Inhibition of connexin 43 prevents trauma-induced heterotopic ossification

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Heterotopic ossification (HO) can result from traumatic injury, surgery or genetic diseases. Here, we demonstrate that overexpression of connexin 43 (Cx43) is critical for the development and recurrence of traumatic HO in patients. Inhibition of Cx43 by shRNA substantially suppressed the osteogenic differentiation of MC-3T3 cells and the expression of osteogenic genes. We employed a tenotomy mouse model to explore the hypothesis that Cx43 is vital to the development of HO. Inhibition of Cx43 by a specific shRNA decreased extraskeletal bone formation in vivo. In addition, we demonstrated that ERK signaling activated by Cx43 plays an important role in promoting HO. ERK signaling was highly activated in HO tissue collected from patient and mouse models. Importantly, de novo soft tissue HO was significantly attenuated in mice treated with U0126. Inhibition of Cx43 and ERK led to decreased expressions of Runx2, BSP and Col-1 in vivo and in vitro. Moreover, HO patients with low Cx43 expression or ERK activation had a lower risk of recurrence after the lesions were surgically removed. Our findings indicate that Cx43 promotes trauma-induced HO formation by activating the ERK pathway and enhances the expression of osteogenic markers.

Heterotopic ossification (HO), which is commonly observed in cases of traumatic injury, severe burns and invasive surgeries, is the aberrant formation of extraskeletal bone in soft tissues such as muscles, tendons, and ligaments1. Traumatic HO occurs in response to injuries such as fractures and fracture-dislocations of joints. Treatment options for HO are limited because the excess bone growth often recurs following surgical resection, and some patients may have HO that is nonresectable because of its location2. For several years, the genetic basis of HO has been known to be similar to that of fibrodysplasia ossificans progressiva (FOP), however, the mechanism of HO development is still unclear. The pathophysiology of HO closely resembles the physiologic process of fracture healing3. In the case of HO formation, the osteogenesis process is thought to arise from the disordered differentiation and proliferation of precursor cells. Previous findings have shown that progenitor cells obtained from traumatized muscle tissue, which is prone to developing HO, are similar to bone marrow-derived mesenchymal stem cells (MSCs) in morphology and biological function4,5.

A number of research groups have reported several types of evidence showing that the gap junction protein connexin 43 (Cx43, also called gap junction protein A1, GJA1) controls bone function and development6,7. Cx43, a gap junction channel protein that mediates the communication between adjacent cells or between a cell and its extracellular environment, is observed to be upregulated under fracture conditions8. In addition, fracture healing is impaired in Cx43-deficient mice, which is consistent with delayed osteoblast differentiation and bone formation9. These observations suggest that Cx43 facilitates new bone formation. Overexpression of Cx43 in osteoblasts promotes the transcriptional activity of Runx2, and Cx43 knockdown suppresses Runx2-dependent transcription10. The signaling molecules mediating the action of Cx43 on osteogenesis include the protein kinase C (PKC) family member PKCδ and the extracellular signal regulated kinases (ERKs), two proteins that were previously reported to regulate Runx2 activity in osteoblasts11,12. Given the correlations between Cx43 and bone formation, we hypothesized that Cx43 may enhance the development of traumatic HO by interacting with ERK signaling.

In this study, we demonstrated that Cx43 contributes to the development of traumatic HO. Cx43 knockdown substantially attenuated the expression of osteogenic marker genes and markedly inhibited the growth of HO

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in a mouse model. Cx43 promoted osteogenesis through the ERK pathway. Inhibition of ERK also resulted in decreased HO development. Moreover, traumatic HO patients with low Cx43 expression and ERK activation showed decreased recurrence after surgical removal of the lesions. The data in the present study may provide potential therapeutic target for traumatic heterotopic ossification.

**Results**

**Expression of Cx43 is increased in human heterotopic ossification tissues.** To investigate the genes that may be involved in the regulation of HO formation, we used genomic data from 5 patients with elbow HO to compare with data from the control bones, which were collected from 5 traumatic amputation patients. In particular, we noted upregulation of Cx43 and osteogenic marker genes in the heterotopic ossification samples (Fig. 1A). Previous reports have underscored the importance of Cx43 in osteoblast differentiation and skeletal development13. Next, the expression levels of Cx43 and osteogenesis-related genes were further confirmed by real-time PCR. We verified that the expression of Cx43 and osteogenic markers, including Runx2, alkaline phosphatase (ALP), bone sialoprotein (BSP) and collagen-1 (Col-1), were increased in HO patients compared with control patients (Fig. 1B). However, no significant difference in Dlx-5 expression was observed between the two groups. Furthermore, soft tissues (joint capsule and ligament) around the HO or near the normal joint (used as
normal control) were collected from the HO or traumatic amputation patients. We observed an elevated expression of Cx43, IL-1, IL-6 and IL-8 in the soft tissue surrounding the HO (Supplementary Fig. 1).

**Cx43 promotes osteoblast differentiation by activating the ERK pathway.** Aberrant osteogenesis plays an important role in the formation of HO. To determine the effect of Cx43 on osteoblast differentiation, expression of Cx43 was inhibited by a Cx43-specific shRNA transfection. The Cx43 shRNA transfection did not affect the viability of MC-3T3 and C2C12 cells (Supplementary Fig. 2). The inhibitory effect was confirmed by western blot (Fig. 2A). Then, MC-3T3 cells transfected with Cx43 shRNA were exposed to osteogenic differentiation medium for 2 weeks. We observed that both calcium deposition and ALP activity were decreased in the Cx43 knockdown group (Fig. 2B,C). Furthermore, our real-time PCR results showed that Cx43 knockdown inhibited osteogenic marker gene expression (Fig. 2D). A similar result was observed in mouse mesenchymal stem cells (Supplementary Fig. 3A–D) Previous studies have suggested that ERK is part of an important downstream pathway of Cx43. We then examined whether the ERK pathway played a role in Cx43-induced osteogenesis.

**Figure 2.** Cx43 promotes osteoblast differentiation by activating the ERK pathway. (A) Small hairpin RNAs (shRNA) were used to knock down Cx43 expression in MC-3T3 cells, and a scrambled sequence was used as a control. The expression of Cx43 was detected by Western blot. (B) MC-3T3 cells transfected with Cx43 shRNA were induced in osteogenic medium. Alizarin Red S (week 2) and ALP (week 1) staining was performed. (C) Calcium deposits and ALP activity were quantified. (D) mRNA expression of osteogenic marker genes was detected by real-time PCR at week 1 and 2. (E) ERK pathway activation and Cx43 expression were examined weekly by Western blot. (F) A densitometric analysis of p-ERK expression was performed (t-test, n = 3, *p < 0.05; NS, not significant).
Our data indicated that ERK signaling was activated during the osteogenesis process, meanwhile expression of Cx43 was increased during osteogenesis. However, when Cx43 was knocked down by shRNA, p-ERK expression was maintained at a steady-state during osteogenesis (Fig. 2E,F). To exclude the effects of osteogenic medium on ERK activation, MC-3T3 cells were incubated in α-MEM without osteogenic induction, and Cx43 was knocked down by shRNA. We observed that activation of the ERK and PKC pathway was inhibited at week 1–3 (Supplementary Fig. 4A,B). However, inhibition of ERK by U0126 or shRNA had no effect on expression of Cx43 (Supplementary Fig. 4C).

Inhibition of Cx43 limits HO formation in vivo. Next, we tested the hypothesis that Cx43 inhibition can prevent HO. For this purpose, mice received a tenotomy were subsequently treated with an injection of Cx43 shRNA at the lesion site. We observed a reduced amount of early endochondral ossification in mice treated with Cx43 shRNA at 3 weeks (Supplementary Fig. 5). Then, the injured limbs were examined by x-ray. The tenotomy mice treated with Cx43 shRNA injection demonstrated a reduction in total HO volume at 4 and 8 weeks after injury (Fig. 3A,B). Consistent with the radiologic data, representative images of 3D μCT reconstructions of a tenotomy mouse showed markedly decreased HO volume after Cx43 shRNA injection (Fig. 3C,D). Moreover, the bone mineral density (BMD) of HO was decreased after the Cx43 shRNA treatment (Fig. 3E). Tissues from the tenotomy site 8 weeks after injury showed diminished levels of Cx43, PERK and osteogenic marker genes, including the early osteogenic gene Runx2 and the later gene BSP (Fig. 3F,G). To test whether Cx43 could inhibit the developed HO, tenotomy mice were maintained for 8 weeks and then treated with Cx43 shRNA injection for another 4 weeks. The x-ray images showed no difference between the two groups at week 12 (Supplementary Fig. 6).

ERK activation is essential for Cx43-induced HO. To investigate whether inhibition of the ERK pathway would limit HO, MC-3T3 cells exposed to U0126 were incubated in osteogenic medium for 2 weeks in vitro. When ERK signaling was inhibited by U0126, Alizarin Red S staining showed less calcium deposition (2 weeks) and less ALP activity (1 week) (Fig. 4A,B). Similarly, inhibition of ERK signaling by a specific shRNA suppressed osteogenic differentiation (Supplementary Fig. 7A,B). Then, tenotomy mice were treated with an injection of U0126 at the lesion sites or intraperitoneal injection (IP). Both injection methods decreased the HO formation, while no difference was observed between the local and intraperitoneal injection groups (Fig. 4C,D). As observed in the mice that received the Cx43 shRNA injection, the 3D μCT reconstruction images showed a decreased HO volume after the U0126 local injection (Fig. 4E,F). We observed a decreased BMD in the HO after the Cx43 shRNA injection (Fig. 4G).

Cx43 enhances the expression of osteogenic markers through ERK. Then, the HO samples from the mice were collected and demineralized. The immunohistochemistry results demonstrated that a Cx43 shRNA injection caused decreased expression of Cx43, p-ERK, BSP and Col-1. Inhibition of ERK signaling by U0126 only decreased the expression of p-ERK, BSP and Col-1, while no change in the Cx43 expression was observed. These data suggest that Cx43 enhances the expression of osteogenic markers by activating ERK (Fig. 5A,B).

Cx43/ERK signaling is critical for clinical outcomes in HO patients. To verify the effects of Cx43/ERK signaling on clinical outcomes, HO lesions were resected from elbow trauma patients (n = 20), and bones from amputation patients were used as a control (n = 15). All bones were demineralized and subjected to an immunohistochemical assay. We observed that HO tissues exhibited increased Cx43 and p-ERK expressions (Fig. 6A,B). According to the Cx43 and p-ERK staining density, the HO patients were divided into weak (<0.5, n = 9) and strong (>0.5, n = 11) groups. The 3D CT showed that strong expression of Cx43 was correlated with increased HO volume (Fig. 6C,D; p = 0.016). Furthermore, we observed that HO patients with strong Cx43 expression showed a higher recurrence rate after surgical treatment (Fig. 6E). Similar results were observed in the strong p-ERK HO patients (Fig. 6F). In addition, tissues from the mice showed that Cx43 was highly expressed in endochondral bone and HO (Supplementary Fig. 8A). Soft tissue around the recurrent HO from the patients exhibited a higher Cx43 baseline than soft tissue around the non-recurrent HO (Supplementary Fig. 8B).

Discussion
HO is a pathologic process in patients with severe musculoskeletal trauma and burns and with genetic mutations that confer hyperactive ossification. Although the exact mechanism of ectopic bone formation is still unknown, progenitor cells capable of osteogenic differentiation are known to play an essential role13. In this study, we leveraged our knowledge to demonstrate that Cx43/ERK signaling represents a route for traumatic ectopic bone formation. The present study demonstrates that Cx43/ERK might be essential for ectopic bone formation in a mouse model, which is highly consistent with our clinical observations. These findings suggest that targeting Cx43/ERK pathway may represent a solution for traumatic HO.

Previous studies have elucidated the genetic basis of HO as fibrodysplasia ossificans progressiva (FOP), a genetic disease that causes progressive HO. Genetic mutations in the ACVR1 gene conferring hyperactivity is regarded as the most important cause for FOP15,16. Although the genetic basis of FOP has been known for several years, the identity of the precursor cells that actually differentiate along the osteogenic lineage to form HO remain unclear17. Several studies have demonstrated that vascular endothelial cells could be a candidate for the cellular origin of HO18,19, while other studies provide strong evidence that tissue-resident MSCs or progenitor cell populations are the original cells in HO20,21. In trauma and wounds, MSCs in the tissue and circulation contribute to wound and fracture healing, which requires osteogenic differentiation of MSCs22. A recent study showed that activated matrix metalloproteinase 9 contributes to HO23. Agarwal S et al. found that pharmacologic inhibition of Hif1α abolished HO formation21.
Gap junctions, which are specialized intercellular membrane channels, are highly dynamic structures that are regulated by kinase-mediated signaling pathways and interactions with other proteins. In the bone tissue, the most abundantly expressed gap junction protein is Cx43, which is expressed in osteoblasts, osteocytes and osteoclasts. Previous studies have shown that alterations in Cx43 expression or function modulate the expression of
osteoblast genes. It has been shown that Cx43 expression is essential for normal osteoblastic gene expression and function in osteoblastic cell lines. Thi et al. demonstrated that mineralization is impaired in osteoblastic cells isolated from Cx43 knockout mice. In addition, osteogenic differentiation markers including Runx2, osteocalcin (OCN), collagen 1 and BSP are decreased in osteoblastic cells isolated from Cx43 knockout mice and in osteoblast cells overexpressing Cx45, which acts as an antagonist for Cx43. Evans KN et al. showed that multiple osteogenesis-related genes are upregulated in traumatic HO. These studies suggested that Cx43 might play...
an important role in HO formation. Using a model of trauma-induced HO, we provided evidence that overexpression of Cx43 plays an important role in the development of HO, in which Cx43 exerts its effects by promoting the expression of the osteogenic markers.

Endochondral ossification is a common process in HO formation. Previous studies showed that activation of β-catenin stimulates endochondral ossification. Consistent with their findings, we observed upregulated Wnt-3a expression in the gene chip. These data suggest that Wnt/β-catenin may play a role in HO formation. Moreover, our histologic analysis showed that Cx43 inhibition decreased early endochondral ossification. However, once the HO was completely developed, inhibition of Cx43 could not decrease the mature ectopic bones. These data suggest that inhibition of Cx43 at the early stage of HO may be benefit for the treatment of HO. Additionally, we demonstrated that HO patients with lower expression of Cx43 exhibit less recurrence after the surgical resection. These results suggested that Cx43 base line may affect the primary and recurrent HO formation.

Cx43-mediated regulation of osteoblast differentiation and gene expression involves transcription factors Runx2 and Osterix (Sp7), both of which are master regulators of osteogenesis. Disruption of the Cx43 channels by overexpression of Cx45 or by specific inhibitors in osteoblast cells decreases activation of ERK signaling. Moreover, decreased ERK activity leads to reduced phosphorylation and DNA binding to the transcription factors, resulting in reduced transcription of the osteogenic marker genes. Similarly, our data showed that ERK signaling was activated and that Runx2 expression was enhanced in the HO tissues. Our findings indicate that Cx43 and osteogenic medium have different effects on ERK activation. On one hand, p-ERK expression was enhanced during osteogenesis, but on the other hand, inhibition of Cx43 decreased the ERK activation. Although Cx43 shRNA did not completely abolish the p-ERK expression, it maintained the p-ERK expression at the un-differentiated level and inhibited osteoblast differentiation. These data indicated that a decreased or no changed p-ERK expression would not induce osteogenesis. Although inhibition of Cx43 or ERK signaling decreased the formation of HO (approximately 50%), it could not completely prevent the HO. Therefore, surgical

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**Figure 5.** Inhibition of Cx43 and ERK signaling decreases osteogenic marker gene expression in vivo. (A) Tenotomy mice were treated by local injection of Cx43 shRNA or intraperitoneal injection of U0126. HO tissues were decalcified, and expression of Cx43, p-ERK, ERK, Runx2 and BSP was detected by immunohistochemistry at week 8. (B) Cx43, p-ERK, Runx2 and BSP expression in the HO tissues was quantified.
Figure 6. Cx43/ERK signaling is critical for clinical outcomes in traumatic HO patients. (A) Normal bones (n = 15) and HO tissues (n = 20) were decalcified. Expression of Cx43 and p-ERK was detected by immunohistochemistry. (B) Quantification of Cx43 and p-ERK expression. (C) A typical radiographic three-dimensional reconstruction CT scan is shown for elbow HO with weak and strong expression of Cx43. (D) The volumes of the HO with weak and strong Cx43 expression were quantified. (t-test, *p < 0.05). (E) The overall recurrence-free survival of patients with weak and strong expression of Cx43 is shown. (F) The overall recurrence-free survival of patients with weak and strong expression of p-ERK is shown. (log-rank test).
reduction of the ectopic bones is necessary in some cases. Moreover, inhibition of Cx43 would result in delayed fracture healing, impaired angiogenesis and tissue regeneration. These observations suggest that application of Cx43 in HO treatment may cause disorders in other organs/tissues. Thus the clinical application of Cx43 would be limited under certain situations.

In summary, we have identified an important mechanism that may determine the progression and development of traumatic HO. We show that increased expression of Cx43 enhances the transcription of Runx2 and osteogenic differentiation in a manner that promotes HO progression. In examining the routes of Cx43-dependent activation of ERK, we have demonstrated the relevant function of Cx43/ERK signaling in HO development. We have demonstrated that pharmacotherapy with Cx43 or ERK inhibitors, such as shRNA or U0126, can markedly reduce extraskeletal bone formation in a traumatic model of HO. Using patient data and human HO specimens, we confirmed that elevated levels of Cx43 and p-ERK increase the risk of recurrence after surgical resection. The results reveal a new mechanism of HO formation and identify a potential therapeutic target.

Materials and Methods

Patients and specimens. Twenty patients who underwent surgical resection for elbow lesions between January 2010 and December 2015 were evaluated in this study, and the control bones were collected from 15 patients who received traumatic amputation (tibia, femur, radius and ulna). The patients, with a median age of 32 years (range: 21–62 years), were healthy without metabolic, inherited or other diseases that may affect the current study. No significant difference was observed in composition regarding age or sex in these two groups (p > 0.05). The mature HO lesions were removed by surgery 8–10 months after the injury. The control bones from the amputation patients were collected 2–4 hours after the injury during surgery. After surgery, all the patients were examined with x-rays or CT monthly, and the follow-up time was 25 weeks. Approval for this study was obtained from the ethics committee of Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, and written informed consent was obtained from the patients or their legal guardians. The experiment was performed in accordance with approved guidelines.

RNA isolation and gene expression profiling. Soft tissues and epiphyses were removed, epiphyses were cut off, and diaphyses were flushed with phosphate-buffered saline solution (PBS) to remove bone marrow and blood. All the samples were shock-frozen in liquid nitrogen, pulverized, and dissolved in Trizol (Life Technologies, Carlsbad, CA, USA). Total RNA was isolated from all bone or HO samples using a Trizol isolation kit according to the manufacturer’s instructions. Quantity and quality measurements were carried out using a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA) and an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, California, USA). All samples were then subjected to downstream genome-wide microarray analysis. All of the hybridization experiments were performed using Affymetrix HG-U133 Plus 2.0 GeneChips according to the manufacturer’s recommendations. The raw data were normalized using Genespring GX11 Software (Agilent Technologies, Santa Clara, CA, USA) with default parameters (MAS5 Summarization Algorithm, median of all samples as baseline transformation).

Cell culture. The preosteoblast cells MC-3T3 and C2C12 skeletal muscle cells were obtained from the Chinese Academy of Sciences (Shanghai, China). Cells were cultured in α-MEM (1000 mg/L glucose) with 10% fetal bovine serum (FBS). After confluency was reached, the culture medium was changed to an osteogenic differentiation medium (OM), which contained 10% FBS, 50 μM L-ascorbic acid, 10 mM glycerol-2-phosphate, and 100 nM dexamethasone.

shRNA transfection. MC-3T3 cells were transfectioned with lentiviral particles loaded with either small hairpin RNA (shRNA) against Cx43 or ERK (Santa Cruz Biotechnology, Santa Cruz, CA; Cat No. sc-35091–SH for Cx43, Cat No. sc-44206-SH for ERK) or a scrambled control (Santa Cruz Biotechnology, Santa Cruz, CA). The shRNA plasmids consisted of a pool of 3 to 5 lentiviral vector plasmids, each encoding target-specific 19–25 nt shRNAs designed to knockdown gene expression. The control shRNA plasmid encoded a scrambled shRNA sequence designed to not lead to the specific degradation of any known cellular mRNA. The cells were subsequently exposed to 3 μg/ml of puromycin (Sigma, USA) for 1 month. The several available clones were expanded and maintained in 1 μg/ml of puromycin to remove cells without shRNA expression.

Cell viability analysis. MT-3T3 and C2C12 cells were seeded in 96-well plates at a density of 1 × 10^4 per well and treated as follows: the culture medium was discarded, and the cells were rinsed 3 times with PBS. Subsequently, the viable cells were quantitated using a cell counting kit-8 (CCK-8, Dojindo, Japan) according to the manufacturer’s instructions. Briefly, the cells were incubated in α-MEM medium containing 10 μl of CCK-8 solution at 37 °C for 2.5 hours. Then, the optical density (OD) at 450 nm was determined using a microplate reader (BIOTEK, Vermont, USA), and the ratio of viable cells was calculated.

ALP and Alizarin Red S staining. For ALP staining, cells were stained with 5-bromo-4-chloro-3-indolyl-phosphate/nitro-blue-tetrazolium solution (Sigma-Aldrich) for 45 minutes at 37 °C to visualize ALP activity. For the Alizarin Red S staining, the cells were washed 3 times with PBS and fixed with 4% paraformaldehyde for 10 min at room temperature. After 3 washes with deionized water, the calcium mineral deposits were stained for 10 min with 2% Alizarin Red S (pH 4.2), rinsed with water, and visualized by light microscopy.

Quantitative measurement of alkaline phosphatase activity. The cells were washed twice with PBS and lysed with 1% Triton X-100 supplemented with protease and kinase inhibitors, followed by freeze–thaw cycles. ALP activity was assayed using p-nitrophenylphosphate (Sigma-Aldrich) as a substrate. Lysates were incubated with 100 μl of 3 mg/ml substrate solution for 30 min at 37 °C, and the reaction was stopped with 20 mM...
NaOH. The protein content was measured by BCA (Thermo Scientific, Rockford, IL) according to the manufacturer’s instructions. ALP activity was then expressed as Sigma unit/min/mg of protein.

Quantitative analysis of mineralization. The calcium deposits from osteoblast cells were washed 3 times with PBS and incubated for 24 h at 4 °C in 0.5 M HCl. Then, the calcium content in the HCl supernatants was measured using the Calcium Colorimetric Assay Kit (BioVision, Mountain View, CA, USA).

Western blot. The cells were washed in ice-cold PBS before lysis with a cell lysis buffer (Cell Signaling Technology, MA, USA). All samples were clarified by centrifugation at 12,000 rpm for 10 minutes at 4 °C. Then, the protein concentrations were determined using the BCA Protein Assay (Thermo Scientific, IL, USA). Equal amounts of total protein lysates were separated by SDS-PAGE, and bands were transferred to a nitrocellulose membrane. The membranes were blotted with the following primary antibodies: p-ERK (Cell Signaling Technology, Cat No. 4370, 42 kD), ERK (Cell Signaling Technology, Cat No. 9102, 42 kD), β-actin (Cell Signaling Technology, Cat No. 4970, 45 kD), Cx43 (Abcam, Cat No. ab11370, 43 kD), Runx2 (Abcam, Cat No. ab23981, 57 kD) and BSP (Abcam, Cat No. ab125227, 35 kD). Bound antibodies were detected with an Odyssey Infrared Imaging System (LI-COR Biosciences, Lincoln, NE, USA). Densitometric analysis of the protein bands was performed with Image-Pro Plus 4.5 software (Media Cybernetics, Silver Spring, MD).

RNA isolation and quantitative real-time PCR. Total RNA was prepared using Qiagen RNeasy Mini Kit (Qiagen, Valencia, CA, USA) for cellular extracts. cDNA was then generated using 1 μg of RNA using the iScript cDNA Synthesis Kit (Bio-Rad, Hercules, CA) according to the manufacturer’s instructions. Quantitative real-time expression analysis was performed using an ABI 7500 Sequencing Detection System and SYBR Premix Ex Taq (Takara, Japan). Relative expression of mRNA was determined after normalization using the ΔCt method. The following primers were used: GAPDH, 5′-ATGAGGGAAGGTGAGTGGC-3′ (forward) and 5′-GGGGTATTGATGGCAACAATA-3′ (reverse); Runx2, 5′-CCGCCCTCATGTATTAGGGC-3′ (forward) and 5′-GGTCTCTGTAATCTGAACCTGCT-3′ (reverse); ALP, 5′-TGAGGGTGTGCTTACCAG-3′ (forward) and 5′-ATGAGGCTTAGGCTTGGCT-3′ (reverse); BSP, 5′-CAGGGAGCAGTACTCTTTCC-3′ (forward) and 5′-AGTTGAAAATGTTGGGCCTT-3′ (reverse); and Col-1, 5′-GAAAAGGTGATCAGGGTGAG-3′ (forward) and 5′-GAACCACATCGATCCTGGT-3′ (reverse).

Mice. The animal experimental protocols were approved by the Animal Research Committee of Shanghai Jiao Tong University Affiliated Sixth People’s Hospital. The experiment was performed in accordance with approved guidelines. Four-week-old male BALB/c mice received an Achilles tenotomy (n = 12 per group) with sharp dissection at the midpoint in the left leg. A 1-cm incision was made on the lateral aspect of the Achilles tendon with a surgical knife. Subsequently, the Achilles tendon was exposed from its origin on the distal end of the gastrocnemius to the insertion at the calcaneus. The Achilles tendon was then divided sharply at its midpoint. The incision was then closed with absorbable sutures. Then, mice received an injection of 2 μl of Cx43 shRNA complexed with 2 μl of Lipo 2000 dissolved in 10 μl of α-MEM. Sham mice received an injection of 2 μl of scrambled shRNA plasmid complexed with 2 μl of Lipo 2000 as a negative control. All mice were treated weekly with an shRNA or scrambled shRNA local injection in the lesion in the tendon. For the in vivo p-ERK inhibition experiment, mice were treated by local injection of U0126 (Calbiochem, CA, USA; 100 μg/mouse) in the lesion in the tendon every 3 days, and DMSO was used as a negative control. The mice were maintained for 8 weeks, and the injured limbs were harvested after euthanasia.

Micro-CT and x-ray analysis. The left tibias dissected from mice were fixed with 4% paraformaldehyde for 24 h, then scanned and analyzed with a SkyScan microcomputed tomography system (Kontich, Belgium) with a 9-μm pixel size. The X-ray voltage and current were set to 80 kV and 80 μA, respectively. The region of interest (ROI) was defined to include the entire tibia, and two-dimensional (2D) image stacks were visually inspected to ensure that all heterotopic bone was included within the ROI. The Nrecon reconstruction software (NRecon v.1.4.4, SkyScan) was used to create 2-D images. For the reconstruction parameters, ring artifact correction and smoothing were fixed at zero and the beam hardening correction was set at 0%. The contrast limits were applied following the manufacturer’s instructions. The lower limit was zero and the upper limit was at the top end of the brightness spectrum. Following the reconstruction, the volume of interest (VOI) was selected within the reconstructed images to calibrate the standard unit of X-ray computed tomography density (Hounsfield unit, HU) using CTan analysis software (v.1.6.0, SkyScan). The HU values of two BMD phantom rods were measured and converted from HU to BMD values (g/cm3). The bone volume of HO was also analyzed using CTan. Analysis of x-ray was performed with a image-pro plus 4.5 software (Media Cybernetics, Silver Spring, MD). The HO area was calculate and normalized to the control group.

Histology. At 8 weeks post-tenotomy, animals were euthanized, and the tenotomized limbs were fixed in 4% paraformaldehyde. HO tissues from patients were collected in surgical operations. All of these specimens were decalcified in a 10% EDTA solution for 1 month, were embedded in paraffin and were cut into 5-μm sections for staining. Immunohistochemical staining was carried out with primary antibodies against Cx43 (Abcam, Cat No. ab11370), p-ERK (Cell Signaling Technology, Cat No. 4370), ERK (Cell Signaling Technology, Cat No. 9102), Runx2 (Abcam, Cat No. ab23981) and BSP (Abcam, Cat No. ab125227) with a 1:1,000 dilution of an appropriate secondary antibody. Protein expression was visualized with a DakoCytomation Envision staining kit. Sections were stained with 0.1% Safranin-O and 0.02% Fast Green (Sigma-Aldrich, Oakville, ON, Canada) according to the manufacturer’s instructions. The mean density of the positive area was measured by Image Pro Plus 6.0 (IPP) image analysis software. Three random slides were selected, and five random fields of images per sample were taken.
Statistical analyses. The data are represented as the mean ± standard deviation (SD). Comparisons between groups were performed using Student’s t-test, and one-way ANOVA was used for multiple comparisons. Recurrence-free survival rates were compared using Kaplan–Meier survival curves. Statistical significance was set at p < 0.05.

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Author Contributions
Bing Tu, Shen Liu and Cunyi Fan conceived and designed the experiments; Guangwang Liu contributed to the performance of experiments; Zhiwei Li and Yangbai Sun performed experiments and analyzed data.

Additional Information
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