA, the present study examined the ability of E2F decoy ODN to inhibit synovial proliferation and production of proinflammatory mediators. We also examined the effects of E2F decoy ODN on cartilage invasion by synovial tissue derived from RA patients.

Materials and methods

Patients. Synovial tissues were obtained from five patients with RA who were undergoing synovectomy at Osaka University Hospital and affiliated facilities after receipt of informed consent. All the patients were diagnosed clinically with RA according to the 1987 revised diagnostic criteria of the American College of Rheumatology (13). Normal synovial tissues were obtained from three patients who were seen for trauma and had no evidence of arthritis.

Synovial cell preparation. The synovial specimens were finely minced into small pieces, soaked in an enzyme cocktail solution containing 0.1% type IV collagenase, 0.1% hyaluronidase, and 0.01% DNase (all from Sigma Chemical Co., St. Louis, MO), and incubated for 2 h at 37˚C in a shaking water bath. After removal of debris by filtration, the cells thus obtained were suspended in Dulbecco’s modified Eagle’s medium (DMEM), washed twice, resuspended in DMEM with 10% fetal calf serum (FCS), and seeded in culture dishes. After overnight culture, non-adherent cells were removed, while adherent cells were re-cultured. Third passage synovial cells were used in the experiments.

Cell proliferation assay. Synovial cells were seeded on to uncoated 24-well tissue culture plates (Corning Inc., Corning, NY) at 4000 cells/well. The cells were then incubated in DMEM with 10% FCS for 48 h. After transfection of decoy ODN, the medium was changed to fresh DMEM with 10% FCS. Four days after transfection, an index of cell proliferation was determined by using sulphonated tetrazolium salt, and a compound that produces a highly water-soluble formazan dye, which makes the assay procedure easier to perform.

Synthesis of ODN and selection of sequence targets. The sequences of phosphorothioate double-stranded ODN against the E2F-binding site and of scrambled ODN used in this study were reported previously (15). The phosphorothioate ODN utilized in this study had the following sequences:

E2F decoy ODN: 5’-CTAGATTTCGCCG-3’
3’-TAAAAGGCCGCTTAG-5’

Scramble decoy ODN: 5’-CTAGATTTCGCCG-3’
3’-TAAAAGGCCGCTTAG-5’

The E2F ODN has been shown to bind the E2F transcription factor (11,15-17). Synthetic ODN were washed in 70% ethanol, dried and dissolved in sterile Tris-ethylene diamine tetra acetic acid (EDTA) buffer (10 mM Tris, 1 mM EDTA). The supernatant was purified over a nucleic acid purification-10 (NAP-10) column (Pharmacia LKB Biotechnology, Piscataway, NJ), and the ODN concentration was quantitated by spectrophotometry. Single-strand ODNs were annealed for 2 h while gradually cooling from 80 to 25˚C.

Transfection using HVJ-liposome method. Phosphatidylserine, phosphatidylcholine, and cholesterol were mixed in a weight ratio of 1:4:8:2 (12,14,18,19). The lipid mixture (10 mg) was deposited on the sides of flask by removal of tetrahydrofuran in a rotary evaporator. Dried lipid was hydrated in 200 ml of balanced salt solution (BBS; 137 mM NaCl, 5.4 mM KCl, 10 mM Tris-HCl, pH 7.6) containing synthetic double-stranded ODN. Liposomes were prepared by shaking and sonication. Purified HVJ (Z strain) was inactivated by ultraviolet irradiation (110 erg/mm²/sec) for 3 min just before use. The liposome suspension (0.5 ml, containing 10 mg of lipids) was mixed with HVJ (10,000 haemagglutinating units) in a total volume of 4 ml of BBS. The mixture was incubated at 4˚C for 10 min and then for 60 min with gentle shaking at 37˚C. Free HVJ was removed from the HVJ-liposomes by sucrose density gradient centrifugation. The top layer of gradient containing purified HVJ-liposomes was collected for use. Synovial tissues were cultured in a serum-free medium 6 h prior to the transfection, then washed 3 times with BBS containing 2 mM CaCl₂. The HVJ-liposome complex (15 mM of encapsulated ODN) was added to the synovial tissues for 30 min at 37˚C. Finally, fresh medium containing 10% FCS was added to the synovial tissues, which were then incubated in a CO₂ incubator.

Estimation of the transfection efficiency. To examine the localization of the transfected FITC-labeled ODN, cryostat sections of synovium transfected with decoy ODN were prepared for fluorescence microscopy. The sections were stained with Hoechst 33342 (Sigma Chemical Co.) and observed under an ultraviolet laser scanning confocal microscope (PCM 2000; Nikon, Tokyo).

RNA extraction and RT-PCR. Twenty-four hours after transfection of E2F decoy ODN, RNA was extracted from synovial tissues by means of RNAzol (Tel-Test Inc., Friendswood, TX). Expression of PCNA, cdk2, and β-actin mRNA were measured by RT-PCR as described previously (11). Total RNA (1 μg) prepared from synovial fibroblasts was first treated with RNase-free DNase. After treatment for 5 min at 94˚C, the samples were subjected to reverse transcription using random hexamer primers (Perkin-Elmer Cetus, Norwalk, CT) and Molony murine leukemia virus reverse transcriptase. The primers for PCNA, cdk2, and β-actin genes used in this study were: The PCNA 5’ primer, 5’-ACTCTGCGGC TCCGAGG-3’; the PCNA 3’ primer, 5’-TCTCCACA ATTAGGCTAAG-3’. The cdk2 5’ primer, 5’-CGCTTC ATGAGAAGCT-3’; the cdk2 3’ primer, 5’-ATGGCA GAAAGCTAGG-3’. The β-actin 5’ primer, 5’-TTGTAA CCAACTGGGAGCATA GG-3’; the β-actin 3’ primer, 5’-GATCCTGATTT CATTCTGGCT-3’. Aliquots of RNA were amplified simultaneously by PCR (30 cycles) performed using random hexamer primers (Perkin-Elmer Cetus, Norwalk, CT).
with the step-cycle program set to denature at 94˚C for 1 min, anneal at 50˚C for 1 min, and extend at 72˚C for 2 min. PCR products were electrophoresed on 2% agarose gels stained with ethidium bromide. We observed a linear increase in the amplification of PCR products with increased amounts of RNA up to 1 mg, as well as with increasing PCR cycle number until 30 cycles, suggesting that our results truly reflect differences in mRNA expression of PCNA and cdk2. We used β-actin as an internal control to standardize the amount of total RNA utilized for RT-PCR. We performed other sets of RT-PCR without RNA samples as negative controls to be certain that there was no artificial amplification.

Gel mobility shift assay. The nuclear extract was prepared from cultured synovial fibroblasts using methods described previously (11,14). In brief, synovial fibroblast pellets were homogenized with a Potte-Elvehjem homogenizer in 4 volumes of ice-cold homogenization buffer [10 mM HEPES pH 7.5, 0.5 M sucrose, 0.5 mM spermidine, 0.15 mM spermin, 5 mM EDTA, 0.25 M ethylene glycol tetra acetic acid (EGTA), 7 mM β-mercaptoethanol, 1 mM phenylmethylsulphonyl fluoride]. After centrifugation at 12,000 g for 30 min at 4˚C, the pellets were lysed and homogenized in a Dounce homogenizer in 1 volume of ice-cold homogenization buffer containing 0.1% NP-40. They were then centrifuged at 12,000 g for 30 min at 4˚C and the pellet nuclei were washed twice with ice-cold buffer containing 0.35 M sucrose. The nuclei were pre-extracted with 1 volume of ice-cold homogenization buffer containing 0.05 M NaCl and 10% glycerol for 15 min at 4˚C. The nuclei were then extracted with homogenization buffer containing 0.3 M NaCl and 10% glycerol for 1 h at 4˚C, following which the concentration of DNA was adjusted to 1 μg/ml. After the nuclear extract was pelleted at 12,000 g for 30 min at 4˚C, the supernatant was brought to 45% (NH₄)₂SO₄ and stirred for 30 min at 4˚C. The precipitated DNA was purified over a nick column (Pharmacia LKB Biotechnology). Binding reactions (10 μl) including the △P-labeled probe (0.5-1 ng, 10,000-15,000 c.p.m.), and 1 μg of polydeoxyinosinic-deoxyctydic acid (Sigma Chemical Co.) were incubated with nuclear extract for 30 min at room temperature and then loaded on a 5% polyacrylamide gel. The gels were subjected to electrophoresis, dried, and pre-extracted with parallel samples 10 min before the addition of the labeled probe.

Organ culture. The synovial tissue specimen was cut into small pieces, washed 3 times in phosphate-buffered saline (PBS), and its wet weight determined. Synovial tissues were cultured on 24-well plates at 150 mg/well in DMEM (Gibco BRL, Grand Island, NY) supplemented with 10% FCS (Hyclone Laboratories, Logan, UT). Synovial tissues were placed for 6 h prior to the transfection in a serum-free medium, then washed 3 times in BSS containing 2 mM CaCl₂. The HVJ-liposome complex (1000 μl) containing 1.3 mg of lipid and 50 μg of encapsulated E2F decoy ODN DNA, and HMG-1 was added to the synovial tissues. The tissues were incubated at 4˚C for 5 min and then at 37˚C for 30 min. After incubation the medium was changed to fresh medium containing 10% FCS.

Preparation of SCID-HuRAg mice. The previously reported SCID mouse model for human RA (20-23) was evaluated as the model for the treatment study. Six-week-old male SCID mice (CB.17/1cr; Japan Clear, Tokyo, Japan) were used for establishment of SCID-HuRAg model. Normal human articular cartilage with subchondral bone was collected from trauma patients with a femoral neck fracture after informed consent at the time of surgery. The complexes of articular cartilage 4.5-mm diameter and RA synovial tissue (100 mg) transfected with E2F decoy ODN or scramble decoy ODN were co-implanted under the skin of SCID mice. The mice were anesthetized with diethyl ether, according to the guidelines established by the Animal Ethics Committee of Osaka University Medical School. A 1-cm incision was made in the middle of the back, and paravertebral muscle was exteriorized. The back muscle was incised, and RA synovial tissue and normal human cartilage were co-implanted. The entire procedure was performed under sterile conditions.

Specimen evaluation. Forty-five days after implantation, implants were removed, immediately cut off the subchondral bone and then snap-frozen in OCT Tissue Tek. To evaluate the effect of E2F decoy ODN on cartilage invasion by RA synovial tissue, the sections were stained with hematoxylin and eosin, and invasion of the articular cartilage and degradation of perichondrocytic cartilage were evaluated by the following previously reported criteria: Invasion score: 0 = no or minimal invasion, 1 = visible invasion (~ >2 cell depths), 2 = invasion (~ >5 cell depths), 3 = deep invasion (~ >10 cell depths). Cartilage degradation scores: 0 = no degradation, 1 = visible degradation, 2 = degradation, 3 = intensive degradation (24).

Statistical analysis. Results are expressed as means ± standard error of the mean (SEM). Mann-Whitney U test was used to determine significant differences. p<0.05 was considered significant. All experiments were carried out at least 3 times.

Results

Transfection of FITC-labeled ODN into cells in the synovial tissue. We first verified that double-stranded ODN tagged with FITC at either the 3’ or the 5’ end could be introduced efficiently into synovial cell nuclei using the HVJ-liposome method. Synovial tissues were fixed 1 and 7 days after transfection and observed by fluorescence microscopy. One day after transfection without HVJ-liposome, little fluorescence was detected in synovial tissue (Fig. 1C). One day after
transfection with the HVJ-liposome method, fluorescence was detected in both the nuclei and the cytoplasm. We detected FITC-labeled ODN in the nuclei of ~50% of the cells (Fig. 1E and F). Even 7 days after transfection with HVJ-liposomes, FITC-labeled ODN were detected in both nuclei and cytoplasm (Fig. 1G and H). No fluorescent signal was seen in the non-transfected cells (Fig. 1A).

**E2F activation in synovial fibroblasts derived from RA.** We examined whether or not E2F was suitable target for inhibition of proliferation of synovial fibroblasts derived from RA (Fig. 2). First, we confirmed the upregulation in E2F-binding activity in synovial fibroblasts derived from patients with RA (lane 2). When RA synovial fibroblasts were stimulated with TNF-α, an increase in E2F-binding activity was observed (lane 3). The gel mobility shift assay demonstrated that
E2F-binding activity was enhanced in synovial fibroblasts from patients with RA, but not in synovial fibroblasts from trauma patients (lane 4). This E2F-binding was eliminated by pre-incubation of nuclear extracts with excess amounts of unlabeled E2F ODN. Pre-incubation with excess amounts of unlabeled double-stranded scrambled ODN did not interfere with detection of E2F-binding in TNF-α-stimulated synovial fibroblasts derived from RA (lanes 5-7).

**Effect of E2F decoy ODN on the inhibition of synovial cell proliferation.** One of the characteristic features of RA is abnormal synovial proliferation leading to joint destruction. We investigated the ability of E2F decoy ODN to inhibit synovial cell proliferation. The level of synovial cell proliferation was determined by using the WST-1 cell counting kit 4 days after transfection. The index of cell proliferation determined by absorbance at 450 nm for synovial cells transfected with E2F decoy ODN or scrambled decoy ODN, and subjected to RT-PCR analysis for cdk2, PCNA and GAPDH. Lane 1, mRNA from synovial tissues from a trauma patient; lane 2, mRNA from synovial tissues with RA; lane 3, mRNA from synovial tissues with RA after transfection with scrambled decoy ODN; lane 4, mRNA from synovial tissues with RA after transfection with E2F decoy ODN using HVJ-liposome; lane 5, mRNA from synovial tissue with RA after transfection with scrambled decoy ODN. Total RNA (30 μg) was used for each blot.

**Effect of E2F decoy ODN on the gene expression of synovial tissues.** The transcription factor E2F plays an important role in the transactivation of the cell cycle regulatory genes. We then examined whether or not E2F inhibition would result in decrease of the downstream cell cycle genes such as PCNA and cdk2. Total RNA was extracted from synovial tissues 24 h after transfection with E2F decoy ODN or scrambled decoy ODN, and subjected to RT-PCR analysis for cdk2, PCNA and GAPDH. The expression levels of PCNA and cdk2 gene in untransfected synovial cells were upregulated. Transfection with E2F decoy ODN resulted in a marked attenuation of PCNA and cdk2 gene expression. The 18S rRNA expression level was not affected by E2F decoy ODN (Fig. 4).

**Effect of E2F decoy ODN on production of inflammatory cytokines.** Since proinflammatory mediators are thought to play a critical role in the pathogenesis of RA, we monitored the production of proinflammatory mediators IL-1β, IL-6 and MMP-1. The protein levels of IL-1β, IL-6 and MMP-1 secreted into the culture medium by RA synovial tissue were
studied 72 h after E2F decoy ODN transfection (n=5 in each group) (Fig. 5). The average level of IL-1ß was 100.2±7.2 pg/ml in untreated group and 104.7±22.5 pg/ml in scrambled decoy group, and 37.4±15.2 pg/ml in E2F decoy group, respectively. The average level of IL-6 was 995±145 pg/ml in untreated group and 1,032±120 pg/ml in scrambled decoy group, and 441±131 pg/ml in E2F decoy group, respectively. The average level of MMP-1 was 775±136 pg/ml in untreated group, 850±372 pg/ml in scrambled decoy group, and 283±76 pg/ml in E2F decoy group, respectively. E2F decoy ODN transfection reduced IL-1ß, IL-6 and MMP-1 production by 54.8±7.4, 42.4±9.5, 28.0±9.9%, respectively. Scrambled decoy ODN had no effect on the production of these mediators.

Marked suppression of invasion of cartilage by RA synovial tissue transfected with E2F decoy ODN. The therapeutic effect of E2F decoy ODN on joint destruction was examined in the severe combined immunodeficient (SCID) mice model for human RA. The volume of RA synovial tissue transfected with E2F decoy ODN was decreased as compared with untreated synovial tissue or synovial tissue transfected with scramble decoy ODN. The articular cartilage co-implanted with RA synovial tissue transfected with scramble decoy ODN showed multiple areas of synovial tissue invasion extending deep into the cartilage. The histological structure of RA synovitis was preserved in these groups. In contrast, the articular cartilage co-implanted with RA synovial tissue transfected with E2F decoy ODN showed marked suppression of invasion by synovial tissue. Infiltrating cells in synovium were decreased in this group (Fig. 6A-F). The grades of cartilage invasion co-implanted with E2F decoy ODN transfected synovial tissue ranged from 1 to 2, with an average of 1.4±0.8 and 1.8±0.4, respectively. The grades of cartilage invasion co-implanted with E2F decoy ODN transfected synovial tissue ranged from 0 to 1, with an average of 0.4±0.5. The grade of cartilage invasion was significantly lower compared with untreated or scrambled decoy ODN group (Fig. 6G).

Discussion

One of the characteristic features of RA is the extensive bone and cartilage erosion caused by the invasive proliferative synovium derived from activated fibroblasts. Thus, cell cycle modulation is an attractive therapeutic target for bone and cartilage erosion in the affected joint of RA. Recent studies demonstrated that suppression of cell cycle in synovial fibroblasts results in inhibition of experimental arthritis (25). Additionally, recent work has found that E2F functions primarily as cell proliferation, suggesting that suppression of transcription of E2F-RB complex suppresses entry into the S phase (26,27). In this study, we used transcription factor E2F decoy oligodeoxynucleotides (ODN) to inhibit synovial cell proliferation. Our data demonstrate that E2F levels in nuclear synovial fibroblast extracts were markedly increased in RA. The specificity of the radiolabeled E2F probe was confirmed by adding a 100-fold excess of unlabeled double stranded E2F decoy ODN to the reaction compared with similar quantities of missense ODN that do not bind E2F. An inhibitory effect of E2F decoy ODN on RA synovial cell proliferation in vitro was shown in this study, as well as in the SCID mice model for human RA. Histological examination showed marked reduction of both the volume and cellularity in synovial tissue transfected with E2F decoy ODN. We showed that nuclear E2F expression in RA synovial tissue was downregulated by transfection of E2F decoy ODN following decreased mRNA expression of the
cell cycle regulatory genes PCNA and cdk2. These results demonstrated that one mechanism by which E2F decoy ODN reduces synovial cell-associated damage is through inducing cell cycle arrest of synovial cells.

The production of proinflammatory mediators such as IL-1ß, IL-6, and MMP-1 was also suppressed by the transfection of E2F decoy ODN. This might be a secondary effect of inhibition of synovial cell proliferation, as synovial cells are the major producers of proinflammatory mediators in RA. However, the biological significance of E2F decoy ODN on proinflammatory mediator production in vivo still remains unclear.

One of the most important strategies for treating RA is prevention of joint destruction. We investigated the inhibitory effect of E2F decoy ODN on cartilage invasion with an in vivo murine model for human RA. Human normal articular cartilage and synovial tissue from patients with RA were co-implanted into SCID mice. On the basis of our in vitro observations, the predicted effect of E2F inhibition in this model would be prevention of cartilage invasion. The results of this study showed that transfection with E2F decoy ODN significantly suppressed cartilage invasion in this model. The exact etiology and pathogenesis have not yet been fully elucidated; however, stimulation of synovial cells by proinflammatory mediators results not only in proliferation but also in a wide variety of biological responses, including alterations in the generation of many effector molecules potentially involved in the pathological process of cartilage destruction. With respect to this point, E2F decoy ODN strategy would be beneficial not only for inducing cell cycle arrest but also for inhibition of proinflammatory mediator secretion.

New classes of technologies, such as antisense ODN, ribozymes (28-31), and RNA interference (32,33), have been adopted by the arthritis field as strategies to inhibit target gene expression in a sequence-specific manner. In the present study, we used a unique molecular strategy: a synthetic double-stranded DNA with high affinity for a target transcription factor is introduced into target cells as a decoy cis element to bind the transcription factors and alter gene transcription. We previously reported a therapeutic strategy to suppress the inflammatory process in synovial cells by transfecting decoy ODN into the affected joints in experimental arthritis models (12). The decoy ODN strategy is more effective than antisense ODN because it blocks multiple transcriptional factors that bind to the same cis element. There are several members of the E2F family, and...
this strategy of using an E2F decoy ODN inhibits all E2F members because the decoy competitively blocks binding to the cis element. This ability of decoy ODN to block all transcription factors binding to a particular cis element is the reason that we focused on this technique and we suggest that this method may be more effective than the manipulation of a single cell cycle regulatory gene.

From the therapeutic point of view, the efficiency, stability, and specificity of decoy ODN in the tissue and cellular delivery are the most important issues to address. Specificity of the E2F decoy ODN was clearly shown in this study by gel mobility shift assay. The main limitation of unmodified ODN is the rapid degradation by nucleases prevalent in sera, cell, and tissue. To rectify this problem, ODNs have been chemically modified with sulfur ions, methyl groups, or other modifications to enhance their resistance against nuclease activity. Although the stability of ODNs is enhanced by these chemical modifications, other problems have been encountered that are attributed to foreign materials (28-30). In this study, we employed a circular dumbbell structure (CD) decoy ODN instead of phosphorothioate-modified decoy ODN to overcome the disadvantages of the chemically modified form; so to enhance their uptake and stability in synovial tissue or synovial fibroblast cells, we used the phosphorothioate-modified decoy ODN and HVJ-liposome delivery system. The effectiveness and usefulness of this delivery system has been demonstrated in various disease model both in vivo and in vitro (35,36). Additional effort is now being applied to use of the circular dumbbell structure decoy ODN instead of phosphorothioate modified decoy ODN to overcome the disadvantages of the chemically modified form (37).

In conclusion, the specificity of the effects of the E2F decoy ODN is supported by the results that only transfection with E2F decoy ODN and not with scrambled decoy ODN could achieve the following effects: a) decreased E2F expression in synovial fibroblasts nuclear extracts, b) decreased mRNA expression of the cell cycle regulatory genes such as cdk2 and PCNA, c) inhibition of cell proliferation, d) decreased production of inflammatory mediators, e) amelioration of cartilage invasion. Since the induction of cell cycle progression appears to be critical for the activation of synovial cells, therapeutic strategies designed to arrest cell cycle progression may prove to be effective in treating cartilage destruction by proliferative synovitis.

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References


