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***Basic Study***

**Formation process of extension knee joint contracture following external immobilization in rats**

Zhou CX *et al*. Extension knee joint contracture of rats

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**Abstract**

BACKGROUND

Current research lacks a model of knee extension contracture in rats.

AIM

To elucidate the formation process of knee extension contracture.

METHODS

We developed a rat model using an aluminum external fixator. Sixty male Sprague-Dawley (SD) rats with mature bones were divided into the control group (*n* = 6) and groups that had the left knee immobilized with an aluminum external fixator for 1, 2, and 3 d, and 1, 2, 3, 4, 6, and 8 wk (*n* = 6 in each group). The passive extension range of motion, histology, and expression of fibrosis-related proteins were compared between the control group and the immobilization groups.

RESULTS

Myogenic contracture progressed very quickly during the initial 2 wk of immobilization. After 2 wk, the contracture gradually changed from myogenic to arthrogenic. The arthrogenic contracture progressed slowly during the 1st week, rapidly progressed until the 3rd week, and then showed a steady progression until the 4rd week. Histological analyses confirmed that the anterior joint capsule of the extended fixed knee became increasingly thicker over time. Correspondingly, the level of transforming growth factor beta 1 (TGF-β1) and phosphorylated mothers against decapentaplegic homolog 2 (p-Smad2) in the anterior joint capsule also increased with the immobilization time. Over time, the cross-sectional area of muscle fibers gradually decreased, while the amount of intermuscular collagen and TGF-β1, p-Smad2, and p-Smad3 was increased. Unexpectedly, the amount of intermuscular collagen and TGF-β1, p-Smad2, and p-Smad3 was decreased during the late stage of immobilization (6-8 wk). The myogenic contracture was stabilized after 2 wk of immobilization, whereas the arthrogenic contracture was stabilized after 3 wk of immobilization and completely stable in 4 wk.

CONCLUSION

This rat model may be a useful tool to study the etiology of joint contracture and establish therapeutic approaches.

**Key Words:** Knee joint; Immobilization; Contracture; External fixator; Rats

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**Core Tip:** Current research lacks a model of knee extension contracture in rats. The study elucidated the formation process and therapeutic strategies of knee extension contracture. To this end, we developed a rat model using an aluminum external fixator. The results showed that the myogenic contracture was stabilized after 2 wk of immobilization, whereas the arthrogenic contracture was stabilized after 4 wk of immobilization. This rat model may be a useful tool to study the etiology of the joint contracture and establish therapeutic approaches.

**INTRODUCTION**

Knee contracture is currently one of the most common clinical diseases and is characterized by joint capsule fibrosis and a restricted range of motion (ROM) secondary to periarticular intermuscular connective tissue hyperplasia[1]. The signature pathology of joint contracture is the proliferation of myofibroblasts (active fibroblasts) and the deposition of proteins in the extracellular matrix in the joint capsule and intermuscular connective tissue[2]. The most common cause of knee contracture is prolonged immobilization, which is clinically used as an acute treatment for musculoskeletal disease to relieve knee pain and reduce inflammation[3,4]. Knee contracture is unsightly and has adverse effects on function and quality of life, affecting daily activities such as ascending and descending stairs, walking, and toileting. Furthermore, knee contracture is very difficult to treat[5]. Despite a large amount of rehabilitation, conservative treatment, and even surgical treatment, it is difficult to completely restore the joint mobility, and this loss of mobility seriously decreases the quality of life of patients and adversely affects the distribution of medical resources in society[6,7]. It is therefore very important to investigate the mechanisms leading to knee contracture.

The contracture mechanism has been explored in many studies. As early as 1993, a rat flexion contracture model was successfully established by fixing the tibia and fibula in complete flexion (150) for up to 7 wk without damaging the joints[8]. In recent years, various immobilization methods have been introduced to create flexion contracture models by fixing the animal knee joint at about 150 of flexion, including hook buckle (a hook-and-loop fastener) immobilization and helix (spiral wire) immobilization[9,10]. By contrast, extension contracture models are rare. However, the extension contracture model is of clinical relevance because it better mimics fracture and bed-associated immobilization than the flexion contracture model.

According to the general international standard, the neutral position of the knee joint is the straightened or extended position, which is defined as 0. The functional position of the knee joint is from 15 to 20 of flexion, while the ROM of the normal knee is 120 to 150 for flexion and 5 to 10 for overextension. Knee injury usually requires immobilization in a straightened or functional position, but this type of immobilization may result in limited knee flexion motion (knee extension contracture); therefore, knee extension contracture is the most common type of knee contracture. Different fixation methods have different effects on muscles. If the muscle is fixed under the condition of being lengthened, its atrophy and the decrease of muscle contraction force will be less, and the ratio of fast and slow muscles of biceps femoris, which is often selected as the detection index of flexion knee contracture model, is also different from that of quadriceps femoris; however, the current literature describes several animal knee contracture models, most of which involve flexion knee contracture. The structure of the knee joint of rats is similar to that of humans and is easier to obtain compared to other animals. Thus, it is necessary to establish a convenient and reliable rat model of knee extension contracture.

In the present study, we studied the process of knee extension contracture formation during external fixation of the knee in a straightened position in a rat model that we developed. To the best of our knowledge, this is the first report of a rat model of knee extension contracture.

**MATERIALS AND METHODS**

***A rat model of knee extension contracture***

Male Sprague-Dawley (SD) rats (age 8 wk and weight 350 g +) were used in this experiment. Aluminum splints (6061; Longkai, Suzhou, China), sponge (33 d; Changzhou, China), and woodworking (BND-2815; Bonida, Guangdong, China) were prepared. The rats were kept under the same conditions without intervention for 2 wk before the experiment (free diet, day and night balance, temperature 20-25 ℃, humidity 50% ± 5%). Each rat was placed on the operating table in the supine position and fixed. The fixing device and fixing schematic diagram is shown in Figure 1. Figure 2A shows the results of applied fixation. Figure 2B shows the anterior X-ray of the rat knee joint, while Figure 2C shows the lateral X-ray of the rat knee joint. Immobilization was performed under general anesthesia achieved with an intraperitoneal injection of 10% chloral hydrate (0.03 mL/kg). A patent application has been made for the self-made aluminum splint (Patent No. ZL202120470158.0).

Measurements of the knee joints of 8-wk-old male rats (*n* = 15) revealed that the average thigh width was 3.23 cm ± 0.21 cm (range 6.38 ± 0.41 to 7.21 cm ± 0.43 cm) and the average calf width was 4.86 cm ± 0.27 cm (range 2.34 ± 0.13 to 5.11 cm ± 0.36 cm). In accordance with the anatomical characteristics of the rats, the immobilization device was shaped using a wire cutting process with an aluminum plate and bonded with a 0.5-cm-thick sponge on the skin to prevent excessive immobilization. The shape of the aluminum plate is shown in Figure 1. The fixation device placed the knee joint in the straightened position and ensured complete external immobilization of the knee joint. The animal experiments described in this study were authorized by the Experimental Animal Ethics Committee of Anhui Medical University (No. LLSC20221126).

***Grouping and specimen collection***

Sixty rats were randomly divided into 10 groups (*n* = 6 in each group). The control group did not do any intervention. while the immobilization groups had the left hindlimb fixed for 1 d (immobilization-1 d group), 2 d (immobilization-2 d group), 3 d (immobilization-3 d group), 1 wk (immobilization-1 wk group), 2 wk (immobilization-2 wk group), 3 wk (immobilization-3 wk group), 4 wk (immobilization-4 wk group), 6 wk (immobilization-6 wk group), and 8 wk (immobilization-8 wk group). All groups were reared in the same environment, and the ROM of the knee joint was measured on the same day as the 8-wk-fixed group after controlling the fixed time. The rats in the appropriate group were euthanized by an excessive intraperitoneal injection of 10% chloral hydrate. After euthanasia, the fixed left hindlimb of the rat was removed at the hip joint. The skin was separated, and the knee mobility was measured using the measurement device designed for this experiment (Figure 3). Then the muscles were separated. The rectus femoris was divided into two parts: one part was frozen at -80 ℃ for protein molecular weight detection, while the other part was fixed in 4% paraformaldehyde for Sirius red staining. Knee mobility was measured after the separation of the muscles. The anterior joint capsule was divided into two parts: one part was frozen at -80 ℃ for protein molecular weight detection, while the other part was fixed in 4% paraformaldehyde for hematoxylin and eosin (H&E) staining. During the experiment, the rats were free to move within the cage with the immobilization device attached.

***Measurement of joint mobility***

A joint mobility meter was used to measure the joint motion of the left knee of 60 SD rats (Figure 3). The Kirschner wire was penetrated from the femoral neck parallel to the femur. Fixed the cruzi needle by magnetic suction removable metal clamp. The distal tibia was secured to the turntable with disposable plastic ties. The digital force gauge was secured to the slide. On the base of the equipment was a rope attached to the groove of the turntable and a digital dynamometer. The turntable moved when the drive wheel was turned to indirectly turn the tibia while the femur was stationary. The applied force was displayed on the screen of the digital force meter, and the angular change between the femur and tibia (the disk radius, the force arm) was constant and was calculated according to the scale of the turntable. Therefore, the force moment and the force size showed a linear relationship. The moment size and the angle also had a corresponding relationship. The applied torque was calculated by multiplying the force by the constant radius of the disk. Knee ROM was measured with 5.3 N-cm as the standard torque. This torque brings the knee close to its physiological limit but does not damage the soft tissue[11-13]. The mobility of each left knee was measured three times by two researchers, giving six measurements. The knee ROM before and after myotomy was measured to yield the total, myogenic, and arthrogenic contracture using a previously described method[14]. (1) Degree of total contracture = ROM before myotomy (knee joint in the control group)-ROM before myotomy (knee joint in the immobilization group); (2) degree of arthrogenic contracture = ROM after myotomy (knee joint in the control group)-ROM after myotomy (knee joint in the immobilization group); and (3) degree of myogenic contracture = degree of total contracture-degree of arthrogenic contracture. A patent application has been made for the self-made joint mobility meter (Patent No. ZL202120996643.1).

***Histological evaluation***

Specimens used for joint mobility assessment were used to evaluate the histology of the knee joint. After the ROM measurements, the left rectus femoris and anterior knee joint capsule were fixed in 4% paraformaldehyde (pH 7.4) at 4 ℃ for approximately 36 h. The specimens were embedded in paraffin. The rectus femoris specimens were cut into 5-µm coronal sections, while the joint capsule specimens were sectioned into 5-µm sagittal sections.

***Sirius red staining***

Rectus femoris sections were stained with Sirius red solution for 1 h (2610-10-8; Solarbio Life Science, Beijing, China) and rinsed with running water to remove the surface dye. Nuclei were stained with Mayer’s hematoxylin solution for 8 to 10 min and flushed with running water for 10 min. Sections were then conventionally dehydrated and sealed with neutral gum. The muscle collagen fiber density and muscle fiber cross-sectional area for each region were assessed using ImageJ software version 1.53a (National Institutes of Health, Bethesda, MD, United States, available at https://imagej.nih.gov/ij/). Histological analysis was performed on six rats in each group, with three slides for each rat.

***H&E staining***

The joint capsule sections were stained with H&E using the following steps. (1) Paraffin sections were dewaxed and then placed in xylene I for 10 min, xylene for 10 min, anhydrous ethanol I for 5 min, anhydrous ethanol for 5 min, 95% alcohol for 5 min, 90% alcohol for 5 min, 80% alcohol for 5 min, 70% alcohol for 5 min, and finally washed with distilled water. (2) Sections were stained with harris hematoxylin for 3-8 min, rinsed with tap water and differentiated with 1% ethanol hydrochloride for several seconds, and then rinsed with tap water again. The sections were returned to blue with 0.6% ammonia and rinsed with running water. (3) Sections were stained in eosin solution for 1-3 min. (4) To attain the dehydration seal, the sections were placed in 95% alcohol I for 5 min, 95% alcohol II for 5 min, anhydrous ethanol I for 5 min, anhydrous ethanol II for 5 min, xylene I for 5 min, and xylene II for 5 min; the sections were then removed from the xylene to dry and were sealed with neutral gum. (5) Microscopic examination, image acquisition, and histological analysis were performed on six rats in each group, with three slides for each rat.

***Immunohistochemistry***

The following steps were used to prepare the articular capsule sections by immunohistochemistry. (1) Paraffin sections were dewaxed in the same way as H&E staining. (2) The 3% hydrogen peroxide was dropped onto the slice tissue, incubated at room temperature for 15 min, and washed with phosphate-buffered saline (PBS) for 3 times, 3 min each time. (3) After wiping the slide dry, the diluted normal goat serum was added and the slide was sealed at room temperature for 30 min. (4) The first antibody was added and incubated in a wet box at 4 ℃ overnight (15 h). (5) After PBS washing, the sections were dried with absorbent paper, and streptavidin-horseradish peroxidase conjugate labeled goat anti rabbit/mouse secondary antibody was added and incubated at 37 ℃ for 30 min. (6) After PBS washing, the PBS solution was removed and 3,3'-diaminobenzidine color developing solution was freshly prepared and added to each slice. (7) Mayer’s hematoxylin re-staining was performed. (8) Finally, dehydration and sealing, microscopic examination, image collection and histological analyses were performed the same way as H&E staining.

***Proteomics analysis of muscle and joint capsule***

Protein immunoblotting (western blotting) was performed as follows. Total protein was extracted from the retained muscle and joint capsule samples. When 50-60 mg rectus femoris muscle was taken, the total tissue protein was extracted with 600 mL radio immunoprecipitation assay (RIPA) reagent (Tris-HCl [pH 7.4], 150 mmol/L NaCl, 1 mmol/L ethyleneiamineetraacetic acid, 1% Triton X-100, 1% sodium deoxycholate, 0.1% sodium dodecyl sulfate [SDS], and 1 mmol/L PMSF), and a protease inhibitor was added to the RIPA. Total proteins were separated by SDS-polyacrylamide gel electrophoresis and transferred to the polyvinylidene difluoride (PVDF) membrane. The PVDF was washed and immersed in 5% skim milk for 4 ℃ overnight. Membranes were incubated with anti-rat monoclonal antibody (1:10000-50000, cell signaling, United States) for 2 h at room temperature, and then washed three times with a Tris-buffered saline with 0.1% Tween 20 (TBST) solution (10 min/wash). The washed PVDF membrane was incubated with horseradish peroxidase-labeled goat anti-rat immunoglobulin G antibody (1: 10000-50000; Cell Signaling, Danvers, MA, United States) for 1 to 2 h at room temperature, washed with TBST (as described above), and then detected with enhanced chemiluminescence light-emitting liquid colored with energy autoexposure. The developing bands were analyzed by IPP software.

***Statistical analysis***

The results are expressed as the mean ± standard deviation. One-way analysis of variance was used to test the difference between groups. *P* < 0.05 was statistically significant. Statistical analyses were performed using IBM SPSS statistics software, version 22 (IBM Corp., Armonk, NY, United States).

**RESULTS**

Three rats in the immobilization-6 wk group experienced slippage of the immobilization device in the 2nd week of immobilization; the immobilization device was fixed the same day as the slippage occurred, and there was no further slippage until the end of the immobilization period. There was no death, lower limb necrosis, or other complications in any group.

***Total contracture, myogenic contracture, and arthrogenic contracture***

After 1 wk of immobilization, the degree of total contracture significantly differed between the control group and the immobilization groups (all *P* < 0.05). Compared with the immobilization-1 d, -2 d, and -3 d groups, the degree of total contracture was significantly greater in the immobilization-1 wk, -2 wk, -3 wk, -4 wk, -6 wk, and -8 wk groups (all *P* < 0.05). Total contracture degree was not significantly different between certain adjacent immobilization groups, *i.e.* between the immobilization-1 d and -2 d, immobilization-2 d and -3 d, immobilization-3 wk and -4 wk, immobilization-4 wk and -6 wk, and immobilization-6 wk and -8 wk groups. The degree of total contracture was increased with the immobilization time (Table 1).

The degree of myogenic contracture was significantly greater in the immobilization-1 wk group than the immobilization-3 d group (*P* < 0.05). The degree of myogenic contracture also significantly differed between the immobilization-1 wk and -2 wk groups (*P* < 0.05), but not between any other adjacent immobilization groups (*P* > 0.05). The results suggested that myogenic contracture occurred after 1 wk of immobilization and gradually stabilized after 2 wk of immobilization, whereas the subsequent joint contractures were mostly arthrogenic (Table 1).

After 1 wk of immobilization, the degree of arthrogenic contracture was significantly greater in the immobilization groups than the control group (*P* < 0.05). The degree of arthrogenic contracture significantly differed between the immobilization-3 d and -1 wk, immobilization-1 wk and -2 wk, and immobilization-2 wk and -3 wk groups (all *P* < 0.05), but not between the other adjacent groups (*P* > 0.05). The results suggested that arthrogenic contracture progressed from 1 to 3 wk of immobilization. However, the progression amplitude of the arthrogenic contracture began to weaken after 3 wk of immobilization, after 4 wk, the arthrogenic contracture is basically stable (Table 1).

***Histological evaluation of the Sirius red-stained sections***

Compared with the control group, the mean diameter of the rectus femoris was significantly decreased with the duration of immobilization in all immobilization groups except the immobilization-1 d and -2 d groups (*P* < 0.05). From 1 d to 3 wk of immobilization, the proportion of collagen fibers in the rectus femoris of the immobilization groups increased with the fixation time. The proportion of collagen fibers significantly differed between the control group and all immobilization groups except the immobilization-1 d, -2 d, and -3 d groups (*P* < 0.05). The proportion of collagen fibers in the rectus femoris stabilized in the immobilization-4w group, but was decreased in the immobilization-6 wk and -8 wk groups (*P* < 0.05 in all cases; Figure 4).

***Evaluation of the hematoxylin-eosin-stained sections and immunohistochemistry***

The largest synovial area in the sagittal plane was analyzed. H&E staining showed that the degree of synovial hyperplasia of the anterior joint capsule in the immobilization groups increased with the fixation time, with significant differences between adjacent groups, *i.e.* between the immobilization-1 d and -2 d, immobilization-3 d and -1 wk, immobilization-1 wk and -2 wk, and immobilization-2 wk and -3 wk groups (*P* < 0.05). The degree of synovial hyperplasia of the anterior joint capsule also significantly differed between the control group and all immobilization groups, except the immobilization-1 d group (*P* < 0.05; Figure 5). Immunohistochemistry showed that the change trend of phosphorylated mothers against decapentaplegic homolog (p-Smad2) was consistent with the degree of synovial hyperplasia of the anterior joint capsule, but the percentage of p-Smad2 in each group was significantly different from that in the control group immobilization-2 d (*P* < 0.05; Figure 6).

***Protein expression in the muscle and joint capsule***

After early joint immobilization, the expression level of transforming growth factor beta 1 (TGF-β1) was significantly increased in the joint capsule and muscles. The expression level of TGF-β1 in the joint capsule was significantly higher in the immobilization-1 wk group than in the immobilization-3 d group. TGF-β1 expression of joint capsule continued to increase after immobilization for up to 8 wk. There were significant differences in TGF-β1 levels in the joint capsule between the immobilization-2 d and -3 d, immobilization-1 wk and -2 wk, and immobilization-2 wk and -3 wk groups (*P* < 0.05). The TGF-β1 content of the anterior joint capsule increased with the immobilization time, with a slight downward trend in the immobilization-8w group that did not reach statistical significance (*P* > 0.05). TGF-β1 expression in the anterior joint capsule was significantly increased in the control group compared with the immobilization groups (*P* < 0.05), except the immobilization-1 d and -2 d groups. Western blot analysis showed that the TGF-β1 content in the quadriceps muscle first increased and then decreased. The TGF-β1 content in the muscle significantly differed between the immobilization-2 d and -3 d, immobilization-3 d and -1 wk, immobilization-1 wk and -2 wk, and immobilization-6 wk and -8 wk groups (*P* < 0.05); The contents of Smad2 and Smad3 and TGF-β1 in quadriceps femoris had the same trend of change, but there were significant differences only between the groups of immobilization-3 d and -1 wk, immobilization-1 wk and -2 wk (*P* < 0.05). In all of the results, with the exception of the fixed-1 d and-2 d groups, there was also a significant difference between the control group and the fixed group (*P* < 0.05) (Figure 7).

**DISCUSSION**

Joint contracture is a relatively common condition that is mainly caused by fibrosis of the joint capsule and skeletal muscle after long-term immobilization, and shows the pathological features of excessive deposition of collagen and connective tissue components[15]. Numerous animal models have been developed to simulate knee flexion contracture; however, few animal models of knee extension contracture have been reported. A previous study successfully established a model of knee extension contracture in New Zealand white rabbits and reported in detail the characteristics of plaster immobilization and the relevant mechanisms[16]. However, it is beneficial to model knee extension contracture in rats rather than rabbits because of the lower cost of studying the pathogenesis of joint contracture and evaluating therapeutic efficacy. Therefore, the present study described a method to establish a model of knee extension contracture in the rat. We demonstrated that this model had significantly limited knee flexion motion and altered expressions of histological and fibrosis-related proteins in the skeletal muscle and anterior joint capsule.

Wang *et al*[17] directly assessed the muscle limitations of rats with immobilized ankle joints and found that the initial flexion contracture of the knee is mainly due to muscle structures and is reversible and can spontaneously resolve. By contrast, long-term contracture is mainly caused by the joint structures and is irreversible[14]. Such arthrogenic contracture cannot be improved, even by aggressive rehabilitation[18]. Several reports suggest that joint contractures occur within 1 wk of immobilization and progress in a time-dependent manner[19,20]. Chimoto *et al*[21] reported that 2 wk of muscle limitation mainly causes myogenic contracture, while long-term contracture (more than 4 wk of immobilization) results in joint contracture. Therefore, prolonged immobilization for longer than 4 wk should be avoided to prevent irreversible joint contracture[22]. In the present study, myogenic contracture was the predominant type of contracture during the first 2 wk of immobilization. From 2 to 3 wk of immobilization, the joint contracture changed from myogenic to arthrogenic. The contracture initiation time in the present study was consistent with previous studies; however, in contrast with previous studies, the arthrogenic contracture stabilized at 3 wk and completely stable at 4 wk. Arthrogenic contracture is primarily a fibrotic response within the joint capsule. The posterior joint capsule is the main contributor to the formation of immobilization-induced knee flexion contracture, while the anterior joint capsule has the greatest impact on knee extension contracture[21]. The synovial layer of the anterior joint capsule is the widest and most complex in the knee joint[23]. In the present study, the degree of synovial hyperplasia continuously increased with the immobilization time; this may explain why knee extension contracture forms earlier than knee flexion contracture.

In the histologic assessment of the present study, the myofiber cross-sectional area, intermuscular collagen deposition, and extent of hyperplasia in the anterior joint capsule supported the biological findings. As the decreased skeletal muscle mass caused by an imbalance in protein metabolism is characterized by a significantly smaller muscle fiber area[24-26]. The amount of intermuscular collagen deposition was significantly greater in the immobilization-1w group than the control group, but did not significantly differ between the immobilization-2 wk and -3 wk groups. Furthermore, there was significantly less intermuscular collagen deposition in the immobilization-6w group compared with the immobilization-4w group. This indicates that intermuscular collagen deposition began to increase in the 1st week of immobilization, but decreased after 4 wk of immobilization. Previous studies have shown that this may be associated with gradual resolution of the inflammatory and fibrotic response after immobilization[11]. This was demonstrated by the reduction in collagen content and TGF-β1 protein expression over time. The anterior joint synovial proliferation was significantly greater in the immobilization-2 d group than the control group, but did not differ between the immobilization-3 wk and -4 wk groups, indicating that the anterior joint synovial proliferation significantly increased during the first 3 wk of immobilization and completely stabilized from 4 wk onwards. Wang *et al*[27] analyzed changes in the synovial membrane caused by anterior articular capsule fibroblasts, microvasodilation, and congestion due to plaster immobilization. Li *et al*[28] and Yao *et al*[29] reported that intra-articular tissue adhesion does not completely cover the cruciate ligament in and around the knee, but originates from synovial fibrosis. The proliferation of intra-articular synovial tissue was responsible for the limited ROM found in the present study.

To further characterize the altered fibrosis of skeletal muscle and the joint capsule, we evaluated the expression of the TGF-β1 and p-Smad protein. There are three subtypes of TGF-β family: TGF-β1, TGF-β2, and TGF-β3. A large number of studies have demonstrated that the expression of TGF-β1 is the most important in fibrosis changes. TGF-β1 can cause differentiation, proliferation and extracellular matrix production, promote the transformation of fibroblasts to myofibroblasts, cause the deposition of collagen and increase the expression of alpha smooth muscle actin (α-SMA), and then cause tissue fibrosis[30,31]. Zhang *et al*[32] found that the expression level of TGF-β1 in the posterior joint capsule was significantly increased *in vivo* experiments. *In vitro* experiments showed that the expression level of TGF-β1 was increased, and the expression of α-SMA and type I collagen was also increased. Mao *et al*[33] applied a TGF-β inhibitor to treat a traumatic rat knee joint contracture animal model and found that after the activity of TGF-β was inhibited, the degree of joint contracture was alleviated, while the protein expression of collagen type I, collagen type III, α-SMA, and p-Smad2 was decreased. Therefore, we used the TGF-β1 and Smad signaling pathway as indicators to prove the formation of contracture. The aggravation of contracture enhances the fibrosis response, characterized by increases in profibrotic genes and proteins (*e.g.,* cytokine TGF-β1 genes, type I and type III collagen genes and proteins), leading to increases in collagen density and joint capsule thickness[34]. Hildebrand *et al*[35] reported the increased expression of type I and III collagen, TGF-β1, p-Smad2, and p-Smad3 in a rabbit post-traumatic flexion contracture model compared with control joint cysts. The increased mRNA levels of TGF-β1 may be related to collagen deposition inside and outside the articular capsule. Joint capsule fibrosis may be associated with the development of joint contracture. Similarly, the present study showed that the expression of fibrosis-related genes increased with prolonged immobilization, but the expression of TGF-β1, p-Smad2, and p-Smad3 in the rectus femoris decreased slightly after 3 wk of immobilization and was significantly decreased at 8 wk. The altered expression levels of TGF-β1 may be due to hypoxia or a reduction in collagen turnover or degradation rates[36]. Among several other roles, one of the adaptive responses of hypoxic cells is the upregulation of hypoxia-inducible factor 1 alpha (HIF-1α). The expression of TGF-β1 and HIF-1α is significantly upregulated during the transformation of fibroblasts to myofibroblasts, leading to the promotion of vascular endothelial growth factor (VEGF) gene expression[37]. The HIF-1α signaling pathway in turn regulates angiogenesis by inducing VEGF expression, thereby improving circulation and reducing the inflammatory response[38].

In the present study, ordinary gypsum and polymer gypsum were initially used to establish the extended knee contracture model. However, the rats inevitably gnawed the gypsum and there was also gypsum slippage. In the process of switching to an aluminum splint, we found that the knee joints of the rats could not be wrapped when fixed because the proximal lower limbs were short and strong. The present shape of the external immobilization device was determined after multiple improvements. The immobilization device comprised of aluminum plate pressurized at the distal end on the back of the foot that was plantarflexed at 60. After the first proximal rectangular aluminum plate was used to fix the knee joint, the second inverted trapezoidal aluminum plate was wrapped around the knee to ensure that the knee was completely immobilized. Although this immobilization method is simple, the pressure strength must be carefully controlled. A pressure that is too high will easily cause poor limb circulation in rats. As aluminum is easy to shape, we were able to adjust the tightness of the external immobilization to resolve any swelling. Tokuda *et al*[39] successfully created a model of flexion contracture outside the knee joint; however, the external immobilization device used in the present study had less effect on the overall activity of the rats and better reflected the clinical situation in which the knee joint is usually fixed in extension after injury, leading to limited knee flexion after long-term immobilization. Therefore, to more closely mimic the clinical situation, we chose to create a model of knee extension contracture.

We demonstrated that the present model is as reliable as other animal models in reproducing the features of human joint contracture, including limited joint mobility, changes in the joint and muscle histology, and changes in the expression of fibrosis-associated proteins in the joint capsule *vs* muscle. The advantage of the present model is that it is easy to replicate because it does not require complex surgical procedures, the tools are easy to use, and the rat anesthesia and immobilization can be performed in a very short period of time. We described the detailed process of establishing a rat model of knee extension contracture, with photographs. The model closely replicates joint contracture caused by complications of immobilization, enabling researchers to investigate the etiology of joint contracture and establish new treatments. This model is a reliable tool, as described earlier. Contractures caused by long-term fixation are mainly caused by joint structure and are irreversible. Moreover, the mechanism by which movement after fixation may exacerbate joint contracture has not been fully explored[40]. As the external fixator is easy to shape and can be removed at any time. Therefore, this model can be used to study the prevention and treatment of knee extension contracture in rats, at the same time, it is possible to change the dressing and keep the wound dry during the fixation of traumatic knee joint contracture.

The present study had some limitations. First, in this model, the ankle joint and the knee joint were inevitably fixed together. Because the lower limb of the rat is shaped like a cone, the ankle joint was plantarflexed at 60 and fixed with the knee joint to prevent slippage of the aluminum splint. Second, the longest immobilization time in the present study was 8 wk. We plan to explore the continuous longer-term changes in fibrosis-related proteins in a future study. Finally, the state of the extension contracture after the removal of the external fixator was also studied and will be reported.

**CONCLUSION**

The results in this study suggest that the myogenic contracture is stabilized after 2 wk, whereas the arthrogenic contracture is stabilized after 3 wk and completely stable in 4 wk.

**ARTICLE HIGHLIGHTS**

***Research background***

There is currently no research on establishing a model of knee joint extension contracture in rats.

***Research motivation***

The extension contracture model is of clinical relevance because it better mimics fracture and bed-associated immobilization than the flexion contracture model.

***Research objectives***

Clarify the formation process of knee joint extension contracture in rats.

***Research methods***

Verify the formation process of extension contracture by observing pathology, detecting fibrotic proteins, and measuring joint range of motion.

***Research results***

All results show that the myogenic contracture tends to stabilize after 2 wk, and the arthrogenic contracture tends to stabilize after 3 wk and completely stable in 4 wk.

***Research conclusions***

The extension contracture model is of clinical relevance because it better mimics fracture and bed-associated immobilization than the flexion contracture model.

***Research perspectives***

This rat model may be a useful tool for studying the etiology of joint contracture and establishing new treatment methods.

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**Footnotes**

**Institutional animal care and use committee statement:** All animal experiments conformed to the internationally accepted principles for the care and use of laboratory animals (Anhui Medical University; protocol no. LLSC20221126, Experimental Animal Ethics Committee of Anhui Medical University, Anhui, China).

**Conflict-of-interest statement:** The authors have no conflicts of interest to declare.

**Data sharing statement:** No additional data are available.

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Grade D (Fair): 0

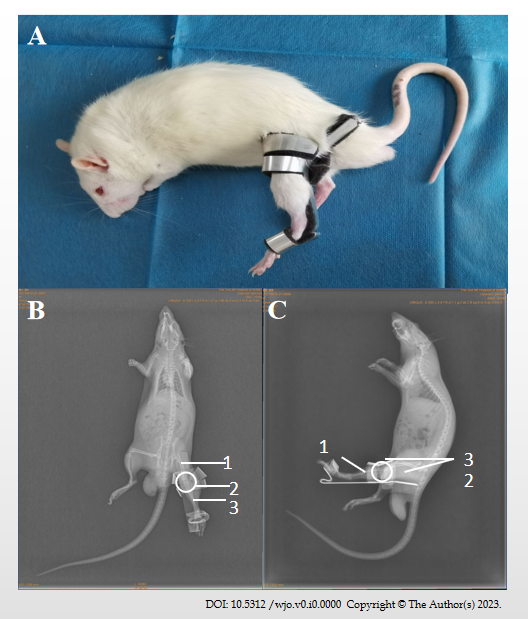
Grade E (Poor): 0

**P-Reviewer:** Malkova TA, Russia; Nambi G, Saudi Arabia; OOMMEN AT, India **S-Editor:** Qu XL **L-Editor:** Filipodia **P-Editor:**

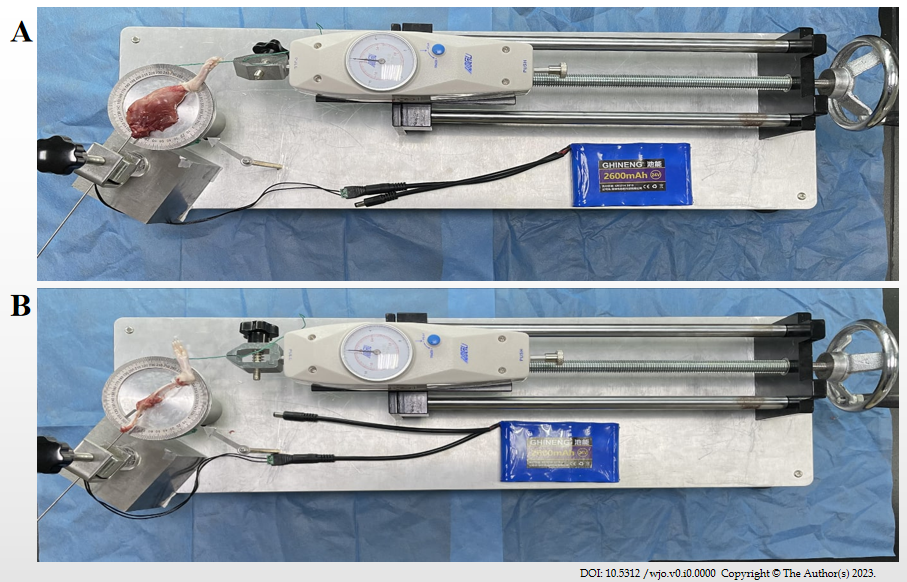
**Figure Legends**



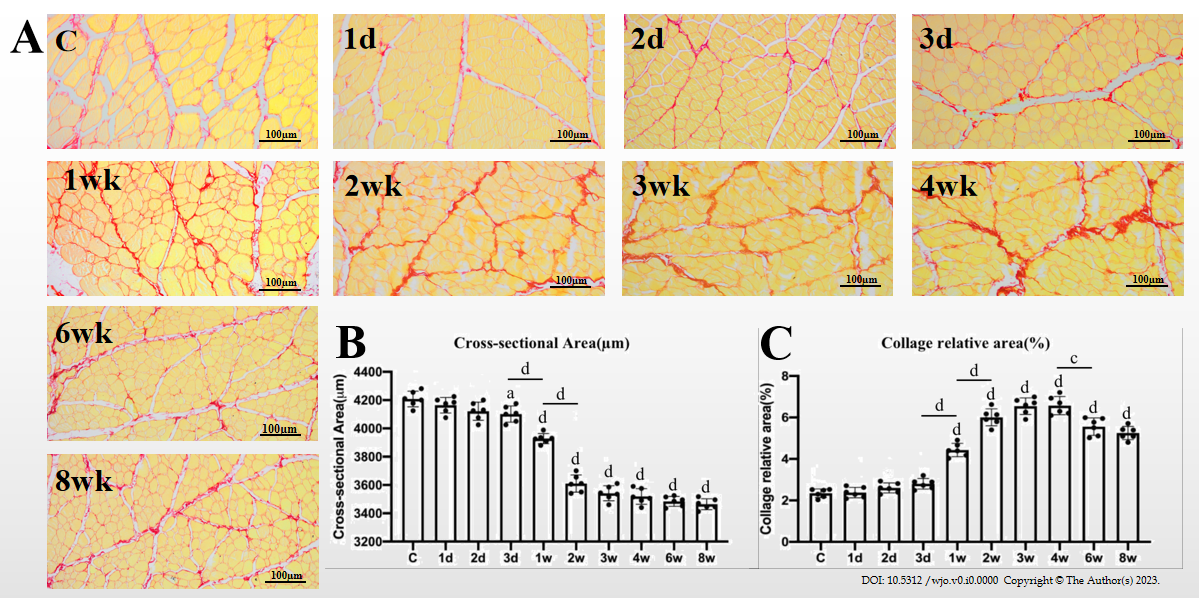
**Figure 1 Knee joint fixation device.** Immobilize the back of the rat’s distal foot to prevent the rat’s lower limb from slipping out of the immobilization device. Immobilize the periphery of the rat knee joint. Above the knee joint of the rat is an inverted conical structure, and this design fixes the periphery of the femur of the rat. Appropriate bending under the rat femur provides a lever fulcrum action.



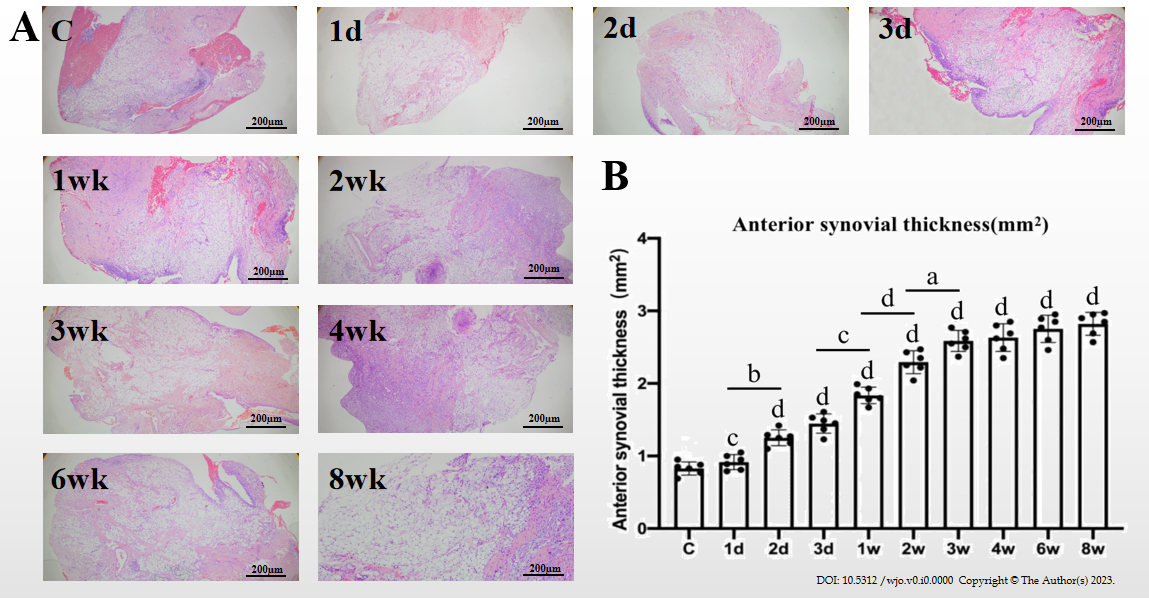
**Figure 2 Fixed schematic diagram.** A: Fixed picture; B: X-ray orthotopic slice of rat knee joint after immobilization; C: X-line lateral tablet of rat knee joint after immobilization. 1Femur; 2Knee; 3Tibia.



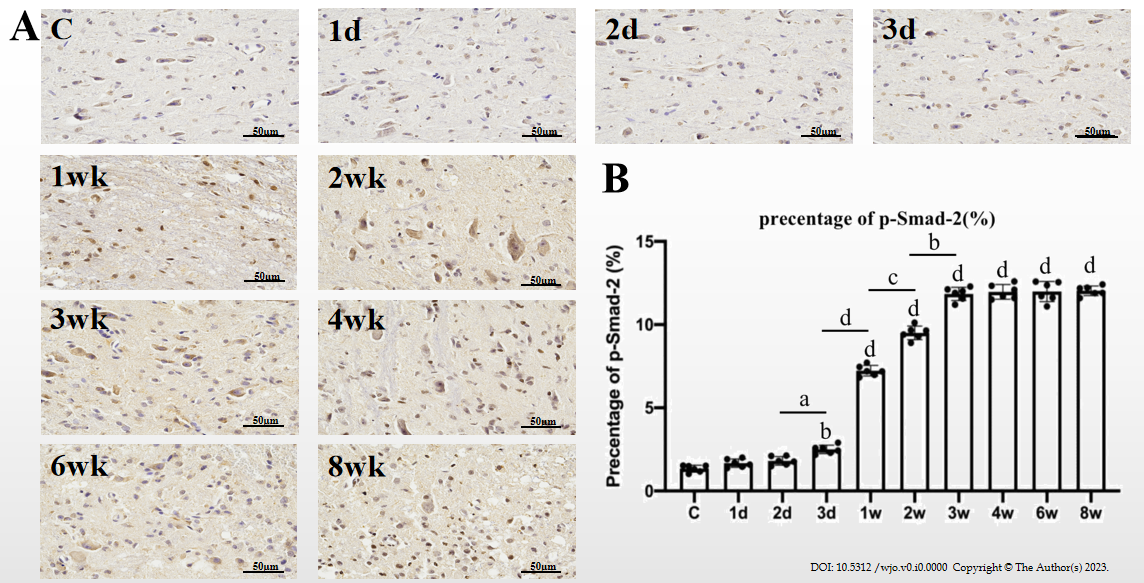
**Figure 3 Joint mobility meter.** A: Peeling range of motion (total contracture) measurement; B: Measuring range of motion after muscle separation (arthrogenic contracture).



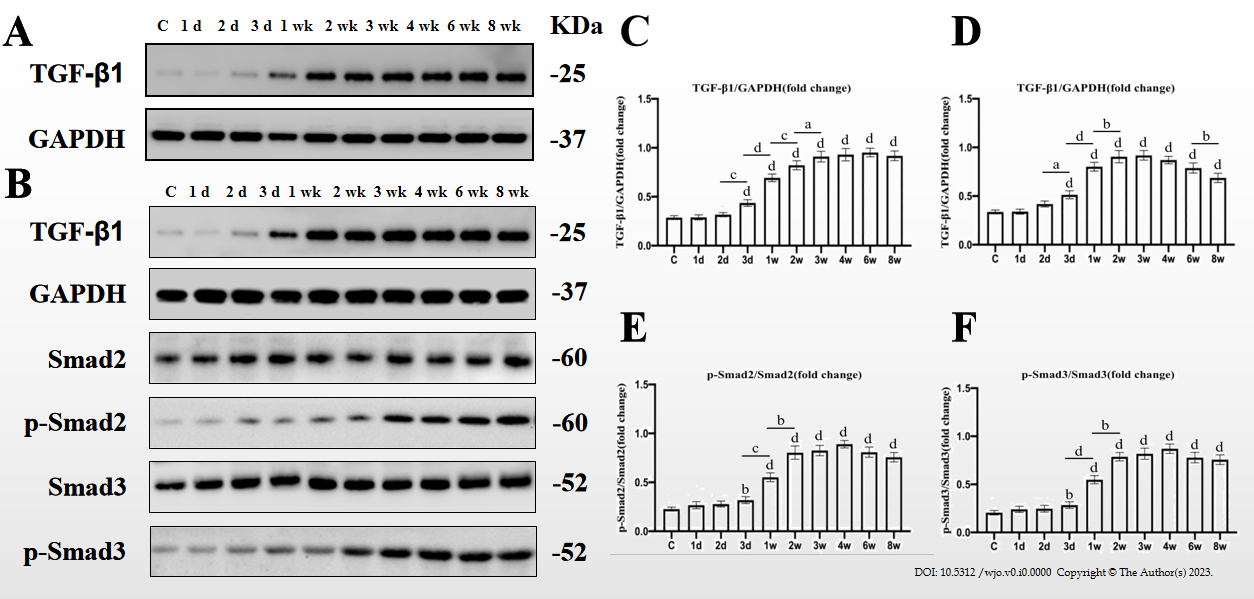
**Figure 4 Sirius red staining findings.** A: Morphological changes of rectus femoris after Sirius red staining, group C (control), 1 d, 2 d, 1 wk, 2 wk, 3 wk, 4 wk, 6 wk, and 8 wk; B and C: Cross-sectional value of rectus femoris fibers. a*P* < 0.05; c*P* < 0.001; d*P* < 0.0001.



**Figure 5 Hematoxylin and eosin staining findings.** A: Morphological changes of the anterior joint capsule; B: The anterior joint capsule thickness value. a*P* < 0.05; b*P* < 0.01; c*P* < 0.001; d*P* < 0.0001.



**Figure 6 Immunohistochemistry findings.** A: Immunohistochemical staining of the anterior joint capsule; B: Percentage of the anterior articular joint capsule phosphorylated mothers against decapentaplegic homolog 2. a*P* < 0.05; b*P* < 0.01; c*P* < 0.001; d*P* < 0.0001.



**Figure 7 Western blot findings.** A: Changes in the intensity of transforming growth factor beta 1 (TGF-β1) and glyceraldehyde 3-phosphate dehydrogenase (GAPDH) bands in the anterior joint capsule; B: Changes in the intensity of TGF-β1/GAPDH, phosphorylated mothers against decapentaplegic homolog 2 (p-Smad2)/Smad2, p-Smad2/Smad2 bands in the quadriceps; C: Graphical representation of the expression level of TGF-β1 in the anterior joint capsule relative to GAPDH; D-F: Graphical representation of the expression level of TGF-β1/GAPDH, p-Smad2/Smad2, p-Smad2/Smad2. a*P* < 0.05; b*P* < 0.01; c*P* < 0.001; d*P* < 0.0001.

**Table 1 Total, myogenic, and arthrogenic contracture after knee immobilization for various time periods in 60 SD rats (mean ± SD)**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Grouping** | **Quantity** | | **Degree of contracture** | | |
| **Total contracture** | **Myogenic contracture** | **Arthrogenic contracture** |
| Control | | 6 | 0.0 ± 0.0 | 0.0 ± 0.0 | 0.0 ± 0.0 |
| Immobilization-1 d | | 6 | 1.1 ± 0.6 | 0.6 ± 0.4 | 0.5 ± 0.3 |
| Immobilization-2 d | | 6 | 2.4 ± 0.7 | 1.5 ± 0.4 | 0.9 ± 0.2 |
| Immobilization-3 d | | 6 | 5.9 ± 1.0 | 4.0 ± 1.0 | 1.9 ± 0.5 |
| Immobilization-1 wk | | 6 | 23.5 ± 2.0a,b,c,d | 15.9 ± 1.9a,b,c,d | 7.6 ± 0.7a,b,c,d |
| Immobilization-2 wk | | 6 | 51.8 ± 1.8a,b,c,d,e | 32.7 ± 1.1a,b,c,d,e | 19.1 ± 1.3a,b,c,d,e |
| Immobilization-3 wk | | 6 | 78.7 ± 2.2a,b,c,d,e,f | 33.9 ± 2.1a,b,c,d,e | 44.5 ± 1.9a,b,c,d,e,f |
| Immobilization-4 wk | | 6 | 79.9 ± 2.8a,b,c,d,e,f | 34.2 ± 2.2a,b,c,d,e | 44.9 ± 2.2a,b,c,d,e,f |
| Immobilization-6 wk | | 6 | 80.8 ± 3.5a,b,c,d,e,f | 35.5 ± 2.9a,b,c,d,e | 45.3 ± 2.9a,b,c,d,e,f |
| Immobilization-8 wk | | 6 | 82.5 ± 3.0a,b,c,d,e,f | 35.9 ± 2.5a,b,c,d,e | 46.6 ± 2.3a,b,c,d,e,f |

a*P* < 0.05 *vs* control group; b*P* < 0.05 *vs* immobilization-1 d group; c*P* < 0.05 *vs* immobilization-2 d group; d*P* < 0.05 *vs* immobilization-3 d group; e*P* < 0.05 *vs* immobilization-1 wk group; f*P* < 0.05 *vs* immobilization-2 wk group.