



2 Extracorporeal shock waves promote healing of
3 collagenase-induced Achilles tendinitis and
4 increase TGF- β 1 and IGF-I expression

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12 **Abstract**

13 Extracorporeal shock waves (ESW) have recently been used in resolving tendinitis. However, mechanisms by which ESW
14 promote tendon repair is not fully understood. In this study, we reported that an optimal ESW treatment promoted healing of
15 Achilles tendinitis by inducing TGF- β 1 and IGF-I. Rats with the collagenase-induced Achilles tendinitis were given a single ESW
16 treatment (0.16 mJ/mm² energy flux density) with 0, 200, 500 and 1000 impulses. Achilles tendons were subjected to biomechanical
17 (load to failure and stiffness), biochemical properties (DNA, glycosaminoglycan and hydroxyproline content) and histological
18 assessment. ESW with 200 impulses restored biomechanical and biochemical characteristics of healing tendons 12 weeks after
19 treatment. However, ESW treatments with 500 and 1000 impulses elicited inhibitory effects on tendinitis repair. Histological
20 observation demonstrated that ESW treatment resolved edema, swelling, and inflammatory cell infiltration in injured tendons.
21 Lesion site underwent intensive tenocyte proliferation, neovascularization and progressive tendon tissue regeneration. Tenocytes at
22 the hypertrophied cellular tissue and newly developed tendon tissue expressed strong proliferating cell nuclear antigen (PCNA) after
23 ESW treatment, suggesting that physical ESW could increase the mitogenic responses of tendons. Moreover, the proliferation of
24 tenocytes adjunct to hypertrophied cell aggregate and newly formed tendon tissue coincided with intensive TGF- β 1 and IGF-I
25 expression. Increasing TGF- β 1 expression was noted in the early stage of tendon repair, and elevated IGF-I expression was persisted
26 throughout the healing period. Together, low-energy shock wave effectively promoted tendon healing. TGF- β 1 and IGF-I played
27 important roles in mediating ESW-stimulated cell proliferation and tissue regeneration of tendon.

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29 **Keywords:** Achilles tendon; Tendinitis; TGF- β 1; IGF-I; Shock waves

30 **Introduction**

31 Insertional tendinitis at the tendon-bone junction is a
32 chronic musculoskeletal disorder that can cause severe
33 pain and impaired function. Repair of a tendon is a
34 complex process requiring inflammatory response, neo-
35 angiogenesis, fibrillogenesis, and matrix remodeling [12].
36 Several methods including growth factor interventions
37 have been used to promote tendon repair [9,11,17].

38 Extracorporeal shock waves (ESW) are generated by
39 high voltage spark discharge under water, which causes
40 an explosive evaporation of water, producing high-en-
41 ergy acoustic waves. Focusing the acoustic waves with a
42 semi-ellipsoid reflector, we can focus primary shock
43 waves to a specific tissue site [22]. ESW has been found
44 to be an effective non-invasive treatment for resolving
45 calcifying tendinitis of shoulder, painful heel syndrome,
46 lateral epicondylitis of the elbow and bony healing of
47 nonunion [18,31-33]. ESW treatment has been found by
48 scintigraphy and sonography to affect local blood flow
49 and metabolism of rabbit bone and Achilles tendon
50 [20,26]. However, the cellular and biochemical mecha-
51 nisms by which ESW enhance tendon repair remains to

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52 be determined. We have recently demonstrated that
53 ESW-promoted osteogenic differentiation of mesenchy-
54 mal stem cells and healing of segmental femoral defect in
55 rats was achieved by the induction of bone morphoge-
56 netic proteins and transforming growth factor (TGF-1)
57 [35-37]. These findings imply that physical ESW may
58 bring about tissue regeneration by triggering anabolic
59 activities in cells.

60 Tenocyte growth and neovascularization have been
61 reported to be critical features in the early stage of
62 tendon healing [15]. Tenocytes have been found to
63 convert biophysical stimulation into a biochemical re-
64 sponse leading to release of growth factors and cellular
65 adaptation [29]. Of the growth factors regulating tendon
66 repair, TGF- β and insulin-like growth factor-I (IGF-I)
67 have been found to promote tendon regeneration by
68 regulating collagen metabolism and tenocyte prolifera-
69 tion [1,2,5,25]. One of our previous studies has demon-
70 strated that ESW treatment increased the number of
71 neovessels at the normal tendon-bone junction in dogs
72 [34]. One of our more recent of our study has shown that
73 ESW promotes neovascularization through the release
74 of anabolic and bio-active materials [38].

75 This study aims to elucidate the effect of various
76 impulses of ESW treatment on the healing of colla-
77 genase-induced Achilles tendinitis, and to investigate bio-
78 chemical and biomechanical properties of healing
79 tendons, and to investigate whether physical ESW pro-
80 motion of tendon repair is linked to increases in tenocyte
81 proliferation and induction of growth factors.

82 Materials and methods

83 Collagenase-induced tendinitis model

84 All procedures and protocols were approved by the Institutional
85 Animal Care and Use Committee of Chang Gung Memorial Hospital,
86 Taiwan. Three-month-old male Sprague-Dawley rats (National
87 Experimental Animals Production Center, Taipei, Taiwan) were caged
88 in pairs and maintained on rodent chow and water ad libitum. Rats
89 were anesthetized by an intraperitoneal injection of pentobarbital so-
90 dium (50 mg/kg; Nembutal[®] sodium, Abbott Laboratories, IL, USA).
91 Hindlimbs of each rat were shaved and washed in Betadine. A total
92 250 unit (30 μ l) of collagenase or 30 μ l saline was injected percutane-
93 ously near the osteotendinous junction of the left Achilles tendon using
94 a 30 G needle, respectively. Collagenase (bacterial collagenase type I;
95 Sigma Chemical Inc. St. Louis, MO, USA) was dissolved in normal
96 saline and sterile filtered through a 0.22 μ m filter. Chloromycetin (10
97 mg/kg, intraperitoneal injection) and analgesia (0.02 mg/kg bupenor-
98 phine, subcutaneous injection; Reckitt and Colman Pharmaceutical
99 Inc., Richmond VA, USA) were administered for 2 days. The injected
100 area and ankle did not display infection, edema and redness after
101 injecting 30 μ l sterile normal saline. The collagenase injection did not
102 appear to have adverse effects on the animals, because all the rats
103 treated with collagenase maintained their weights or gained weight
104 during the study period.

105 ESW treatment

106 Each rat with an Achilles tendinitis was re-anesthetized and placed
107 in prone position with the left heel up. The shock wave tube was fo-
108 cused on the Achilles tendon near the insertion site to the heel bone

using a C-arm image intensifier and the treatment device's control
guide. The ESW treatment (0.16 mJ/mm² energy flux density, 1 Hz;
HMT High Medical Technologies GmbH, Kreuzlingen, Switzerland)
was used 3 days after collagenase injection. Ultrasound transmission
gel (Pharmaceutical Innovations Inc, Trenton, NJ, USA) was used as
contact medium between the ESW apparatus and skin.

Experimental design

Experiment I

The total acoustic energy of ESW treatment applied to tissue is
presented as the number of pulses multiplied by the energy per pulse
[22]. In a pilot study, 12 weeks after ESW treatment (0.16 mJ/mm²
energy flux density, 1 Hz, 200 impulses), rats with the Achilles tendi-
nitis were found to restore mechanical properties of tendons. In the
present study, we examined the effects of ESW energy on biochemical
and biomechanical characteristics of tendons. Seventy-five rats were
employed for studies. Sixty rats with an Achilles tendinitis were
randomly divided into four groups. Rats in each group received ESW
treatment (0.16 mJ/mm² energy flux density, 1 Hz) for control
($n = 15$), 200 impulses ($n = 15$), 500 impulses ($n = 15$) and 1000 im-
pulses ($n = 15$), respectively. The remaining rats ($n = 15$) received 30 μ l
sterile saline and received no ESW treatment were used as vehicle
group. Rats in each group were killed with an overdose of pentobarbi-
tal sodium 12 weeks after ESW treatment. Twelve Achilles tendons
were harvested for biochemical and biomechanical assessment and
three Achilles tendons were subjected to histomorphometry.

Experiment II

Forty-eight rats with the tendinitis were randomly divided into
ESW and control groups. Twenty-four rats were given ESW at 0.16
mJ/mm² for 200 impulses. These rats were killed in 3 days ($n = 4$) after
collagenase injection, 1 week ($n = 4$), 2 weeks ($n = 4$), 4 weeks ($n = 4$),
6 weeks ($n = 4$) and 12 weeks ($n = 4$) after ESW treatment. The
remaining 24 rats received no ESW treatment. Four rats were killed at
the same time point as mentioned for the ESW group and were dis-
sected for histological assessment.

Biomechanical properties determination

Tendons were excised through two incisions, one distal to the
musculotendinous junction and the other one proximal to the calcaneal
bone. The biomechanical properties of the specimens were assessed
using a Material Testing System QT-10 (MTS Corp. Minneapolis,
MN, USA) with custom designed cryoclamps. Tendons were loaded at
a rate of 20 mm/min. During this test, specimens were kept moistened
with normal saline. For each sample, load-displacement curves were
plotted, and load-to-failure and stiffness were calculated for each
sample.

Glycosaminoglycan, DNA and hydroxyproline content measurement

After biomechanical assessment, each specimen containing both
rupture tendons was ground with a mortar and pestle under liquid
nitrogen. Each ground specimen was divided into two portions.
Specimens were digested in 0.5% papain at 65 $^{\circ}$ C for 4 h. Glycos-
aminoglycan contents in tendons were spectrophotometrically deter-
mined using dimethylmethylene blue as previously described [13].
Papain digests were incubated at 65 $^{\circ}$ C for a further 20 h. DNA
contents in mixtures were determined using bisbenzimidazole-fluorometric
assay [24]. Specimens were acid hydrolyzed with 6 N HCl at 110 $^{\circ}$ C for
24 h and derivatized with phenylisothiocyanate. Hydroxyproline con-
tent was determined using a high performance liquid chromatography
(LC-10AD, Shimadze, Tokyo, Japan) equipped with a reverse phase
column (4.6 mm internal diameter \times 250 mm length; TSK-gel, ODS-
80TM). The column was eluted with 0.4% ammonium acetate (pH 7.4)
and 75% acetonitrile at a flow rate of 2 ml/min through an isocratic
pump. The hydroxyproline concentration in each sample was detected
using a fluorescence detector (RF-10AXL, Shimadze, Tokyo, Japan)
and integrated from the retention time and area under the eluting peak.
Results were normalized with protein concentration in each sample.

173 Immunohistochemistry

174 Specimens were fixed in 4% buffered paraformaldehyde for 48 h
175 and then decalcified in PBS-buffered 10% EDTA. Decalcified tissues
176 were embedded in paraffin. Tendon specimens were cut longitudinally
177 into 5- μ m sections and transferred to poly-lysine-coated slides for
178 conventional hematoxylin-eosin staining (Sigma Chemicals Inc, St.
179 Louis, MO, USA). Immunoreactivity in specimens was demonstrated
180 using a horseradish peroxidase (HRP)-3', 3'-diaminobenzidine (DAB)
181 cell and tissue staining kit (R&D Systems Inc, Minneapolis, MN,
182 USA) according to manufacturer's instructions. Sections were immu-
183 nostained for proliferating cell nuclear antigen (PCNA), TGF- β 1 and
184 IGF-1 (Upstate Biotechnology, Lake Placid, NY, USA). Sections were
185 further incubated with biotinylated secondary antibodies with strep-
186 tavidin conjugated to HRP, followed by chromogen solution and
187 counterstaining with hematoxylin. Sections were dehydrated and
188 mounted. Sections without primary antibodies were enrolled as nega-
189 tive controls for the immunostaining.

190 Histomorphologic assessment

191 Five areas of tendon were randomly chosen for observation from
192 three sections of four rats. To quantify immunostaining, the sections
193 were analyzed using a Zeiss Axioskop 2 plus microscope (Carl Zeiss,
194 Gottingen, Germany). An area (3 mm²) with positive cells was selected
195 for analysis. Three random images of 0.75 mm² from each selected area
196 (3 mm²) were then taken under 400 \times magnifications. All the images of
197 each specimen were captured using a Cool CCD camera (SNAP-Pro
198 c.f. Digital kit; Media Cybernetics, Sliver Spring, MD, USA). Images
199 were analyzed using an Image-Pro⁺ Plus image-analysis software
200 (Media Cybernetics, Sliver Spring, MD, USA). The number of the
201 hematoxylin-stained nuclei in each image was counted using an Image-
202 Pro⁺ Plus image-analysis software (Media Cybernetics, Sliver Spring,
203 MD, USA). A professional pathologist, blinded to the treatment reg-
204 imen performed measurement on all sections.

205 Statistical analysis

206 All values were expressed as mean + standard error. Wilcoxon test
207 was used to evaluate the difference between the sample of interest and
208 its respective control. For analysis time course, a multiple range of
209 ANOVA was used. A *P*-value of <0.05 was considered significant.

210 Results

211 Animal activity

212 The injected area and ankle displayed edema and
213 redness 1 day after injection. Rats walked with lameness
214 within 3 days after collagenase injection. ESW with 200
215 and 500 impulses did not induce skin hemorrhage or
216 ecchymosis, but ESW with 1000 impulses immediately
217 caused skin ecchymosis after treatment. Improvement of
218 lameness and edema of ankle varied with ESW dose.
219 ESW with 200 impulses completely improved lameness
220 and edema of ankle (*n* = 15) 4 weeks after treatment.
221 ESW with 500 impulses was found to resolve lameness
222 to some extent (*n* = 5) 8 weeks after treatment. No
223 improvement in lameness was observed in the control
224 and 1000-impulse groups of animals during the study.

225 Biomechanical and biochemical properties

226 All specimens displayed a typical load-displacement
227 curve, with an upward linear slop and a failure response

at the point of failure (Fig. 1). Compared with the
vehicle group, ESW treatment with 200 impulses sig-
nificantly restored mechanical load-to-failure and stiff-
ness of healing tendons. ESW treatment with 500
impulses and 1000 impulses significantly decreased bio-
mechanical properties of tendons (Table 1). We further
determined the influences of ESW on biochemical
properties of tendons. ESW treatment with 200 impulses
reversed DNA, glycosaminoglycan and hydroxyproline
contents of healing tendons, bringing them back to
levels comparable to those of normal tendons. ESW
treatment with 500 impulses and 1000 impulses elicited
inhibitory effects on biochemical characteristics of ten-
dons (Table 1). Histological observation also showed
that injured tendons subjected to ESW treatment with
200 impulses displayed well-aligned tendon fiber bundle
arrangement 12 weeks after ESW (Fig. 2A). In the 500-
impulse group (Fig. 2B) and 1000-impulse group (Fig.
2C), injured tendons were filled with loose fibrous tis-
sues. In control group, the lesion site was degenerative
and surrounded by fibrous and connective tissue (Fig.
2D). Results indicated that ESW treatment with 200
impulses promoted tendinitis healing, whereas ESW
treatment with greater than 200 impulses may suppress
tendon repair. ESW treatment with 200 impulses was
then used for the proceeding experiments.

254 Kinetic histological changes in ESW-promoted healing of
255 Achilles tendinitis

256 Three days after collagenase injection, areas of sur-
257 rounding the tendon displayed obvious edema, erythro-
258 cyte extravasation and massive inflammation cell
259 infiltration. The tendon integrity was disrupted and the
260 lesion site displayed intensive tendon fiber swelling and

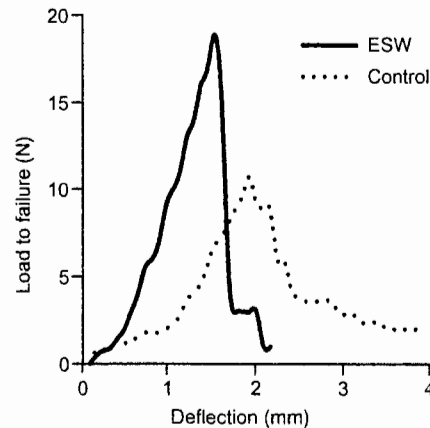


Fig. 1. Representative load failure curves of control and ESW-treated tendons. Achilles tendinitis without or with ESW treatment (0.16 mJ/mm², energy flux density, 1 Hz, 200 impulses were subjected to bio-mechanical determination 12 weeks after ESW).

Table 1
Biochemical and biomechanical properties of collagenase-induced Achilles tendinitis with and without ESW treatment

	Vehicle ^a	Control ^b		ESW at 0.16 mJ/mm ² energy flux density					
		Control	<i>P</i> -value ^c	200 impulses	<i>P</i> -value ^c	500 impulses	<i>P</i> -value ^c	1000 impulses	<i>P</i> -value ^c
Failure load (N)	20.6 ± 3.2	11.8 ± 1.4	0.027	18.6 ± 2.3	0.682	12.5 ± 1.8	0.016	8.1 ± 1.5	0.0032
Stiffness (N/mm)	14.8 ± 1.9	9.3 ± 1.3	0.013	13.4 ± 2.1	0.824	6.3 ± 1.2	0.012	5.6 ± 1.4	0.0074
DNA content (µg/mg)	4.1 ± 0.4	2.7 ± 0.2	0.023	3.9 ± 0.3	0.518	2.0 ± 0.2	0.036	1.7 ± 0.3	0.016
Hydroxyproline (pg/µg)	1.45 ± 0.2	0.95 ± 0.16	0.028	1.33 ± 0.18	0.364	0.85 ± 0.18	0.028	0.78 ± 0.15	0.021
Glycosaminoglycan (µg/mg)	25.7 ± 2.9	17.8 ± 1.7	0.019	23.4 ± 2.7	0.617	15.4 ± 1.4	0.011	13.2 ± 0.9	0.015

Results are presented with mean ± standard errors calculated from 12 rats.

^a Achilles tendons were given 30 µl saline and treated without ESW.

^b Achilles tendons were given collagenase and treated without ESW.

^c Comparisons between vehicle group, control group and ESW groups, respectively.

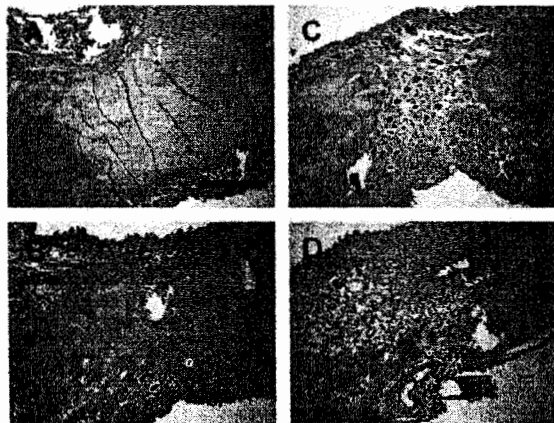


Fig. 2. Histological photographs of collagenase-induced Achilles tendinitis 12 weeks after with or without ESW treatment. (A) Lesion site displayed well-aligned tendon fiber bundle arrangement after 200 impulses of ESW. (B) In the 500-impulse group and (C) 1000-impulse group, injured tendons were filled with loose fibrous tissues. (D) In the control group, the injured tendon was degenerative and surrounded by fibrous and connective tissue. Specimens were stained with conventional hematoxylin-eosin and observed in magnification = 5× and bar scale = 2 mm.

261 necrosis (Fig. 3A). One week after ESW treatment,
262 inflammation was gradually resolved and intensive
263 blood capillaries and extracellular matrix production
264 were observed in lesion site (Fig. 3B). Four weeks after
265 ESW treatment, lesion site appeared to be fused by a
266 fibrous bridge. The granulation tissue and inflammation
267 cell infiltration was completely improved. Well-aligned
268 tendon fiber bundles gradually formed parallel to the
269 long axis of the tendon (Fig. 3C). Six weeks following
270 ESW, fibrous tissue in lesion site was replaced by newly
271 developed tendon tissue. Tendon fiber density was in-
272 creased and matrix remodeling was in process (Fig. 3D).
273 We further assessed histological changes in sections
274 using high power microscopy. Three days after colla-
275 genase injection, lesion site was largely filled with loose

276 fibrous tissue, granulation tissue and connective tissue
277 and intensive inflammatory cells accumulation (Fig. 3E).
278 One week after treatment, a large number of plum-
279 shaped and spindle-shaped fibroblasts from peritendon
280 recruited into the lesion site and turned into hypertro-
281 phied cellular tissue 1 week after ESW treatment (Fig.
282 3F). Four weeks after ESW, spindle-shaped tenocytes
283 gradually oriented into tendon bundles. Intensive ten-
284 don fibril continued to be produced. Fibril increased in
285 size to become histologically visible thin wavy forms
286 (Fig. 3G). Six weeks following ESW, newly formed
287 tendon tissue displayed increasing vascularity within the
288 new tendon and the longitudinal orientation of the
289 collagen bundles (Fig. 3H).

Immunohistochemistry

290

291 We examined the effect of ESW treatment on the
292 tenocyte proliferation. Proliferating cells were immu-
293 nostained for PCNA, a protein associated with the S-
294 phase of a dividing cell. The nuclei of cells with positive
295 PCNA expression appeared as brown color. Proliferat-
296 ing cells were concentrated at the lesion site. Within 1
297 week and 6 weeks after ESW, cuboidal-, plum-shaped
298 (Fig. 4A) and spindle-shaped tenocytes (Fig. 4B) adja-
299 cent to hypertrophied cellular tissue and newly formed
300 tendon tissue matrix were found to have undergone
301 intensive PCNA expression. Cells at the fibrous tissue in
302 the control group displayed slight PCNA expression
303 throughout the study period (Fig. 4C and D). We
304 compared the number of proliferating cells in injured
305 tendon of both groups and found ESW treatment able
306 to stimulate a significant increase in cell proliferation
307 within 6 weeks of treatment (Table 2).

308 We further investigated immunohistochemically
309 whether ESW promotion of tenocyte proliferation and
310 tendon healing was linked to TGF-β1 and IGF-I
311 induction. One week after ESW, plum-shaped tenocytes
312 at hypertrophied cellular tissue were found to express
313 intensive TGF-β1 (Fig. 5A). Four weeks after treatment,

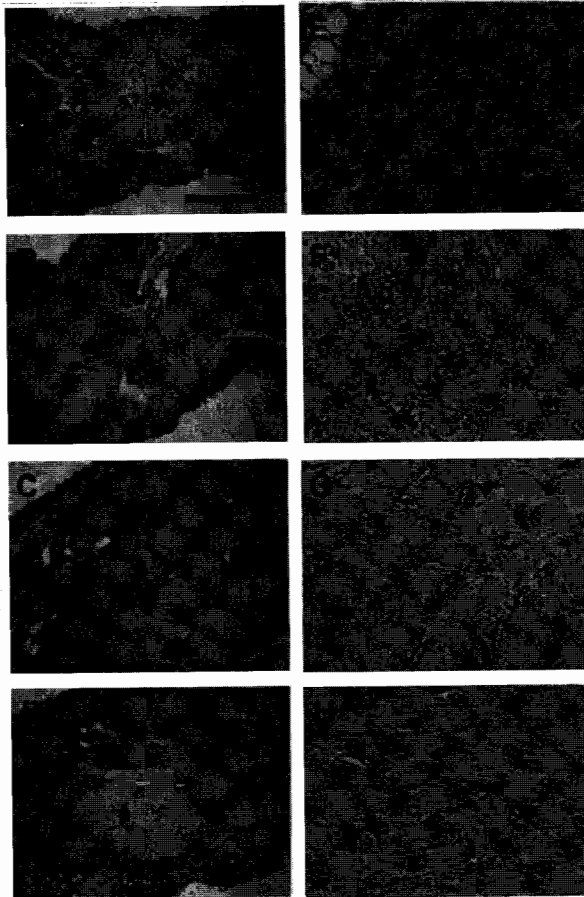


Fig. 3. Histological changes in ESW promotion of Achilles tendinitis repair. (A) Tendon displayed evident swelling, edema, and partially ruptured 3 days after percutaneous injection of collagenase. (B) Inflammation was clearly resolved and extracellular matrix was produced 1 week after ESW treatment. (C) The ruptured end filled with well-aligned tendon fibrils within 4 weeks after treatment. (D) The lesion site was replaced by newly formed tendon fiber 6 weeks after ESW treatment. (E) Lesion site displayed intensive inflammatory cell infiltration 3 days after percutaneous injection of collagenase. (F) Plum-shaped tendocytoid aggregated as hypertrophied cellular tissue was adjunct to the injured tendon 1 week after ESW treatment. (G) Intensive tendon fibril production and formed into wave forms 4 weeks after ESW treatment (H) Newly developed tendon tissue appeared to increase vascularity within the repair neotendon 6 weeks after ESW treatment. Rats with the Achilles tendinit were given a single ESW treatment (0.16 mJ/mm² energy flux density, 1 Hz, 200 impulses) 3 days after injection of collagenase. Specimens were stained with conventional hematoxylin-eosin and observed in magnification = 5 \times , bar scale = 2 mm (A, B, C, and D) and in magnification = 20 \times , bar scale = 100 μ m (E, F, G, and H).

314 spindle-shaped tenocytes at the newly developed tendon
315 tissue expressed strong TGF- β 1 (Fig. 5B). Mature
316 spindle-shaped tenocytes in the dense tendon fiber
317 showed weak TGF- β 1 expression in the later stage of
318 healing process (Fig. 5B). In the control sections,
319 cuboidal- and spindle-shape fibroblastic cells located at
320 the lesion site exhibited slight TGF- β 1 expression during
321 experiments (Fig. 5C and D). One week after treatment,

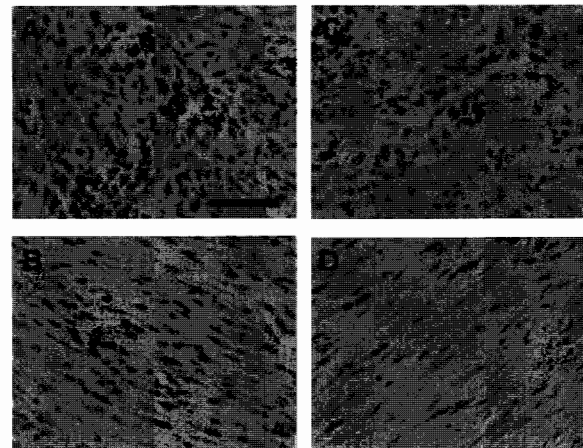


Fig. 4. PCNA expression in the repair of collagenase-induced tendinitis with and without ESW treatment. Within 1 week and 6 weeks after ESW, a large number of plum-shaped and spindle-shaped tenocytes at (A) hypertrophied cellular tissue and (B) newly developed tendon tissue displayed intensive PCNA expression after ESW treatment. Plum- and spindle-shaped tenocytes at (C) lesion site and (D) fibrous tissue of the controls without ESW treatment displayed weak PCNA immunoreactivity. The positive PCNA immunostained cells showed brown color in the nuclei of cells. Specimens were observed in magnification = 40 \times and bar scale = 50 μ m.

tenocytes adjacent to the hypertrophied showed intensive IGF-I expression (Fig. 6A). Four weeks after ESW, cells at newly formed tendon bundles showed intensive IGF-I expression (Fig. 6B). In control sections, cells at the loose fibrous tissue expressed weak IGF-I expression (Fig. 6C and D). Intensities of TGF- β 1 and IGF-I expression varied with the stages of tendon regeneration. Table 2 summarizes the number of tenocytes that exhibited positive TGF- β 1 and IGF-I expression during ESW-promoted tendon repair. There were significant increases in TGF- β 1 expression within 4 weeks of ESW treatment. Elevating IGF-I expression was noted throughout the study period.

Discussion

In this study, we demonstrate that physical ESW treatment promotes repair of collagenase-induced Achilles tendinitis. Injured tendons receiving ESW treatment resolved inflammation, increased tenocyte proliferation and restored tendon integrity, leading to restoration of biomechanical properties. While a number of reports implicating ESW treatment have been used for tendinitis, little has been done to define the role of growth factor expression in the repair processes. The findings in this study provide the first evidence that ESW promotion of Achilles tendinitis repair coincides with increases in TGF- β 1 and IGF-I. We propose that it is the increased mitogenic and anabolic responses of ten-

Table 2

Expression of PCNA, TGF-β1 and IGF-I in collagenase-induced Achilles tendinitis with and without ESW treatment

		Week(s) after ESW treatment				
		1	2	4	6	12
PCNA	ESW ^a	932 ± 183	1024 ± 198	814 ± 206	498 ± 102	376 ± 92
	Control	306 ± 86	324 ± 89	287 ± 64	275 ± 53	283 ± 57
	P-value ^b	<0.001	<0.001	<0.001	0.016	0.37
TGF-β1	ESW	614 ± 102	532 ± 93	417 ± 79	318 ± 53	312 ± 48
	Control	226 ± 43	257 ± 51	235 ± 46	231 ± 51	242 ± 47
	P-value	0.0074	0.011	0.018	0.023	0.46
IGF-I	ESW	562 ± 67	654 ± 98	736 ± 105	514 ± 82	467 ± 58
	Control	265 ± 58	249 ± 63	247 ± 72	238 ± 54	256 ± 52
	P-value	0.013	0.0034	0.0078	0.021	0.016

^aResults are presented with mean ± standard errors calculated from five tendon of areas from three sections of four rats after ESW treatment at 0.16 mJ/mm² energy flux density, 1 Hz, 200 impulses.

^bComparison between ESW and control groups.

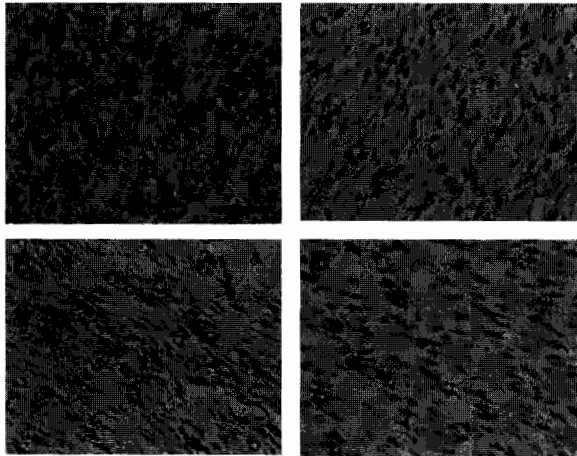


Fig. 5. Expression of TGF-β1 in collagenase-induced tendinitis with and without ESW treatment. Cells adjunct to (A) hypertrophied cellular tissue 1 week after treatment and (B) newly developed tendinous tissue were found to have intensive TGF-β1 expression after 4 weeks after ESW treatment. In control sections, plum- and spindle-shaped tenocytes at (C) lesion site and (D) fibrous tissue expressed weak TGF-β1. Specimens were observed in magnification = 40× and bar scale = 50 μm.

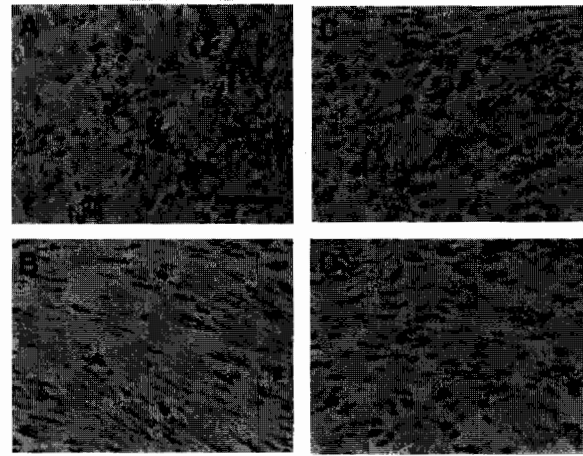


Fig. 6. Expression of IGF-I in collagenase-induced tendinitis with and without ESW treatment. Cells adjunct to (A) hypertrophied cellular tissue 1 week after treatment and (B) newly developed tendinous tissue were found to have intensive IGF-I expression after 4 weeks after ESW treatment. In control sections, plum- and spindle-shaped tenocytes at (C) lesion site and (D) fibrous tissue of the controls displayed weak TGF-β1 immunoreactivity. Specimens were observed in magnification = 40× and bar scale = 50 μm.

349 don tissue that bring about the clinical success of ESW
350 treatment in resolving tendinitis.

351 The etiologies of tendinitis are multifactorial includ-
352 ing avascular changes, degenerative change and meta-
353 bolic disturbance, neural factors and neovascularization
354 [3,4,9]. Previous reports have demonstrated the effect of
355 ESW treatment on pathophysiology of tendon [19,26].
356 However, such studies were performed on normal tend-
357 ons, and their observation should not be extrapolated
358 to explain ESW-promoted tendinitis repair. Collage-
359 nase-induced model is a well-established model for the
360 studies of tendinitis. The acute swelling, inflammation
361 and matrix destruction in tendons are similar to those
362 seen in naturally occurring tendon injuries [1,9,10].

363 Histological observation in the present study revealed
364 that lesion sites in injured tendons displayed massive
365 inflammation cell infiltration and tendon degeneration.
366 This model also clearly showed that the physical ESW
367 treatment stimulated the regeneration of tendon. Our
368 study finds the use of collagenase-induced Achilles ten-
369 dinitis to an excellent model for clarifying the cellular
370 and histological mechanism that biophysical strategies
371 such as ESW treatment uses to stimulate tendon repair
372 in vivo. In contrast to previous studies demonstrating
373 injured tendons healed eventually over time after ten-
374 dotomy [7,23], our current study showed that collage-
375 nase-induced Achilles tendinitis displayed incomplete
376 healing or degenerative tendinitis. We speculate that this

377 discrepancy that tendinitis healing may depend on the
378 model system and experimental animals used. There
379 may be a chronic tendinopathy that does not respond to
380 normal healing time.

381 Our present study indicates that high-energy ESW
382 treatment impairs DNA, extracellular matrix synthesis
383 and biomechanical characteristics of tendons. Mechan-
384 ical or physical impact on tissues exposed to ESW has
385 been found to depend on tissue impedance. ESW-in-
386 duced damage has been reported as a function of the
387 total amount of acoustic energy absorbed in a finite
388 volume [8]. Previous studies have demonstrated that
389 high-energy ESW treatment can induce fibrinoid
390 necrosis, paratenon fibrosis and inflammatory cell infil-
391 tration in normal rabbit Achilles tendons [26] and im-
392 paired tensile strength of tendons [19]. Histological
393 findings of our study showed that loose fibrous con-
394 nective tissue and degenerative tendon formed in the
395 500-impulse and 1000-impulse groups, indicating high-
396 energy ESW treatment is not beneficial for Achilles
397 tendinitis repair in our model.

398 ESW with 200 impulses restored biomechanical and
399 biochemical characteristics of tendons. Histological
400 findings showed that 200 impulses improved edema and
401 inflammatory cell infiltration in injured tendon and
402 progressively promoted tendon regeneration. Improve-
403 ment of lameness also provides an explanation for the
404 trend toward tendon healing promoted by low-energy
405 ESW treatment. Our findings suggest that optimal ESW
406 energy magnitude is beneficial for collagenase-induced
407 Achilles tendinitis healing in rats. In contrast to these
408 findings in rats, human subjects treated for chronic
409 plantar fasciitis and calcifying tendinitis with ESW of
410 more than 200 impulses improved without side effects
411 [27,28]. It has also been our clinical experience that
412 patients treated for painful heel and calcifying tendinitis
413 improved without complications [31,32]. Our findings
414 suggest that ESW stronger than 0.16 mJ/mm², 500 im-
415 pulses should not be used for treating Achilles tendinitis
416 in rats. Soft tissue in rats may be more sensitive to ESW.
417 The impact of high-energy ESW treatments on soft tis-
418 sue in experimental animals may not be readily extrap-
419 olated to humans. Effects of ESW energy magnitude on
420 treating Achilles tendinitis have not been clarified. In
421 one more recent study demonstrated that patients with
422 lateral epicondyliti receiving low-energy ESW (0.07-0.09
423 mJ/mm²) did not improve [16]. More research is needed
424 in ESW energy magnitude required for treating Achilles
425 tendinitis in patients.

426 Increasing fibroblast proliferation and biosynthesis of
427 extracellular matrix and collagen are crucial stage for
428 the return of normal tendon strength [9,10]. In the
429 present study, ESW treatment significantly reversed
430 DNA, glycosaminoglycan and hydroxyproline contents.
431 Histological findings also showed that intensive PCNA
432 expression and new tendon fibril formation were

prominent findings in our study of ESW treatment for
tendon repair. These findings indicate that ESW treat-
ment raises mitogenic and morphogenic responses,
which stimulates tenocyte growth and typical tendon
microstructure formation.

Increased tenocyte growth and tissue regeneration
indicated that some growth factors were responsible for
initiating the ESW-induced mitogenic and morphogenic
responses of injured tendon. Intensive PCNA expression
of tenocytes and new tendon bundle formation coin-
cided with intensive TGF-β1 and IGF-I expression after
ESW treatment. These findings suggest that TGF-β1
and IGF-I, at least in part, are involved in ESW pro-
motion of tendon repair. These growth factors have
been found to up-regulate extracellular matrix biosyn-
thesis of tenocytes [1,2,30]. Tenocytes responded to
mechanical stimulation by increasing TGF-β1 gene
expression [6]. Our findings indicate that tendon tissue
can convert ESW stimulation into biochemical signals
via release of TGF-β1 and IGF-I for tendinitis repair.
Moreover, TGF-β1 has been reported to act as a potent
inhibitor of macrophages-induced extracellular matrix
degradation and inflammation during the healing of a
wound [14,39]. Intensive TGF-β1 expression in the early
stage of ESW-augmented tendon repair may explain the
improvement in inflammation. Our findings suggest that
TGF-β1 and IGF-I are actively involved in tendinitis
healing. Each growth factor has a distinct temporal
expression pattern and each has a potentially unique
role in repair of tendinitis. In summary, our research
findings provide evidence that an optimal ESW treat-
ment promotes tendon regeneration, with TGF-β1 and
IGF-I playing important mitogenic and anabolic roles in
signaling physical ESW stimulation.

Uncited reference 467

[21] 468

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