RNA-Sequencing Profiling Analysis of Pericyte-Derived Extracellular Vesicle-Mimetic Nanovesicles-Regulated Genes in Primary Cultured Fibroblasts From Normal and Peyronie's Disease Penile Tunica Albuginea

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Research article

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Abstract

Background

Peyronie's disease (PD) is a severe fibrotic disease of the tunica albuginea that causes penis curvature and leads to penile pain, deformity, and erectile dysfunction. The role of pericytes in the pathogenesis of fibrosis has recently been determined. Extracellular vesicle (EV)-mimetic nanovesicles (NVs) have attracted attention regarding intercellular communication between cells in the field of fibrosis. However, the global gene expression of pericyte-derived EV-mimetic NVs (PC-NVs) in regulating fibrosis remains unknown. Here, we used RNA-sequencing technology to investigate the potential target genes regulated by PC-NVs in primary fibroblasts derived from human PD plaque.

Methods

Human primary fibroblasts derived from normal and PD patients was cultured and treated with cavernosum pericytes isolated extracellular vesicle (EV)-mimetic nanovesicles (NVs). A global gene expression RNA-sequencing assay was performed on normal fibroblasts, PD fibroblasts, and PD fibroblasts treated with PC-NVs. Reverse transcription polymerase chain reaction (RT-PCR) was used for sequencing data validation.

Results

A total of 4135 genes showed significantly differential expression in the normal fibroblasts, PD fibroblasts, and PD fibroblasts treated with PC-NVs. However, only 91 contra-regulated genes were detected among the three libraries. Furthermore, 20 contra-regulated genes were selected and 11 showed consistent changes in the RNA-sequencing assay, which were validated by RT-PCR.

Conclusion

The gene expression profiling results suggested that these validated genes may be good targets for understanding potential mechanisms and conducting molecular studies into PD.

Background

Peyronie’s disease (PD) is caused by excessive fibrosis and scar tissue formation in the tunica albuginea (TA), resulting in penile pain, abnormal curvature, and erectile dysfunction (ED) [1, 2]. Although the existence of PD has been known for a long time, the pathophysiology of PD has not been studied as widely as fibrosis in other organs, such as the kidneys, liver, or lungs. Currently, the most available medical therapy is collagenase and interferon injection and surgical intervention [3, 4]. However, these treatments can cause glandular hypoesthesia and a high risk of new onset ED [5]. Therefore, the identification of novel therapeutic targets related to PD fibrosis is required.
Pericytes play a fundamental role in vascular contractility and stability, regulation of vascular development, and as a storage vault of mesenchymal stem cells [6, 7]. In vitro studies have shown that pericytes exhibit fibrogenic potential [8, 9] and transition to myofibroblasts [10]. Moreover, the inhibition of angiogenesis may be effective in the suppression of fibrosis [11]. However, recently studies have shown that the inhibition of angiogenesis may aggravate fibrosis [12, 13]. These findings suggest that different antiangiogenic and molecular targets produce different results in the treatment of fibrosis.

Extracellular vesicles (EVs) were previously believed to be cell excretions. However, a number of studies have shown that EVs contain proteins, lipids, and RNA, which can affect the physiological and pathological communications between cells [14, 15]. Many studies regarding the potential role of EVs have been conducted for human diseases, including strokes [16], tumor metastasis [17], and kidney disease [18]. Therefore, clarifying the role of EVs in fibrosis would be beneficial to aid in the understanding of fibrosis mechanisms. However, one of the major limitations of EVs is the low production yield [19]. Therefore, to maximize the production of vesicles, we used a mini extruder system and extracted more than 100-fold greater EV-mimetic NVs from pericytes. The cell-derived EV-mimetic NVs showed similar characteristics to the natural EVs [20]. These studies suggest that pericyte-derived EV-mimetic NVs (PC-NVs) may be beneficial for the functional study of fibrosis.

Gene expression profiling analysis in physiological and pathological conditions can provide a foundation for studying the mechanisms of fibrosis in PD. In the present study, we performed an RNA-sequencing assay on normal fibroblasts, PD fibroblasts, and PD fibroblasts treated with PC-NVs.

**Methods**

**Ethics statement and Study design**

All TA tissues and animals used in this study were approved by the Institutional Review Board (IRB No: 2007-730) and the Institutional Animal Care and Use Committee of our University (approval number: 171129-527), respectively. The plaque tissue of a patient with PD (48 years old) and the normal TA tissue from control patients (undergoing penoplasty for congenital curvature, 21 years old) were used for the human fibroblast culture study. In addition, 10 adult male C57BL/6J mice (8 weeks old, Orient Bio, Korea) were used for the mouse cavernous pericytes (MCPs) primary culture.

**Primary culture and characterization of human fibroblasts**

The TA tissues were used for the primary fibroblast culture as described previously [21, 22]. Briefly, PD plaque and normal TA tissues were maintained in sterile vials with Hank's balanced salt solution (HBSS, Gibco, Carlsbad, CA, USA) and washed three times with phosphate-buffered saline (PBS). The TA tissues were cut into 1-2 mm sections and incubated in 12.5 mL Dulbecco's modified Eagle's medium (DMEM, Gibco) supplemented with 0.06% collagenase A (Sigma-Aldrich, St. Louis, MO, USA) at 37°C for 1 h in a 5% CO₂ atmosphere. The cells and tissue fragments were collected by centrifugation (400 g for 5 min), washed with fresh culture medium, and placed in 100 mm cell culture dishes (Falcon-Becton Dickinson...
Labware, Franklin Lakes, NJ, USA) with DMEM containing 10% fetal bovine serum (FBS), penicillin (100 U/mL), and streptomycin (100 μg/mL) at 37°C in a 5% CO₂ atmosphere. Media were changed every 2 days and the cells were characterized as previously described [21, 22]. Passages 5 to 8 were used for the experiments.

To determine cell type, the cells were cultured on sterile cover glasses, placed into 12-well plates and grown until nearly confluent. The cells were stained with antibody to CD90 (fibroblast marker, R&D Systems Inc., Minneapolis, MN, USA; 1:100), Vimentin (fibroblast marker, R&D Systems Inc., 1:50), NG2 chondroitin sulfate proteoglycan (NG2, pericyte marker, Millipore, San Francisco, CA, USA; 1:50), CD31 (endothelial cell marker, Chemicon, Temecula, CA, USA; 1:50), or DAPI (nucleus marker; Vector Laboratories Inc., Burlingame, CA, USA), as previously described [23]. Signals were visualized and digital images were obtained using a confocal fluorescence microscope (K1-Fluo, Nanoscope Systems, Inc., Daejeon, Korea).

**Primary culture of MCPs**

The primary cultures of MCPs were performed as described previously [24, 25]. Shortly, 8 weeks old male C57BL/6J mice were anesthetized with ketamine (100 mg/kg) and xylazine (5 mg/kg) intramuscularly, and sacrificed by cervical dislocation. Then, the penis tissues were harvested and maintained in sterile vials with HBSS (Gibco). After washing three times with PBS, the urethra and dorsal neurovascular bundle were removed, and only the corpus cavernosum tissues were used. The corpus cavernosum tissues were cut into approximately 1-2 mm sections and settled via gravity into collagen I-coated 35 mm cell culture dishes with 300 μL complement DMEM (GIBCO) at 37°C for 20 min in a 5% CO₂ atmosphere. Thereafter, 900 μL of complement medium was added and incubated at 37°C with 5% CO₂. The complement medium contained 20% FBS, 1% penicillin/streptomycin, and 10 nM human pigment epithelium-derived factor (PEDF; Sigma-Aldrich). The medium was changed every 2 days, and after approximately 10 days sprouting cells were sub-cultured into collagen I (Advanced BioMatrix, San Diego, CA, USA)-coated dishes. Cells from passages 2 to 3 were used for the experiments.

**Preparation and characterization of MCP-derived EV-mimetic nanovesicles (NVs)**

MCP-derived EV-mimetic NVs (PC-NVs) were prepared using a mini extruder system (Avanti Polar Lipids, Birmingham, AL, USA), as described previously [26, 27]. Briefly, MCPs were washed three times with PBS, detached with 0.25% Trypsin-EDTA (Invitrogen, Carlsbad, CA, USA) and re-suspended in 4-(2-Hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES) buffer solution (Gibco). The cell suspension was sequentially extruded 10 times across 10, 5, and 1 μm pore-sized polycarbonate membranes (Nuclepore, Whatman Inc., Clifton, NJ, USA), respectively. Next, ultracentrifugation was performed at 100,000 g for 2 h at 4°C with a step gradient, which was formed with 50% iodixanol (1 mL; Axis-Shield PoC AS, Oslo, Norway) overlaid with 10% iodixanol (2 mL) and topside with the extruded samples (7 mL). PC-NVs were filtered with a 0.45 μm filter and stored at -80°C until further use. To quantify the PC-NVs, the EXOCET
exosome quantitation assay kit (System Biosciences, Palo Alto, CA, USA) was used, and 1 µg/µL of the final concentration of the PC-NVs was prepared for all experiments.

**Western blotting**

For the immunoblot analyses of PC-NVs, equal protein amounts (10 µg) of purified PC-NVs and whole cells extracted using RIPA lysis buffer (Sigma-Aldrich) were separated by SDS-PAGE (12% gel) and transferred to polyvinylidene fluoride (PVDF) membranes. Each blot was blocked and incubated with antibodies to GM130 (BD Biosciences, San Jose, CA, USA; 1:1000), CD9 (Abcam, Cambridge, UK; 1:1000), CD81 (Novus Biologicals; 1:1000), or TSG101 (Novus Biologicals; 1:500).

**RNA-sequencing assay**

For the RNA-sequencing study, the normal and PD TA-derived fibroblasts were cultured and treated with PC-NVs (n = 4 per group). The RNA-sequencing assay was performed by E-Biogen Inc. (Korea). Briefly, total RNA was isolated 24 h after exposure to PC-NVs using TRIzol reagent (Invitrogen). RNA quality was assessed using an Agilent 2100 Bioanalyzer (Agilent Technologies, Amstelveen, The Netherlands), and RNA quantification was performed using an ND-2000 Spectrophotometer (Thermo Inc., DE, USA).

**Library sequencing and data analysis**

Libraries were prepared from total RNA using the SMARTer Stranded RNA-Seq Kit (Clontech Laboratories, Inc., USA). The isolation of mRNA was performed using the Poly(A) RNA Selection Kit (LEXOGEN, Inc., Austria). Indexing was performed using the Illumina indices 1-12. The enrichment step was performed using PCR. Subsequently, libraries were checked using the Agilent 2100 Bioanalyzer (DNA High Sensitivity Kit) to evaluate the mean fragment size. Quantification was performed using the library quantification kit using a StepOne Real-Time PCR System (Life Technologies, Inc., USA). High-throughput sequencing was performed as paired-end 100 sequencing using HiSeq 2500 (Illumina, Inc., USA).

Quality control of the raw sequencing data was performed using FastQC (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Adapter and low-quality reads (<Q20) were removed using FASTX_Trimmer (http://hannonlab.cshl.edu/fastx_toolkit/) and BBMap (https://sourceforge.net/projects/bbmap/). Then, the trimmed reads were mapped to the reference genome using TopHat [28]. Gene expression levels were estimated using RC (read count) and FPKM (fragments per kb per million reads) values by BEDTools [29] and Cufflinks [30]. The expression values were normalized with the Quantile normalization method using EdgeR within R (https://www.r-project.org). Data mining and graphic visualization were performed using ExDEGA (E-Biogen, Inc., Korea). The RNA-sequencing data have been deposited in the Gene Expression Omnibus database (www.ncbi.nlm.nih.gov/geo accession no. GSE146500).

**Validation of sequencing data by RT-PCR**
Total RNA was extracted from cultured cells using TRIzol (Invitrogen) following the manufacturer's protocols. Reverse transcription was performed using 1 µg of RNA in 20 µL of reaction buffer with oligo dT primer and AccuPower RT Premix (Bioneer Inc., Korea). The PCR reaction was performed with denaturation at 94°C for 30 s, annealing at 60°C for 30 s, and extension at 72°C for 1 min in a DNA Engine Tetrad Peltier Thermal Cycler. For the analysis of PCR products, 10 µL of each PCR product was electrophoresed on a 1% agarose gel and detected under ultraviolet light. GAPDH was used as an internal control [31].

All digital image, western blot, and PCR band densitometry analyses were performed using an image analyzer system (National Institutes of Health [NIH] ImageJ 1.34, http://rsbweb.nih.gov/ij/).

Statistical analysis

All data are expressed as means ± standard errors. Statistical analysis was performed using Student t-test. P values less than 0.05 were considered statistically significant.

Results

Identification of human fibroblasts

The fibroblasts were isolated from human normal and PD plaque tissues. Representative images showed high positive staining for CD90 and Vimentin (fibroblast markers) of more than 95%, but not for pericyte (NG2) or endothelial cell (CD31) markers (Fig. 1a, b).

PC-NV preparation and characterization

PC-NVs were prepared from MCPs according to previous methods [26]. Western blot analysis showed that PC-NVs displayed positive exosomes markers, including CD9, CD81, and TSG101, but not for negative marker GM130 (Fig. 1c, d).

Transcriptional profiling and gene ontology (GO) category analysis

For this study, three gene libraries for the normal fibroblast (NF), PD fibroblast (PF), and PC-NVs-treated PF (PFPC) groups were constructed for an RNA-sequencing assay (n = 4 for each group). In total, 25737 genes were detected in three libraries. Significant gene selection was performed with three conditions: fold-change > 2.0, log2 > 4, and p-value < 0.05. Among all detected genes, 3961 showed significant differential expression in the PF group compared with the NF group, and 174 were significantly differentially expressed in the PFPC group compared with the PF group (Fig. 2a, b, c). Only 91 contra-regulated genes (Supplementary Table S1) were detected between PF/NF and PFPC/PF through Venn diagram analysis (Fig. 2d).

To evaluate the GO categories, all significantly differentially expressed genes (DEGs) were classified in 16 GO categories by ExDEGA software (E-Biogen, Inc., Korea). Among these, extracellular matrix (33.61%),
angiogenesis (32.33%), and fibrosis (31.07%) made up the largest proportion in the PF group compared with the NF group (Fig. 3a, b). However, fibrosis (3.88%), extracellular matrix (2.92%), and angiogenesis (2.59%) made up the largest proportion in the PFPC group compared with the PF group (Fig. 3c, d).

**Validation of RNA-sequencing results by RT-PCR**

To validate the RNA-sequencing results, we selected 20 genes (Supplementary Table S2) from 91 contra-regulated DEGs, and 11 (primers as shown in Supplementary Table S3) showed results consistent with the RNA-sequencing assay by RT-PCR. Among these genes, **MMP3, AKR1C1, SMOC1, ANGPTL2, SEMA3A, TRIM15, EGR1, and BMP2** were downregulated in the PF group compared with the NF group, and were significantly recovered in the PFPC group (Fig. 4a, c). Only **TFPI2, SFRP4, and SERPINE1** were induced in the PF group and recovered in the PFPC group (Fig. 4b, d).

**Discussion**

The accurate physiological and pathological mechanisms of PD remain poorly understood. To date, most gene expression studies have focused on human PD plaque at a tissue level **in vivo** [32, 33] but not at a cellular level **in vitro**. Therefore, to investigate the exact mechanisms and potential target genes for PD, we cultured human fibroblasts from human PD plaque and performed RNA-sequencing assays.

EVs display a potential role in kidney fibrosis and other fibrotic diseases [18, 34]; however, little is currently known regarding the detailed mechanisms. Considering the low yield of EVs, we extracted more than a 100-fold greater EV-mimetic NVs from MCPs, which were primarily cultured from mouse corpus cavernosum tissues. Many previous studies have found that pericytes display diverse features in relation to fibrosis that are dependent on different molecular targets [8, 9, 11]. In this study, human PD fibroblasts exposed to PC-NVs were compared with human PD fibroblasts to investigate the regulation of gene expression by PC-NVs in PD.

From the RNA-sequencing assay, 3961 DEGs were detected, and the 16 top GO categories were assessed in this study. GO analysis showed that significantly altered genes were enriched in the extracellular matrix, angiogenesis, and fibrosis. The extracellular matrix is a driver of progressive fibrosis [35], and angiogenesis is closely associated with chronic liver fibrosis [18]. These data suggest that our DEG detection is credible. However, the molecular basis of PC-NVs in regulating the extracellular matrix or angiogenesis pathway in PD remains largely unknown. In this study, only 91 contra-regulated genes were identified from the three libraries (NF, PF, and PFPC). After precision screening, 20 genes were selected and validated by RT-PCR in same conditions. However, only 11 genes were validated to be consistent with the RNA-sequencing results. These genes may be the key to understanding how PC-NVs regulate the extracellular matrix, angiogenesis, and fibrosis mechanisms in PD.

To the best of our knowledge, this is the first study to demonstrate the systematic profiling of gene alterations in NF, PF, and PFPC. However, the present study has some limitations. First, a small number of cultured human fibroblast samples were used in target gene validation and age differences existed...
among groups. Second, we were unable to demonstrate the network of these validated genes in the extracellular matrix, angiogenesis, and fibrosis pathways. Third, mouse corpus cavernous pericytes were used for EV-mimetic NVs isolation, and further studies are required to evaluate the role of human corpus cavernous pericytes isolated with EV-mimetic NVs in PD.

**Conclusion**

In summary, we profiled the DEGs of human TA cultured fibroblasts in NF, PF, and PFPC groups. We hypothesize that these validated genes are good candidates for the study of the mechanism of PC-NVs in PD. Further studies exploring the effect of these target genes will be beneficial to further our understanding of the detailed mechanisms of the extracellular matrix, angiogenesis, and fibrosis in PD.

**Abbreviations**

**PD**: Peyronie’s disease  
**EV**: Extracellular vesicle  
**NVs**: Nanovesicles  
**PC-NVs**: Pericyte-derived EV-mimetic NVs  
**RT-PCR**: Reverse transcription polymerase chain reaction  
**TA**: Tunica albuginea  
**ED**: Erectile dysfunction  
**MCPs**: Mouse cavernous pericytes  
**HBSS**: Hank’s balanced salt solution  
**PBS**: Phosphate-buffered saline  
**DMEM**: Dulbecco’s modified Eagle’s medium  
**FBS**: Fetal bovine serum  
**PEDF**: Pigment epithelium-derived factor  
**HEPES**: 4-(2-Hydroxyethyl)piperazine-1-ethanesulfonic acid  
**PVDF**: Polyvinylidene fluoride  

**Declarations**
Ethics approval and consent to participate

All TA tissues donors provided written informed consent and the procedures were approved by the Institutional Review Board (IRB No: 2007-730). And all male C57BL/6J mice (8 weeks old, Orient Bio, Korea) used in this study were approved by the Institutional Animal Care and Use Committee of our University (approval number: 171129-527), respectively. All authors of the manuscript have read and agreed to its content of the abstract.

Availability of data and material

Not applicable

Consent for Publication

Not applicable

Competing interests

The authors declare that they have no competing interests.

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Author's contributions

GNY, SGP, JKS and JKR conception and design of the research. GNY, SGP, JYO, MHK performed the experiments. GNY, SGP, ZYL, LW, JYO, DKK, analyzed the data. GNY, SGP prepared the figures and drafted the manuscript. YSG, JKS and JKR edited and revised the manuscript. GNY, SGP, ZYL, LW, JYO, MHK, DKK, YSG, JKS and JKR have read and approved the final version of the manuscript. All authors are accountable for all aspects of work.

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References
1. Gholami SS, Gonzalez-Cadavid NF, Lin CS, Rajfer J and Lue TF (2003) Peyronie's disease: a review. J Urol 169:1234-1241
2. Hellstrom WJ and Bivalacqua TJ (2000) Peyronie's disease: etiology, medical, and surgical therapy. J Androl 21:347-354
3. Duncan MR, Berman B and Nseyo UO (1991) Regulation of the proliferation and biosynthetic activities of cultured human Peyronie's disease fibroblasts by interferons-alpha, -beta and -gamma. Scand J Urol Nephrol 25:89-94
4. Russo GI, Milenkovic U, Hellstrom W, Levine LA, Ralph D and Albersen M (2018) Clinical Efficacy of Injection and Mechanical Therapy for Peyronie's Disease: A Systematic Review of the Literature. Eur Urol 74:767-781
5. Levine LA, Greenfield JM and Estrada CR (2005) Erectile dysfunction following surgical correction of Peyronie's disease and a pilot study of the use of sildenafil citrate rehabilitation for postoperative erectile dysfunction. J Sex Med 2:241-247
6. Armulik A, Abramsson A and Betsholtz C (2005) Endothelial/pericyte interactions. Circ Res 97:512-523
7. Armulik A, Genove G and Betsholtz C (2011) Pericytes: developmental, physiological, and pathological perspectives, problems, and promises. Dev Cell 21:193-215
8. Crisan M, Yap S, Casteilla L et al (2008) A perivascular origin for mesenchymal stem cells in multiple human organs. Cell Stem Cell 3:301-313
9. Cai X, Lin Y, Friedrich CC et al (2009) Bone marrow derived pluripotent cells are pericytes which contribute to vascularization. Stem Cell Rev Rep 5:437-445
10. Lin SL, Kisseleva T, Brenner DA and Duffield JS (2008) Pericytes and perivascular fibroblasts are the primary source of collagen-producing cells in obstructive fibrosis of the kidney. Am J Pathol 173:1617-1627
11. Yoshiji H, Kuriyama S, Yoshii J et al (2003) Vascular endothelial growth factor and receptor interaction is a prerequisite for murine hepatic fibrogenesis. Gut 52:1347-1354
12. Patsenker E, Popov Y, Stickel F et al (2009) Pharmacological inhibition of integrin alphavbeta3 aggravates experimental liver fibrosis and suppresses hepatic angiogenesis. Hepatology 50:1501-1511
13. Stockmann C, Kerdiles Y, Nomaksteinsky M et al (2010) Loss of myeloid cell-derived vascular endothelial growth factor accelerates fibrosis. Proc Natl Acad Sci U S A 107:4329-4334
14. Colombo M, Raposo G and Thery C (2014) Biogenesis, secretion, and intercellular interactions of exosomes and other extracellular vesicles. Annu Rev Cell Dev Biol 30:255-289
15. Record M, Carayon K, Poirot M and Silvente-Poirot S (2014) Exosomes as new vesicular lipid transporters involved in cell-cell communication and various pathophysiologies. Biochim Biophys Acta 1841:108-120
16. Zhang ZG and Chopp M (2016) Exosomes in stroke pathogenesis and therapy. J Clin Invest 126:1190-1197
17. Li K, Chen Y, Li A, Tan C and Liu X (2019) Exosomes play roles in sequential processes of tumor metastasis. Int J Cancer 144:1486-1495
18. Zhang W, Zhou X, Zhang H, Yao Q, Liu Y and Dong Z (2016) Extracellular vesicles in diagnosis and therapy of kidney diseases. Am J Physiol Renal Physiol 311:F844-F851
19. Kim OY, Lee J and Gho YS (2017) Extracellular vesicle mimetics: Novel alternatives to extracellular vesicle-based theranostics, drug delivery, and vaccines. Semin Cell Dev Biol 67:74-82
20. Jang SC, Kim OY, Yoon CM et al (2013) Bioinspired exosome-mimetic nanovesicles for targeted delivery of chemotherapeutics to malignant tumors. ACS Nano 7:7698-7710
21. Piao S, Choi MJ, Tumurbaatar M et al (2010) Transforming growth factor (TGF)-beta type I receptor kinase (ALK5) inhibitor alleviates profibrotic TGF-beta1 responses in fibroblasts derived from Peyronie's plaque. J Sex Med 7:3385-3395
22. Ryu JK, Kim WJ, Choi MJ et al (2013) Inhibition of histone deacetylase 2 mitigates profibrotic TGF-beta1 responses in fibroblasts derived from Peyronie's plaque. Asian J Androl 15:640-645
23. Yin GN, Ryu JK, Kwon MH et al (2012) Matrigel-based sprouting endothelial cell culture system from mouse corpus cavernosum is potentially useful for the study of endothelial and erectile dysfunction related to high-glucose exposure. J Sex Med 9:1760-1772
24. Neng L, Zhang W, Hassan A et al (2013) Isolation and culture of endothelial cells, pericytes and perivascular resident macrophage-like melanocytes from the young mouse ear. Nat Protoc 8:709-720
25. Yin GN, Das ND, Choi MJ et al (2015) The pericyte as a cellular regulator of penile erection and a novel therapeutic target for erectile dysfunction. Sci Rep 5:10891
26. Kwon MH, Song KM, Limanjaya A et al (2019) Embryonic stem cell-derived extracellular vesicle-mimetic nanovesicles rescue erectile function by enhancing penile neurovascular regeneration in the streptozotocin-induced diabetic mouse. Sci Rep 9:20072
27. Oh K, Kim SR, Kim DK et al (2015) In Vivo Differentiation of Therapeutic Insulin-Producing Cells from Bone Marrow Cells via Extracellular Vesicle-Mimetic Nanovesicles. ACS Nano 9:11718-11727
28. Trapnell C, Pachter L and Salzberg SL (2009) TopHat: discovering splice junctions with RNA-Seq. Bioinformatics 25:1105-1111
29. Quinlan AR and Hall IM (2010) BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26:841-842
30. Roberts A, Trapnell C, Donaghey J, Rinn JL and Pachter L (2011) Improving RNA-Seq expression estimates by correcting for fragment bias. Genome Biol 12:R22
31. Yin GN, Jin HR, Choi MJ et al (2018) Pericyte-Derived Dickkopf2 Regenerates Damaged Penile Neurovasculature Through an Angiopoietin-1-Tie2 Pathway. Diabetes 67:1149-1161
32. Wang Z, Liu JF, Zhou ZH et al (2004) [Gene expression profiles and effects of transforming growth factor-beta1 intervention in Peyronie's disease]. Zhonghua Wai Ke Za Zhi 42:182-186
33. Sampaio FJ (2004) Comparison of gene expression profiles between Peyronie's disease and Dupuytren's contracture. Int Braz J Urol 30:349-350

34. Asef A, Mortaz E, Jamaati H and Velayati A (2018) Immunologic Role of Extracellular Vesicles and Exosomes in the Pathogenesis of Cystic Fibrosis. Tanaffos 17:66-72

35. Bruno S, Porta S and Bussolati B (2016) Extracellular vesicles in renal tissue damage and regeneration. Eur J Pharmacol 790:83-91