circ_001653 Silencing Promotes the Proliferation and ECM Synthesis of NPCs in IDD by Downregulating miR-486-3p-Mediated CEMIP

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Functional changes of nucleus pulposus cells (NPCs) are considered to be the initiating factors of intervertebral disc degeneration (IDD). In this study, we investigated whether circular RNA homo sapiens (hsa)_circ_001653 (circ_001653) could bind to microRNA-486-3p (miR-486-3p) to regulate the biological properties of NPCs and the synthesis of extracellular matrix (ECM) in IDD. Initially, circ_001653 was highly expressed in isolated NPCs and degenerative NP tissues in close relation to the severity of IDD. To evaluate the effects of circ_001653 on cellular processes, we performed experiments in vitro and in vivo with altered expression of circ_001653 and miR-486-3p. An increased expression of circ_001653 in the NPCs and the degenerative NP tissues was directly associated with elevated apoptosis and an imbalance between anabolic and catabolic factors of the ECM. miR-486-3p regulated NPC proliferation and inhibited the expression of CEMIP, the cell migration-inducing hyaluronan binding protein. circ_001653 regulated miR-486-3p expression, functioning in NPCs to upregulate CEMIP, whereas circ_001653 silencing alleviated IDD in the mouse model. Altogether, circ_001653 downregulation could potentially alleviate NPC apoptosis and the metabolic imbalance of the ECM through the miR-486-3p/CEMIP axis. These mechanistic insights may present new therapeutic targets for the treatment of IDD.

INTRODUCTION

Previous work shows that intervertebral disc degeneration (IDD) is a main pathology responsible for chronic lower back pain.¹ ² Although factors such as gender, aging, obesity, and diabetes mellitus also have an association with IDD, the pathological mechanisms involved in IDD progression remain unknown.³ The intervertebral disc (IVD) is comprised of the inner nucleus pulposus (NP), which is encircled by the annulus fibrosus as well as cartilaginous endplates lying between the IVD and the adjacent vertebral bodies.⁴ The extracellular matrix (ECM) of the inner disc is composed of collagens and proteoglycans, which enable the normal functioning of the IVD.⁵ ECM in healthy NP contains a high concentration of proteoglycans within a loose network of principally type II collagen (collagen II) fibers.⁶ IDD is a complicated pathological process, which is attributed to a series of degenerative events, ultimately leading to apoptotic signaling pathways and cell death.⁷ For the present, the medical treatments of IDD include infusions of protein-based and cell-based therapies. The protein-based therapies aim to enhance ECM synthesis, thus delaying degeneration or inhibiting inflammation, whereas the cell-based therapies are intended to replace necrotic or apoptotic cells or otherwise diminish cell death.⁸

Circular RNAs (circRNAs), a type of noncoding RNA derived from exons (exonic circRNA) or introns (intronic circRNA), are distinctive RNAs with independent mechanisms of biosynthesis.⁹ Importantly, circRNAs can bind to microRNA-7 (miR-7), thus acting as a critical regulator of neurons, and participating in the pathogenesis of neurodegenerative disorders and the progression of brain tumors.¹⁰ It follows that circRNAs may potentially bind to microRNAs (miRNAs) to modulate the progression of diseases. For instance, circ_104670 is overexpressed in IDD tissues and regulates the expression of matrix metalloproteinase-2 (MMP2) by binding directly to miR-17-3p.¹¹ Additionally, circ_VMA21 protects nucleus pulposus cells (NPCs) against IDD by targeting miR-200c and X-linked inhibitor-of-apoptosis protein.¹² Another study has revealed effects of the homo sapiens (hsa)_circ_0016788/miR-486/cyclin-dependent kinase 4 (CDK4) axis in tumorigenesis of hepatocellular carcinoma (HCC), thus providing a novel therapeutic target for HCC.¹³ There was a negative correlation between the levels of ECM1 and miR-486-3p in cervical cancer tissues.¹² Interestingly, miR-486 has low expression in IDD specimens,¹⁴ which supports our hypothesis that miR-486 downregulation may be involved in the progression of IDD. In the present context of IDD, it seems relevant that cell migration-inducing hyaluronan binding protein (CEMIP), also known as KIAA1199, is closely associated with cellular invasion, proliferation, as well as motility.¹⁵ Given its higher expression in the progeria syndrome relative to healthy persons and in mortal human cells relative to immortalized cells, CEMIP may help to explain the close association between cellular and...
organismal aging. In the present study, we examined the function of circ_001653 and miR-486-3p in IDD NPCs. With the use of microarray analysis, we identified 7294 circRNAs that were differentially expressed in degenerated human IDD NPCs. Since circRNAs regulate the viability, degradation, apoptosis, and oxidative stress in NPCs, they may be positioned to present a molecular target for gene therapy of IDD. Our in silico analysis led us to predict that circ_001653 has a potential binding site of miR-486-3p, which substantiates the correlation between circ_001653 and miR-486-3p in NPCs. Further, we provide new evidence for the involvement of miR-486-3p and CEMIP in IDD. Taken together, we provided a novel experimental and theoretical basis for the rationally targeted therapy of IDD.

RESULTS
Identification of Differentially Expressed circRNAs
The differentially expressed circRNAs were screened from the IDD-related microarray chip GSE67566 using the limma package in R language with |logFC| >1 and p <0.05 set as the threshold for significance. As the annotation platform GPL19978 showed, there were ten samples, including five IDD samples and five healthy controls, in the Agilent-069978 Arraystar Human CircRNA Microarray V1, among which, there were 339 upregulated circRNAs and 289 downregulated circRNAs. The circRNA circ_001653 had especially high expression in GSE67566 (Figure 1A). In addition, some studies have shown that circ_001653 plays an important role in the progression of IDD, but its detailed mechanism in IDD is still unclear. As shown in Figure 1B, a volcano plot was drawn by analyzing the chip with the R language protocol. The red dots represent the significantly upregulated circRNAs, and the green dots show downregulated ones, whereas the black dots indicate circRNAs without significantly different expression in NP tissues of IDD cases.

Enhanced Expression of circ_001653 in NP Tissues of IDD Is Correlated with the Degree of Degeneration
We verified the isolated NPCs: the primary cells were plated into culture bottles and initially formed spherules in the culture medium. The cells began to adhere to the vessel wall on the 3rd to 5th day and then gradually proliferated, extended out pseudopodia, and became spindle or polygonal shaped. The cell confluence reached about 90% after 20–30 days of culture. When the cell confluence rate was sufficiently high enough, the cell masses were confluent in the shape of stone or whirlpool with rich cytoplasm, strong refraction, oval nucleus, and clear structure. After passage, the adherence time of the cells was significantly shorter than that of the primary generation, such that most cells became adherent to the wall within 12 h (Figures 2A–2C). We tested for expression of collagen type II alpha 1 (COL2A1) and aggrecan using double immunofluorescence, which showed as green and red staining in the NPC cytoplasm, which was absent in the nucleus (Figures 2D and 2E). The positive rate of CD24 was 71.43% ± 6.48% (Figure 2F). Taken together, these results confirmed that the isolated cells were NPCs.

We measured and compared the transcription levels of hsa_circ_001653 in IDD tissues and control tissues by quantitative reverse transcriptase polymerase chain reaction (qRT-PCR), finding a significantly higher expression of circ_001653 in the degenerative NP tissues compared with spinal cord injury (SCI) tissues (Figure 2G). In addition, Pearson’s correlation analysis showed that the expression
of circ_001653 was positively correlated with the clinical severity of IDD ($R^2 = 0.4036$, $p = 0.0003$, $n = 28$; Figure 2H).

circ_001653 Is Involved in Proliferation, Apoptosis, and ECM Synthesis of NPCs

We investigated the effects of circ_001653 on the regulation of biological functions of the NPCs by circ_001653 silencing. The relative expressions of circ_001653 and linear 001653 in each group of cells were determined to assess the specificity of small interfering RNA (siRNA) circ_001653 (si-circ_001653) targeting the circular structure of circ_001653 (Figure 3A). This analysis revealed that circ_001653 expression was downregulated by the siRNA, whereas expression of the linear 001653 was unaffected relative to the treatment of si-negative control (NC), thus indicating success of the si-circ_001653 interference. In addition, we evaluated the cell proliferation of NPCs by the cell counting kit-8 (CCK-8) assay, which revealed higher proliferation of NPCs treated with si-circ_001653 than that in the si-NC group (Figure 3B). Likewise, downregulation of circ_001653 in the NPCs decreased apoptosis and the number of cells in the G1 phase (Figures 3C–3H). We investigated the role of circ_001653 in the synthesis of ECM in NPCs by determining the mRNA and protein expressions of apoptosis and ECM synthesis-related factors (Figures 3I–3L). This analysis revealed that treatment of cells with si-circ_001653 led to a dramatic enhancement in the expression levels of miR-486-3p, collagen II, and aggrecan, whereas the expression levels of caspase-3, MMP3, MMP13, a disintegrin and metalloproteinase with thrombospondin motifs 4 (ADAMTS4), and ADAMTS5 were markedly reduced. These results indicated that circ_001653 silencing could promote the synthesis of ECM and proliferation in NPCs of IDD.

circ_001653 Competitively Binds to miR-486-3p

We predicted the miRNAs binding to circ_001653, as well as the secondary structure of circ_001653 (Figure 4A). The yellow-marked sequence in Figure 4B was the sequence of circ_001653 expected to bind to miR-486-3p. We measured the abundance of circ_001653 and linear 001653 in NPCs, with or without treatment of the RNase R enzyme, which showed that circ_001653 had strong resistance to RNase R treatment and could be enriched, whereas almost all linear RNA was digested (Figure 4C). Additionally, six miRNAs were obtained through the intersection of the top 500 miRNAs shown in the microarray GSE116726 analysis (with log transformation) and the predicted miRNAs (Figure 4D). The relative expression of differentially expressed miRNAs in NPCs in IDD was determined by qRT-PCR, which showed that miR-486-3p expression was significantly downregulated in NPCs in IDD (Figure 4E).

Furthermore, the binding sites between circ_001653 and miR-486-3p were identified in keeping with predictions (Figure 4F). To determine whether circ_001653 could indeed bind to miR-486-3p, we identified the localizations of circ_001653 and miR-486-3p in cells by the fluorescent in situ hybridization (FISH) assay (Figure 4G). The FISH assay showed that circ_001653 and miR-486-3p were mainly
circ_001653 Impairs Proliferation and ECM Synthesis of NPCs while Promoting Their Apoptosis by Binding to mir-486-3p

The role of miR-486-3p in the proliferation of NPCs in IDD was measured by the CCK-8 assay, which showed that proliferation of miR-486-3p mimic-treated NPCs was significantly faster than that of NC mimic-treated cells, whereas proliferation was significantly decreased after addition of overexpression (OE)-circ_001653; the effect of combined treatment with OE-circ_001653 and miR-486-3p mimic was similar to that of NC mimic (Figure 5A). Therefore, overexpression of circ_001653 could rescue the increase of cell proliferation induced by miR-486-3p mimic. Meanwhile, the effects of

expressed in the cytoplasm; here, green staining represents circ_001653, red staining the miR-486-3p, and blue staining shows the nuclei. Upon merging the three different fluorescences, we see that circ_001653 and miR-486-3p were mainly localized in cytoplasm. Furthermore, the binding relationship between circ_001653 and miR-486-3p was confirmed by the Dual Luciferase Reporter Gene Assay. The luciferase activity of cells transfected with pmirGLO-circ_001653 (pcirc)_001653-wild type (WT) was significantly repressed by miR-486-3p mimic relative to NC mimic, but luciferase activity for transfection with pcirc_001653 mutant (mut) was unchanged in both groups (Figure 4H), thus confirming that circ_001653 could be targeted and degraded by miR-486-3p. In addition, we performed enrichment of circ_001653 by the Argonaute2 (AGO2) immunoprecipitation assay, which showed that circ_001653 pulled down with AGO2 was enriched in NPCs transfected with miR-486-3p (Figure 4I), thus indicating that miR-486-3p could directly target circ_001653 in an AGO2-dependent manner. Next, we detected the linking between circ_001653 and miR-486-3p by northern blot analysis (Figure 4I) and measured the expression of miR-486-3p in NP tissues in IDD and SCI tissues by qRT-PCR (Figure 4K). These results showed that the expression of miR-486-3p in degenerative NP was significantly lower than that in healthy SCI tissues. The correlation analysis showed an inverse relationship between the expressions of circ_001653 and miR-486-3p (Figure 4L). The aforementioned results suggested that circ_001653 could bind directly to miR-486-3p in NPCs.

**Figure 3. circ_001653 Regulates the Biological Functions of NPCs in IDD**

The relative expressions of circ_001653 and linear 0016553 in cells were determined by qRT-PCR after treatment with siRNA (A). Proliferation of NPCs examined by the CCK-8 assay (B). NPC apoptosis detected by fluorescence staining. Scale bars, 25 μm (C). Quantification of NPC apoptosis evaluated by TUNEL assay (D). The cell cycle distribution in each group by flow cytometry (E and F). Representative dot plots of apoptosis detection after annexin V-FITC/PI double staining (G and H). The expression levels of miR-486-3p in cells upon treatment with si-circ_001653 were determined by qRT-PCR (I). The mRNA expression levels of collagen II, aggrecan, caspase-3, MMP3, MMP13, ADAMTS4, and ADAMTS5 in response to the treatment with si-circ_001653, as determined by qRT-PCR (J). The protein expression levels of collagen II, aggrecan, caspase-3, MMP3, MMP13, ADAMTS4, and ADAMTS5 in response to the treatment with si-circ_001653, as measured by western blot analysis (K and L). *p < 0.05 versus the NPCs treated with si-NC. Statistical data were measurement data described as mean ± standard deviation. The experiment was repeated three times independently. Data between two groups were analyzed by independent sample t test and at different time points by repeated-measurement analysis of variance.
Figure 4. circ_001653 Regulates miR-486-3p in NPCs

Prediction of miRNAs with binding sites to circ_001653 through the bioinformatics programs (A). Sequence fragments of circ_001653 binding to miR-486-3p (B). The abundance of circ_001653 and linear 001653, with or without treatment of RNase R enzyme (C). Results of the top 500 differentially expressed miRNAs in microarray GSE116726 and miRNAs predicted by bioinformatics. Intersecting miRNAs were hsa-miR-18b-5p, hsa-miR-4651, hsa-miR-150-3p, hsa-miR-4688, hsa-miR-486-3p, and hsa-miR-221-3p (D). Relative expression of miRNAs detected by qRT-PCR (E). The binding sites of circ_001653 to miR-486-3p analyzed by the bioinformatics programs (F). circ_001653 and miR-486-3p were located in the cytoplasm of NPCs, as detected by RNA FISH; scale bars, 25 μm (G). Identification of interaction between circ_001653 and miR-486-3p by the Dual Luciferase Reporter Gene Assay (H). AGO2 immunoprecipitation in degenerative NPCs transfected with miR-486-3p mimic or NC and the quantification of miR-486-3p expression determined by qRT-PCR (I). Northern blot analysis of miR-486-3p after being pulled down by a circ_001653 probe or a random probe (J). The relative expression of miR-486-3p in NP tissues of IDD and SCI tissues evaluated by qRT-PCR (K). The correlation between circ_001653 and miR-486-3p assessed by Pearson’s correlation analysis (L). *p < 0.05 versus NC mimic or SCI tissues. Statistical data in Figures 5J and 5K were measurement data described as mean ± standard deviation; n = 30 for the SCI tissues, and n = 36 for the NP tissues from the IDD patients. The experiment was repeated three times independently. Data between two groups were analyzed by t test. The one-way ANOVA was adopted for comparison among multiple groups, followed by Tukey’s post hoc test.
Figure 5. circ_001653 Inhibits Proliferation and ECM Synthesis of NPCs while Promoting their Apoptosis by Binding to miR-486-3p

NPCs were cotransfected with miR-486-3p, with or without OE-circ_001653. The proliferation of NPCs was examined by CCK-8 (A). The apoptosis measured by the fluorescence staining; scale bars, 25 μm (B). Quantification of NPC apoptosis evaluated by TUNEL staining (C). The cell cycle distribution in each group recorded by flow cytometry (D). Histogram of cell cycle distribution in each group by flow cytometry (E). The representative dot plots of apoptosis detection by flow cytometry (F). The apoptotic rate of each group detected by flow cytometry (G). The mRNA expression levels of collagen II, aggrecan, caspase-3, MMP3, MMP13, ADAMTS4, and ADAMTS5 determined by qRT-PCR (H). The protein bands of collagen II, aggrecan, caspase-3, MMP3, MMP13, ADAMTS4, and ADAMTS5 measured by western blot analysis (I). The protein levels of collagen II, aggrecan, caspase-3, MMP3, MMP13, ADAMTS4, and ADAMTS5 measured by western blot analysis (J). *p < 0.05 versus the NPCs treated with NC mimic; #p < 0.05 versus the NPCs treated with miR-486-3p mimic. Statistical data were measurement data described as mean ± standard deviation. The experiment was repeated three times independently. Data among multiple groups were analyzed by one-way analysis of variance and at different time points by repeated-measurement analysis of variance, followed by Tukey’s post hoc test.
miR-486-3p on cell apoptosis and cell cycle progression of NPCs were determined by terminal deoxynucleotidyl transferase (TdT)-mediated 2’-deoxyuridine 5’-triphosphate (dUTP) biotin nick end-labeling (TUNEL) staining and flow cytometry. In comparison with the NC group, the treatment with miR-486-3p mimic significantly enhanced the number of G2-phase cells and significantly decreased cell apoptosis, whereas the application of OE-circ_001653 reversed effects induced by miR-486-3p mimic (Figures 5B–5G). Furthermore, we determined the levels of the apoptosis and ECM synthesis proteins collagen II, aggrecan, caspase-3, MMP3, MMP13, ADAMTS4, and ADAMTS5. When transfected with miR-486-3p mimic, the cells displayed promoted expression levels of collagen II and aggrecan, but significantly decreased levels of caspase-3, MMP3, MMP13, ADAMTS4, and ADAMTS5. The enforced expression of circ_001653 elevated the apoptotic and catabolic effects of these cytokines and restrained the expression of collagen II and aggrecan (Figures 5H–5J). Collectively, these data indicated that circ_001653 might inhibit the proliferation and ECM synthesis of NPCs by binding to miR-486-3p.

miR-486-3p Upregulates Proliferation and ECM Synthesis of NPCs in IDD through Targeting CEMIP

The positive correlation between circ_001653 and CEMIP and the negative correlation between miR-486-3p and CEMIP were identified by Pearson’s correlation analysis (Figures 6A and 6B), and the binding sites between miR-486-3p and CEMIP were predicted (Figure 6C). The targeting relationship was further verified by the Dual Luciferase Reporter Gene Assay. This analysis showed that the luciferase activity of the CEMIP-WT reporter was suppressed by miR-486-3p, whereas the introduction of targeted mutations abolished the inhibitory effect of miR-486-3p (Figure 6D). We next measured the expression levels of CEMIP in the SCI tissues and the degenerative NP tissues, finding a decline in the expression of CEMIP in the degenerative NP tissues compared with the expression in SCI tissues (Figure 6E).

We assessed the effect of CEMIP on the proliferation of NPCs in IDD by the CCK-8 assay (Figure 6F), which showed no significant differences between groups in the optical density (OD) value on the 1st day after transfection. Furthermore, the NPCs treated with overexpressed CEMIP displayed a significantly inhibited OD value on the 2nd, 3rd, and 4th days after transfection versus those treated with OE-NC. In comparison with the cells treated with OE-CEMIP alone, the OD value of cells with combined treatment of si-circ_001653 or miR-486-3p mimic with OE-CEMIP increased significantly on the 2nd, 3rd, and 4th days after transfection. These results indicate that transfection of OE-CEMIP could reverse the increase of cell proliferation induced by si-circ_001653, whereas overexpression of CEMIP inhibited the proliferation induced by miR-486-3p mimic. Furthermore, we investigated the effects of miR-486-3p and CEMIP on the cell apoptosis and cell cycle of NPCs as well as ECM synthesis (Figures 6G–6O), finding that CEMIP upregulation induced apoptosis of NPCs and reduced the expression levels of ECM synthesis-related cytokines. Furthermore, the presence of CEMIP markedly counteracted the effects of miR-486-3p mimic and circ_001653 silencing apoptosis, cell cycle, and ECM synthesis of NPCs, indicating that miR-486-3p and circ_001653 exerted their functions in NPCs through CEMIP.

circ_001653 Silencing Alleviates IDD in the Mouse Model

The IDD mouse model was established to confirm the effect of circ_001653 in vivo. The modeled mice were treated with si-circ_001653 or OE-circ_001653 + miR-486-3p agomir. Computed tomography (CT) scanning showed persistent narrowing of the disc space at the 1st, 5th, and 9th weeks in all IVD-punctured mice. The value of disc height index (DHI) of all IVD-injected mice was significantly decreased at the 1st week after operation relative to baseline values (p < 0.01). The DHI of the IVD-injected mice at each time point after injection was significantly lower than that of the mice that received no in vitro fertilization (IVF) puncture (p < 0.01; Figure 7A). At the 9th week after operation, we performed a magnetic resonance imaging (MRI) examination, which showed significantly lower degeneration scores of the IVDs in the si-circ_001653-injected mice than that in the noninjected control mice (p < 0.05; Figure 7B). With the undertaking of Safranin O staining, the mucopolysaccharides, chondrocytes, and granules of mast cells presented as orange-red. Examination of the IVD showed a clear framework with red-stained NP, fibrous rings, and cartilage plates, which indicated abundant proteoglycan in the control group. In contrast, we saw atrophy of NP, narrowed IVD with unclear framework, less red-stained tissue, and smaller amounts of proteoglycan in the noninjection group. As shown in the model mice treated with si-circ_001653 and...
Figure 7. circ_001653 Silencing Alleviates IDD in the Mouse Model

A total of 24 mice were randomly divided into four groups: nonpuncture group (control), noninjection with puncture group (noninjection), si-circ_001653 injection with puncture group (si-circ_001653), and OE-circ_001653 plus miR-486-3p agomir injection with puncture group (OE-circ_001653 + miR-486-3p agomir). Changes of intervertebral DHI after injection (A). MRI scores of mice after 9 weeks of injection (B). Safranin O staining in IVD of mice (400×) (C). Apoptosis conditions in IDD NPCs of mice (legend continued on next page)
OE-circ_001653 + miR-486-3p agomir, the NP of IVD showed an ordered and clear framework. Besides, there were densely distributed island chondrate cells and fibrous rings with clear boundaries, along with dark-red staining with moderate hyperchome (Figure 7C). TUNEL staining and immunohistochemistry showed that circ_001653 silencing could attenuate degenerative changes of NPCs, such as increased cell apoptosis and catabolic responses of the ECM and decreased ECM composition in the mouse model of IDD (Figures 7D–7F). Taken together, these results revealed positive or ameliorative effects in vivo of decreasing circ_001653 expression in attenuating NPC apoptosis, inhibiting ECM catabolism, and promoting anabolism in the NP.

DISCUSSION

The NP of the intervertebral disc, which provides compressive load support in the spine, contains cells that are critical in the generation and maintenance of this tissue. The role of circRNAs in IDD has been previously emphasized, including their effects in the modulation of NPC death and ECM synthesis. In this study, we used bioinformatics to identify the top 20 differentially expressed circRNAs in IDD. Among these, we predicted a critical role for circ_001653 in IDD. Collectively, the results of this present study demonstrate that circ_001653 indeed regulates the expression of CEMIP by binding to miR-486-3p, which mediates the proliferation, apoptosis, and ECM synthesis of NPCs in IDD.

miRNAs have an emerging potential for the prevention and treatment of IDD, specifically for modulating IDD-related cell processes, such as NPC proliferation and apoptosis. For example, restored transcription of miR-486 exerts a profoundly positive effect in alleviating progression in dystrophic skeletal muscle. In metastatic colorectal cancer, CEMIP (KIAA1199) is highly expressed in CD44+ cancer cells relative to paired primary tissues and is positively linked to the mesenchymal phenotype and numbers of circulating tumor cells, while also predicting a briefer, progress-free survival time. Furthermore, a finding of enhanced expression of CEMIP may have prognostic significance in patients with pancreatic ductal adenocarcinoma and is correlated with increased pancreatic ductal adenocarcinoma cell migration. In the present study, we confirmed that miR-486-3p has binding sites with circ_001653 and CEMIP, respectively. Similarly, circ_001678 has been implicated in hepatocellular carcinoma tumorigenesis through binding to miR-486 to regulate CDK4.

The present study showed that circ_001653 downregulation significantly decreased apoptosis and promoted proliferation of NPCs and ECM synthesis by regulating the miR-486-3p/CEMIP axis. Consistent with our study, overexpressed miR-10b induced NPC proliferation, which may be a possible mechanism in the progression of IDD. In conclusion, notochordal cell conditioned medium can suppress the apoptosis of bovine NPCs and their degradative effects via inhibition of induced caspase-3/7 and a matrix-degrading gene (MMP3), while promoting the expression of anabolic/matrix-protective genes (aggrecan and collagen II). Moreover, polymorphisms in aggrecan and MMP3 appear to be well positioned for a connection with disc degeneration (DD) since an association of these genes has been confirmed in various ethnic populations in relation to the individual’s vulnerability to DD. Furthermore, IDD-related catabolic cytokines were remarkably enhanced in severe IDD, along with diminished ECM and increased expression of inflammatory cytokines. Along with the general degradation of ECM that occurs during DD, the pattern of synthesis and the conformation of collagens also change in the diseased disc. Zou et al. illustrated that circRNAs may modulate the expression of genes that encode IDD-related nucleic acid binding proteins, enzyme regulators, as well as transferases, revealing the involvement of circRNAs in the ECM metabolism. Specifically, chondrocyte ECM-related circRNA (circRNA-CER) silencing exerts an inhibitory effect on MMP13 expression and a promotive effect on ECM synthesis. ADAMTS1 and ADAMTS5 are observed to be overexpressed in murine aortas, and reduced ADAMTS5 may lead to the accumulation of aggrecan. The controlled synthesis and degradation of hyaluronic acid are important for maintaining the homeostasis of ECM. Liang et al. proved that CEMIP-induced metabolism of hyaluronic acid may be responsible for carcinogenesis through the activated neovascularization and invasion in tumor tissues. In vitro, CEMIP has been validated to promote proliferation and invasive capacity, as well as epithelial-to-mesenchymal transition of cancer cells. Besides, the promotive role of miR-486 in proliferation and suppressive role in apoptosis have been emphasized in myeloid cells as well. Here, we have demonstrated that miR-486-3p stimulated by circ_001653 silencing suppressed the expression of CEMIP, thereby decreasing apoptosis and increasing proliferation and ECM compositions of NPCs in vitro.

In conclusion, the present study demonstrates that circ_001653 downregulation facilitated NPC proliferation and ECM synthesis, while reducing apoptosis of NPCs by downregulating CEMIP obtained through upregulation of miR-486-3p (Figure 8). These findings are invaluable for obtaining a better understanding of the molecular mechanisms underlying the pathophysiology and molecular pathway of IDD and have important clinical significance in the prevention and treatment of degenerative discogenic diseases. However, in our review of literature, few studies have hitherto illustrated the role of circ_001653 in IDD or other related degenerative conditions, and the relation between miR-486 and miR-486-3p remains to be investigated further. In addition, it is possible that circ_001653 may function in NPCs through other miRNAs or other unknown mechanisms, in addition to miR-486-3p. Thus, further studies are essential to validate these findings and expand the translational potential of this direction.

assessed by TUNEL staining (400×) (D). Apoptosis ability detection (E). The expression of ECM synthesis-associated proteins (100×) in mice IVD cells detected by immunohistochemistry (F). *p < 0.05 versus the control group; #p < 0.05 versus the noninjection group; &p < 0.05 versus the si-circ_001653 group. Statistical data were measured as mean ± standard deviation. The experiment was repeated three times independently. Data among multiple groups were analyzed by one-way analysis of variance and at different time points by repeated-measurement analysis of variance, followed by Tukey’s post hoc test.
MATERIALS AND METHODS

Ethics Statement
This study was approved and supervised by the Ethics Committee of Shengjing Hospital of China Medical University. Great efforts were made to minimize the number of animals used in the experiments and their suffering. Written, informed consents were acquired from all patients enrolled for the use of their tissues for research purposes.

Study Subjects
We collected 36 cases of NP tissues of IDD patients from April 2015 to May 2017 at Shengjing Hospital of China Medical University, among which, there were 22 men and 14 women aged 21–74 with a mean age of 53 years (body mass index: 19.0–26.0). Based on the Pfirrmann grading system, IDD patients were assigned to grades I–III. Enrolled patients were free of lumbar, leg, or spinal canal surgery history before IDD, and all patients had suffered chronic disease with a course of 2.5–12 years (average course of IDD was 11 years). Patients with cervical spinal injury, tumor, inflammation, and tuberculosis were excluded from the study. An additional 30 cases of IVD tissues without apparent degeneration were obtained as control material from patients (21 men and 9 women aged 19–47 years) who were free of cervical or shoulder trauma. The tissues were instantly frozen by immersion in liquid nitrogen and then stored at −80°C.

Isolation and Culture of NPCs
The NP tissue was removed under aseptic conditions, immediately immersed in D-Hanks solution, then separated by ophthalmic shears on an ultra-clean workbench at the junction of the annulus fibrosus, and cut into 1 mm³ cubes. The tissue blocks were treated by 0.25% type II collagenase for 1–2 h at 37°C and washed twice with phosphate-buffered saline (PBS), whereupon the tissue residue was removed by passing through a 200-mesh filter. The detachment was terminated by resuspension in Dulbecco’s modified Eagle’s medium/Ham’s F-12 medium (DMEM/F12) culture medium, supplemented with 20% fetal bovine serum (FBS). After counting, the cells were seeded into 25 cm² culture dishes and incubated at 37°C. The primary culture was started in a 5% CO₂ incubator, and the medium was changed every 3 days. When the confluence of cells attained 90%, the cells were detached with 0.25% trypsin at 37°C for 2 min, which was then terminated by the culture medium. Cells were then prepared into cell suspension and passaged at a ratio of 1:2. The cells at the primary, second, and third passages were observed and photographed under an inverted phase-contrast microscopy. The cells at passage 3 were used for subsequent experiments.

Immunofluorescence Staining
Primary monolayer NPCs were collected and cultured on the slide conventionally. When cell confluence reached 50%–60%, the culture medium was removed, after which, the cells were rinsed using PBS for three times, fixed with 4% paraformaldehyde for 30 min at room temperature, and put in 0.2% Triton X-100 at 37°C to break the membrane for 30 min. Next, the cells were blocked using goat serum at 37°C for 30 min and incubated overnight at 4°C with an appropriate concentration of mouse monoclonal antibodies against COL2A1 (ab150771, 1:200; Abcam, Cambridge, MA, USA) and aggrecan (ab3773, 1:200; Abcam, Cambridge, MA, USA). The next day, the cells were balanced for 30 min and incubated overnight at 4°C with an appropriate concentration of mouse monoclonal antibodies against COL2A1 (ab150771, 1:200; Abcam, Cambridge, MA, USA) and aggrecan (ab3773, 1:200; Abcam, Cambridge, MA, USA). The next day, the cells were rinsed with PBS for 3 times (5 min/time), and rebrodf with goat anti-mouse fluorescein isothiocyanate (FITC) (a0568, 1:500; Beyotime Biotechnology, Shanghai, China) and fluorescent secondary goat anti-mouse Cy3 (A0521, 1:500; Beyotime Biotechnology, Shanghai, China), respectively, at 37°C in the dark for 30 min. After PBS rinse for 3 times (5 min/time), the cells were stained with 4’,6-diamidino-2-phenylindole (DAPI) (nucleus), blocked with fluorescence decay-resistant medium, and observed under a fluorescence microscope (Leica, Germany).

Cell Transfection and Treatment
The NPCs at a logarithmic growth phase were used for transfection. The siRNA targeting circ_001653 (si-circ_001653), circ_001653 OE
plasmid (circ_001653), miR-486-3p mimic, CEMIP overexpression plasmid (OE-CEMIP), or their controls (si-NC, NC mimic, and OE-NC) were obtained from Dharmacon (Lafayette, CO, USA).

RNA Isolation and Quantification
The total RNA in cells 24 h postculture was extracted using the Trizol reagent (Shanghai Haling Biological Technology, Shanghai, China). The concentration, purity, and integrity of RNA were determined using NanoDrop ND-1000 spectrophotometry and 1% agarose gel electrophoresis. The complementary DNA (cDNA) was synthesized according to the instructions of the EasyScript First-Strand cDNA Synthesis SuperMix (AE301-02; Transgene Biotech, Beijing, China) with a 20-μL reaction volume. The fluorescence quantitative real-time PCR was carried out with the reference to the instructions of the SYBR Premix Ex Taq II kit (TaKaRa, Dalian, Liaoning, Japan) on the ABI 7500 quantitative real-time PCR instrument (ABI, Oyster Bay, NY, USA). The primers used are listed in Table 1, and the fold changes were calculated by means of relative quantification (2^ΔΔCt method).

Western Blot Analysis
The degenerative NPCs at logarithmic growth phase were lysed with the prepared radioimmunoprecipitation assay (RIPA) lysis buffer on ice for 30 min. Afterward, the protein concentration was determined by the bichinchoninic acid (BCA) kit, and the proteins were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). Next, the proteins were transferred onto the nitrocellulose membrane by the wet transfer method and blocked with 5% bovine serum albumin (BSA) at room temperature for 1 h. The membrane was then incubated with diluted primary rabbit antibodies against CEMIP (ab107073, 1:500), cleaved caspase-3 (ab32042, 1:500), collagen II (ab34712, 1:1,000), MMP3 (ab53015, 1:1,000), MMP13 (ab51072, 1:500), ADAMTS4 (ab185722, 1:1,000), and ADAMTS5 (ab41037, 1:250) and mouse anti-aggrecan antibody (ab3778, 1:100) at 4°C overnight. All of these antibodies were purchased from Abcam (Cambridge, MA, USA). The membrane was then incubated with the secondary antibody, horseradish peroxidase (HRP)-conjugated goat anti-rabbit immunoglobulin G (IgG) (1:5,000; ZSGB-Bio, Beijing, China), at room temperature or at 4°C for 1 h. At room temperature, the membrane was reacted with enhanced chemiluminescence (ECL) solution (ECL808-25; Biostica, San Diego, CA, USA) for 1 min, then compacted with the dedicated software (National Institutes of Health, Bethesda, MD, USA). The ratio of the gray value of the target protein to that of the internal control, β-actin, was taken as the relative expression of the protein.

CCK-8 Assay
Cell proliferation was measured by the CCK-8 assay. The transfected cells were seeded into a 96-well culture plate at a density of 1,500 cells/well. When the cell confluence reached approximately 80%, the cells were transfected according to the instructions of the Lipofectamine 2000 reagent (Invitrogen, Carlsbad, CA, USA). The culture medium was changed 6 h after transfection, and the cells were incubated at 37°C, 5% CO2 for 2 h. The CCK-8 assay was performed according to the manufacturer’s instructions, with 5 μL of CCK-8 solution (Dojindo Molecular Technologies, Kumamoto, Japan) added at baseline (0 h) and at 24 h, 48 h, 72 h, and 96 h, and incubated at 37°C with 5% CO2 for 2 h. The OD value of each well was measured using a microplate reader (Tecan, Mannedorf, Switzerland) at the wavelength of 450 nm.

Flow Cytometry
The third passage of NPCs in good growth conditions was detached with trypsin. After the adjustment of the concentration into 1 × 10^6 cells/mL, a 1-μL portion of cells was washed with PBS and mixed with 100 μL mouse anti-human CD24-phycocerythrin (PE) monoclonal antibody, and the isotype control was set. The cells were incubated at room temperature in darkness for 30 min. After being washed with 2 mL of PBS, the cells were counted using the flow cytometer. The positive rate of CD24 in NPCs was recorded according to the fluorescence intensity, with the isotype control serving as the negative cell group.

### Table 1. The Primer Sequences for qRT-PCR

| Gene          | Primer Sequences (5’–3’)                          |
|---------------|---------------------------------------------------|
| circ_001653 F | AGCCAGGAGAGACCTTGACT                               |
| circ_001653 R | GAGGCTTTACAGTGTCCTCC                               |
| miR-486-3p F  | CGGCGCCGATCGTACAGGAT                               |
| miR-486-3p R  | CATGCTTGAGTAGTAGGACGAT                             |
| CEMIP F       | GCTCTTGAGTTGACATGGAC                               |
| CEMIP R       | ACCGCGTTCAATATTGAC                                 |
| Collagen II F | AGAATCTGTTGAGCACGCAAGAGA                           |
| Collagen II R | AGCACGCGTAGGAGGGTCAT                                |
| Aggrecan F    | TGGAGTACAGGGAGCTTGGG                               |
| Aggrecan R    | TGGAGTACAGGGAGCTTGGG                               |
| MMP3 F        | TTTCTTGGATTGGAGGTTGAC                              |
| MMP3 R        | AGCCCTGGAGAATGTTGAGTGG                             |
| MMP13 F       | CCGAACCAACACTCCAAC                                 |
| MMP13 R       | AAACAGCTCCGGCATCAACC                               |
| ADAMTS4 F     | ACCCACACTCCAGATC                                   |
| ADAMTS4 R     | TGGCCACCATCGCATAC                                   |
| ADAMTS5 F     | GACAGTTCCAAGCAGCAAGCACC                            |
| ADAMTS5 R     | TTTCTTGGAGGAT                                      |
| β-Actin F     | AGGGACATCCCTCCAAAGATT                              |
| β-Actin R     | GGGCCAGAAGGTCATCATT                                 |

qRT-PCR, quantitative reverse transcriptase polymerase chain reaction; F, forward; R, reverse; circ_001653, circular RNA has_circ_001653; miR-486-3p, microRNA-486-3p; CEMIP, cell migration-inducing hyaluronan binding protein; MMP, matrix metalloproteinase; ADAMTS, a disintegrin and metalloproteinase with thrombospondin motifs.
A total of $1 \times 10^6$ cells in the logarithmic growth phase were washed two times with PBS, fixed with precooled 70% alcohol, and stained with 1 mL of 50 μg/mL propidium iodide (PI) for 30 min in darkness. The cell cycle was detected using the flow cytometer (FACSCalibur, Becton Dickinson, Franklin Lakes, NJ, USA) and analyzed by the ModFit software.

A total of $1 \times 10^6$ cells in the logarithmic growth phase were washed two times with precooled PBS, resuspended with 1 × annexin buffer, and allowed to stand with 5 μL of annexin V-FITC (Becton Dickinson, Franklin Lakes, NJ, USA) at room temperature for 10 min. After being washed with PBS, the cells were resuspended with 300 μL of 1 × annexin buffer, and the cell apoptosis rate was determined using the flow cytometer.

**TUNEL Staining**

We adopted TUNEL staining to detect the apoptotic DNA fragments. The degenerative NPCs were incubated with 3% H2O2 and 0.1% Triton X-100 for 10 min and washed three times with PBS. Subsequently, the TUNEL cell staining was conducted based on the manufacturer’s instructions in the in situ cell death detection kit (F. Hoffmann-La Roche, Basel, Switzerland) by fixing with 4% paraformaldehyde and staining with DAPI. Cell apoptosis was observed under the optical microscope (BX61; Olympus, Tokyo, Japan).

**FISH**

FISH was utilized to identify the localization of circ_001653 in cells, according to the instructions of the Ribo IncRNA FISH Probe Mix (Red) (Guangzhou RiboBio, Guangzhou, Guangdong, China). The cells in logarithmic growth phase were seeded into the six-well culture plates at a density of $3 \times 10^4$ cells per well and cultured for 2 days until the confluence reached about 80%. The cells were fixed at room temperature with 1 mL of 4% paraformaldehyde for 10 min; treated with Proteinase K (2 μg/mL), glycine, and ethylphthalein; and incubated with the 250 μL of prehybridization solution at 42°C for 1 h. The cells were then hybridized with 250 μL of hybridization solution containing probe (300 ng/mL) at 42°C overnight. The next days, the cells were seeded into 24-well plates and stained for 5 min with DAPI prepared with PBS containing Tween 20 (PBST) and sealed by addition of an anti-fluorescence quenching agent. Five different visual fields were randomly selected and photographed under the fluorescence microscope (Olympus, Tokyo, Japan).

**Dual Luciferase Reporter Gene Assay**

The binding sites between circ_001653 and miR-486-3p and between miR-486-3p and CEMIP were analyzed using the biological prediction websites (https://circinteractome.nia.nih.gov/). The full length of circ_001653 and the 3’ untranslated region (UTR) of CEMIP were cloned into the luciferase vector of pmirGLO (E1330; Promega, Madison, WI, USA), which we named pcirc_001653-WT and pCEMIP-WT. The site-directed mutations were conducted in the regions between miR-486-3p and circ_001653 and between miR-486-3p and CEMIP, and the pcirc_001653-mut and pCEMIP-mut vectors were constructed. The pRL-TK was applied to adjust for differences in the number of cells and transfection efficiency. miR-486-3p mimic and NC were cotransfected into 293T cells, respectively, with the luciferase reporter vectors. The cells were collected 48 h after transfection, lysed, and measured according to the manufacturer’s instructions of the Dual Luciferase Reporter Gene Assay Kit (GM-040502A; Qcbio Science & Technologies, Shanghai, China). The ratio of the relative luminescence units (RLU) of firefly to that of the renilla was used to determine the binding intensity.

**AGO2 Immunoprecipitation**

miR-486-3p and miR-486-3p-mut were separately transfected into NPCs. After 48 h, AGO2 was immunoprecipitated with the AGO2-specific antibody, where the antibody to IgG served as NC. The cells were dissolved in 150 mM KCl, 25 mM Tris-HCl (pH 7.4), 5 mM ethylenediaminetetraacetic acid (EDTA), 0.5% Triton X-100, and 5 mM dithiothreitol (DTT) + Ribonlock (Fermentas MBI, Pittsburgh, PA, USA), along with the proteinase inhibitor cocktail (Roche Applied Science, Mannheim, Germany). The lysate was mixed with the antibody-coupled Sepharose microballs, rotated at 4°C for 4 h, and then washed six times with the lysis buffer prior to extraction of the RNA with the Trizol reagent (Invitrogen, Carlsbad, CA, USA).

**Northern Blot Analysis**

Northern blot analysis was performed with the Northern Blot Kit (Ambion, Austin, TX, USA). In brief, portions of total RNA (30 μg) were denatured in formaldehyde, followed by electrophoresis in 1% agarose formaldehyde gel. RNA was then transferred onto the Hybond-N + nylon membrane (Beyotime Biotechnology, Shanghai, China) and hybridized with a biotin-labeled DNA probe. Detection of bound RNA was performed using the biotin assay kit (Thermo Fisher Scientific, Waltham, MA, USA). Finally, the membrane was developed and analyzed by Image Lab software (Bio-Rad Laboratories, Hercules, CA, USA).

**Establishment of a Mouse Model of IDD**

A total of 24 C57BL/6J mice (Shanghai SLAC Experimental Animal, Shanghai, China; aged 10–14 weeks) were included in the experiment. Eighteen mice received surgery, and the remaining six mice were free of surgical intervention, thus serving as the NC group. For induction of anesthesia, the mice were injected intraperitoneally with ketamine (90 mg/kg) and dimethylrazine (10 mg/kg) and placed in a prone position. Under the guidance of fluoroscopy, the dorsal sides of coccygeal (Co) 6/7, 8/9, and 10/11 IVDs were punctured with a 20-gauge needle. The needle passed through the center of the disc, penetrating to the opposite side, rotated 180°, and was held in place for 10 s. Both Co 7/8 and 9/10 were left intact. After the operation, the wound was covered with the gauze and sutured according to postoperative standards of care.

1 week after IVD puncture, the mice were randomly assigned to treatment groups with injection of si-circ_001653 or OE-circ_001653 + miR-486-3p agomir or no injection (n = 8). After inducing anesthesia, a small incision was made to expose the previously punctured disc.
from the left side. A 33-gauge needle tip (Hamilton, Bonaduz, Switzerland), filled with 2 μL of virus vectors containing human circ_001653 and OE-circ_001653 + miR-486-3p agomir, was inserted within the disk capsule intradiscally and connected to a microsyringe (Hamilton, Bonaduz, Switzerland) for infusion of the virus. The treatment was repeated 4 weeks later.

The IVD height was measured at the 1st, 5th, and 9th weeks after treatment was repeated 4 weeks later. All IVD tissues from mice were sectioned, dewaxed, and stained with hematoxylin for 3 min. After being washed three times, sections were counter-stained with hematoxylin for 1 min, mounted, and photographed under a microscope. The positive expression showed as granules of yellow or brownish yellow. The proportion of positive cells was calculated in five randomly selected fields, each containing a total of 200 cells.

Safranin O Staining

IVD tissues from mice were sectioned, dewaxed, and stained with hematoxylin for 3 min. After being washed three times, sections were differentiated with 1% hydrochloric acid in ethanol for 5 s. Then, sections were stained with 0.02% fast green solution for 10 min, washed in 1% acetic acid, and counterstained with 0.1% Safranin O solution for 3 min. After dehydration through an alcohol series and being rendered transparent with xylene, sections were mounted with neutral balsam and observed under a microscope.

Statistical Analysis

All data were processed using SPSS 21.0 statistical software (IBM, Armonk, NY, USA). The measurement data with normal distribution were expressed as mean ± standard deviation. The statistical significance of data between two groups was calculated using the independent t test. Comparisons between groups were conducted by one-way analysis of variance. The repeated-measures analysis of variance was applied for the comparison of data at different time points, followed by Tukey's post hoc test. The relationship between circ_001653 and IVD tissue degeneration was analyzed by Spearman rank correlation analysis. The correlation between circ_001653 and miR-486-3p and between miR-486-3p and CEMIP in IVD tissues was analyzed by Pearson's correlation analysis. p < 0.05 indicates the significantly statistical difference.

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