

Prediction and analysis of osteoarthritis hub genes with bioinformatics

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Background: Osteoarthritis (OA) is the most common type of arthritis. OA can cause joint pain, stiffness, and loss of function. The pathogenesis of OA is not completely clear. Moreover, there is no effective treatment, and clinical management is limited to symptomatic relief or joint surgery. This study utilized bioinformatics to analyze normal and OA articular cartilage samples to find biomarkers and therapeutic targets for OA.

Methods: The GSE169077 gene chip dataset was downloaded from the public gene chip data platform of the National Biotechnology Information Center. The dataset included 6 samples of OA tissues and 5 samples of healthy cartilage tissues. Differentially expressed genes (DEGs) were screened using the R language "limma" function package under the threshold of $\log_2[fold change (FC)] \ge 2$ and a P value <0.05. The Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) signal pathways of the target genes were enriched and analyzed using the database for annotation, visualization, and integrated discovery (DAVID), and a protein-protein interaction (PPI) network was further constructed using the search tool for the retrieval of interacting genes/proteins (STRING) database. The coexpression relationship of the genes in the module was visualized and screened with Cytoscape.

Results: A total of 27 DEGs were identified, including 9 downregulated genes and 18 upregulated genes. GO signal pathway enrichment analysis showed involvement in hypoxic response, fibrous collagen trimer, and extracellular matrix structural components. KEGG analysis demonstrated associations with protein digestion and absorption, extracellular matrix receptor interaction, and the peroxisome proliferator-activated receptor signal pathway, among several other pathways. A PPI network was obtained through STRING analysis, and the results were imported into Cytoscape software. The 27 DEGs were sequenced by the cytoHubba plug-in by various calculation methods, and 5 hub genes (*COL1A1, COL1A2, POSTN, BMP1*, and *MMP13*) were finally selected. These genes were analyzed by PPI again and annotated with GO and KEGG in different colors.

Conclusions: Bioinformatics technology effectively identified differential genes in the knee cartilage tissue of healthy controls and patients with OA, providing opportunities to further explore the mechanism and treatment of OA on a transcriptional level.

Keywords: Osteoarthritis (OA); differentially expressed genes (DEGs); hub gene; enrichment analysis

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Introduction

Osteoarthritis (OA) is the most common form of arthritis worldwide, affecting approximately 40% of people over 70 years old (1). According to the World Health Organization, the number of people with OA will double by 2025 and again by 2050. OA can cause joint pain, stiffness, and loss of function. It is characterized by cartilage erosion, osteophyte formation, subchondral bone formation, and synovial inflammation due to biomechanical and biochemical joint changes (2). Since the first description of hip OA in 1793 (3), a large amount of in-depth research has been done on its pathophysiology (4-6). To date, the "gold standard" for clinical diagnosis of OA is still imageology. However, there is a lag in imaging examinations, which often require 1–3 years to obtain reliable information on progression (7).

As high throughput sequencing technology has advanced and costs have fallen, bioinformatics analysis is now widely used to identify disease-specific biomarkers (7-10) for screening, diagnosis, prediction, and identification of drug targets of tumor genes. In recent years, researchers have reported the use of bioinformatics and microarray technology to screen the genes and signal transduction pathways of OA, providing theoretical support for the pathogenesis of OA and new therapeutic targets (11,12). Although the pathogenesis of OA is not fully known, we can refer to its main risk factors (13). Cartilage tissue in OA may be the first to undergo pathophysiological changes, including cartilage matrix proteolysis, cartilage cell erosion, and fibrosis, while the release of collagen fragments and

Highlight box

Key findings

• Analysis of cartilage samples for osteoarthritis based on known biological information identified differential genes, providing potential biomarkers and therapeutic targets for osteoarthritis.

What is known and what is new?

- Osteoarthritis still has no definitive biomarkers for a definitive diagnosis, and there are no definitive drugs available for its treatment.
- Comparative access to biomarkers of osteoarthritis using biological information from known transcriptomes may be an effective solution to this problem.

What is the implication, and what should change now?

• The discovery of biomarkers for osteoarthritis can help drug development and clinicians to detect osteoarthritis at an early stage for early diagnosis and treatment.

proteoglycans into synovial fluid leads to synovial tissue inflammation (14). Data from current bioinformatics studies on OA are mostly from synovial tissue (15,16) and peripheral blood (11), and some scholars believe that although inflammatory response as a major mechanism may be involved in the pathogenesis of OA (17), local or systemic reactions are affected by other factors.

Many driving factors (18-20) have been found to induce OA development, including mechanical injury, aging, cartilage degeneration, and synovitis, among others. A major challenge for researchers is applying bioinformatics to find the switch that drives OA initiation in a wide variety of sample types so that it can be used to improve the early diagnosis and prediction of OA. Therefore, we speculated that looking for "the Big Bang" might be the key to better understanding OA. Further, bioinformatics is usually calculated by the cytoHubba plug-in of Cytoscape software, which contains many kinds of calculation methods. Most previous research has involved a single method (15,21), and it is unclear whether there are differences in the hub genes obtained by the various calculation methods.

To address the above problems, we employed biological information analysis of OA cartilage tissue rather than synovium and blood samples. We then performed cluster analysis of differential genes, and grouped and compared functional paths. In addition, we selected multiple computational models for the screening of hub genes to avoid the bias of using a single model. Finally, proteinprotein interaction (PPI) analysis of the acquired hub genes was conducted to provide a more intuitive understanding of their interrelationships. *Figure 1* shows a workflow of the steps undertaken in this study. We present the following article in accordance with the STREGA reporting checklist (available at https://atm.amegroups.com/article/ view/10.21037/atm-22-6450/rc).

Methods

Data sources

The GSE169077 dataset was downloaded from the National Biotechnology Information Center's Gene Expression Omnibus (GEO) public database (https://www.ncbi.nlm. nih.gov/geo/) (22). The chip platform used was the GPL96 (HG-U133A) Affymetrix Human Genome U133A Array. The study used data sets from 11 knee replacement surgery patients. The patient data in this work were obtained from a public database, which included informed consent and

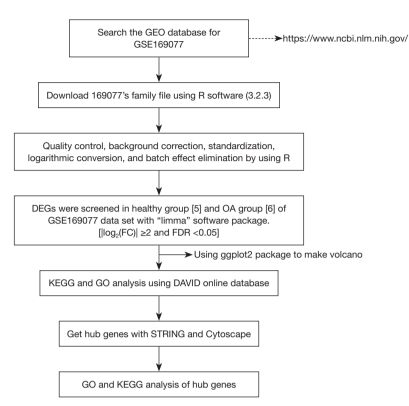


Figure 1 Workflow for this study. The workflow demonstrates the important steps undertaken in this study, including KEGG and GO analysis, obtaining hub genes, and secondary analysis of hub genes. KEGG, Kyoto Encyclopedia of Genes and Genomes; GO, Gene Ontology.

ethical approval. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

Screening of differentially expressed genes (DEGs)

R version 3.2.3 (The R Project for Statistical Computing, Vienna, Austria) was used to analyze and read the chip and download the matrix file. The "affy" (23) and "limma" packages (24) were employed for quality control, background correction, standardization, logarithmic conversion, and batch effect elimination. Samples without clinical data were filtered, and the rest of the data were analyzed. The "limma" software package was used to screen DEGs among 5 normal controls and 6 OA samples from the GSE169077 dataset. An absolute value of the log₂ difference multiple [fold change (FC)] \geq 2 and false discovery rate (FDR) <0.05 were set as the thresholds for screening DEGs (25). The "ggplot2" package was utilized to draw the volcano map for the obtained DEG data (26).

Statistical analysis

After gene expression quantification is completed, statistical analysis of their expression data is required to screen the samples for genes with significantly different expression levels in different states. First, to correct for the sequencing depth, we normalized the original readcount; then the statistical model was used to calculate the probability of hypothesis testing (P value), and finally multiple hypothesis testing was performed to correct for the FDR.

Enrichment analysis of the DEGs

The online database for annotation, visualization, and integrated discovery (DAVID, https://david.ncifcrf.gov) (27) was used to perform gene ontology (GO) (28) and Kyoto Encyclopedia of Gene and Genomes (KEGG) pathway enrichment analysis (29). GO functional enrichment data were annotated and classified according to the biological

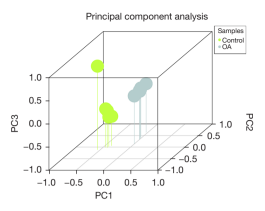


Figure 2 GSE169077 principal component analysis. The data of 11 samples from the control and OA groups were analyzed by PCA, and there were significant differences between the 2 groups. OA, osteoarthritis; PCA, principal component analysis.

pathway (BP), cellular composition (CC), and molecular function (MF). KEGG enrichment analysis was performed to identify the possible biological pathways. Using the Expression Analysis Systematic Explorer (EASE) score statistical method (30), the screening conditions of the GO and KEGG data were set at a P value <0.05. The upregulated and downregulated DEGs were explored with GO and KEGG to analyze the effects of the two kinds of DEGs on OA. The "ggplot2" package was used to draw the visualization chart based on results of the analyses.

DEG network construction and hub gene screening

The search tool for the retrieval of interacting genes/ proteins (STRING version 11.5, https://string-db.org/), an online analysis tool (31), was utilized to present and evaluate the protein interaction (protein-protein interaction, PPI) network. The 27 DEGs selected in this study were imported into the STRING website, and the potential relationships among these DEGs were further explored using STRING analysis tools. The filter condition was set as follows: network type selected; "full-STRING network"; confidence \geq medium confidence (0.4). The calculation results from STRING were imported into Cytoscape version 3.8.2 (Cytoscape Consortium, San Diego, CA, USA) (32). The hub genes (33) in key positions in the PPI network were screened with a variety of algorithms through the cytoHubba plug-in of Cytoscape (34). The obtained hub genes were reintroduced into the STRING website to determine the PPI interaction (35).

Results

Results of the downloaded data

The analyzed data included 11 samples, including 6 OA cartilage tissues and 5 normal cartilage tissues. The fresh OA cartilage tissue was washed with aseptic phosphatebuffered saline and immediately frozen in liquid nitrogen. The samples were preserved at -140 °C until the extraction of RNA. Healthy cartilage tissue was acquired from the National Center for Disease Research and Exchange (National Disease Research Interchange, NDRI). Eigenvalues of the main components in each sample were analyzed through covariance matrix analysis (36), and 3-dimensional principal component analysis diagrams were drawn (*Figure 2*).

DEG screening results

A total of 31 DEGs from the OA and control groups were identified. After weight removal, 27 DEGs were retained (Table S1), including 18 and 9 upregulated and downregulated genes, respectively (*Table 1*). A volcanic map (*Figure 3*) was used to analyze and observe the $|\log_2(FC)|$ distribution of the DEGs showing changes of more than 2 times. The upregulated and downregulated genes were marked in red and blue, respectively. The genes with $|\log_2(FC)|$ values less than 2 or P<0.05 were categorized as indifferent genes and marked in gray. The DEGs in each sample were depicted by a heat map, which presented upregulated and downregulated genes as red and green, respectively (*Figure 4*).

Results of GO and KEGG analysis of the DEGs

The results of GO and KEGG signal pathway enrichment analysis of the DEGs using DAVID are shown in *Figure 5*. In the BP category, DEGs were mainly correlated with oxygen, nutrition, and corticosteroid levels. In the CC category, DEGs were mainly enriched in fibrous collagen trimers, banded collagen fibrils, and extracellular matrices containing collagen. In the MF category DEGs were mainly involved in extracellular matrix structural components for tensile strength and platelet-derived growth factor binding. GO enrichment showed that the upregulated DEGs were mainly correlated with hypoxic response and mitochondrial matrix and magnesium ion binding. In comparison, the downregulated DEGs were mainly correlated with the extracellular matrix structure. KEGG analysis demonstrated

	Table 1 Li	st of differentially expressed genes	
	Gene ID	Description	Log ₂ (FC)
	DDIT4	DNA damage inducible transcript 4	4.248
	TSC22D3	TSC22 domain family member 3	3.665
	TMOD1	Tropomodulin 1	3.632
	HILPDA	Hypoxia inducible lipid droplet associated	3.578
	CYP4B1	Cytochrome P450 family 4 subfamily B member 1	3.572
	PCK1	Phosphoenolpyruvate carboxykinase 1	3.511
	PDK4	Pyruvate dehydrogenase kinase 4	3.281
	PDE2A	Phosphodiesterase 2A	3.193
	ADM	Adrenomedullin	3.135
	GAB2	GRB2 associated binding protein 2	2.831
	C10orf10	Chromosome 10 open reading frame 10	2.828
	STC2	Stanniocalcin 2	2.811
	ACADL	Acyl-CoA dehydrogenase, long chain	2.762
	TXNIP	Thioredoxin interacting protein	2.736
	GLRX	Glutaredoxin	2.689
	GLUL	Glutamate-ammonia ligase	2.666
	SCNN1A	Sodium channel epithelial 1 alpha subunit	2.654
	GADD46B	Growth arrest and DNA damage inducible beta	2.626
	MXRA5	Matrix remodeling associated 5	-2.937
	EPHB2	EPH receptor B2	-2.977
	SERPINF1	Serpin family F member 1	-2.991
	POSTN	Periostin	-3.115
	BMP1	Bone morphogenetic protein 1	-3.206
	COL1A2	Collagen type I alpha 2 chain	-3.518
	MMP13	Matrix metallopeptidase 13	-3.578
	COL1A1	Collagen type I alpha 2 chain	-3.622
_	HLA-DRA	Major histocompatibility complex, class II, DR alpha	-3.83

Table 1 List of differentially expressed genes

FC, fold change.

that upregulated genes were mainly related to the peroxisome proliferator-activated receptor (PPAR) signal pathway, whereas downregulated genes were associated with protein digestion and absorption, extracellular matrix receptor interaction, and other signal pathways (*Figure 6*).

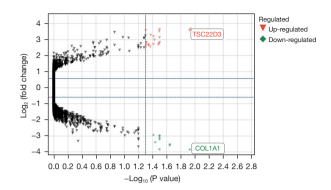


Figure 3 Volcano map of DEGs. The volcano plot for DEGs in OA and normal sample. Genes with adjusted $|\log(FC)| \ge 2$ and FDR <0.05 values were considered DEGs. Red dots indicate upregulated genes, green dots indicate downregulated genes, and black dots indicate genes without difference. DEGs, differentially expressed genes; OA, osteoarthritis; FC, fold change; FDR, false discovery rate.

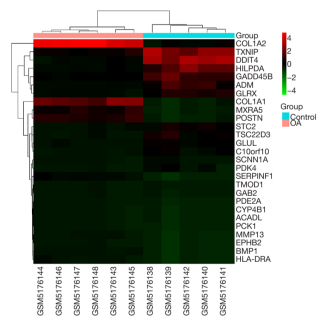


Figure 4 Heatmap of differentially expressed genes. Heatmap of DEGs of 11 samples in GSE169077. Red represents upregulated DEGs and green represents downregulated DEGs. OA, osteoarthritis; DEGs, differentially expressed genes.

Results of the PPI network analysis of the DEGs

STRING analysis demonstrated that the PPI network consisted of 27 nodes and 22 edges. The average degree of the nodes among the networks was 1.63. The P value was

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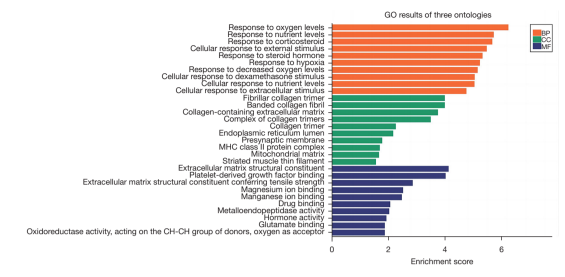


Figure 5 GO enrichment analysis of all DEGs. The abscissa represents the percentage of genes enriched in the functional area, and the ordinate represents different functional areas. The longer the line, the more significant P is. GO, Gene Ontology; DEGs, differentially expressed genes; CH-CH, Hypermethyl structure.

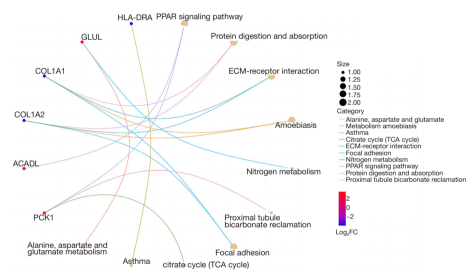


Figure 6 KEGG enrichment analysis of all DEGs. The abscissa represents the percentage of genes enriched in the functional area, and the ordinate represents different functional areas. The longer the line, the more significant P is. The circle size represents the number of enriched genes, the color represents P value. The whiter the color, the more significant P is. GO, Gene Ontology; KEGG, Kyoto Encyclopedia of Genes and Genomes; DEGs, differentially expressed genes; TCA, tricarboxylic acid.

6.33e-15, and the degrees for *COL1A2*, *COL1A1*, *BMP1*, *MXRA5*, and *SERPINF1* ranked the top 5. In order to better explore the pathogenesis of OA, 27 clustered DEGs were analyzed by k-means (*Figure 7*). These DEGs were visualized by hiding the non-interacting genes. Clusters were distinguished by 3 colors: the green group (14 genes)

was primarily involved in the glucocorticoid receptor pathway, the blue group (7 genes) was involved in the relaxin signaling pathway, and the red group (6 genes) was associated with the estrogen receptor pathway (*Table 2*).

The results of the PPI network were imported into Cytoscape, and the 27 DEGs were sequenced by the

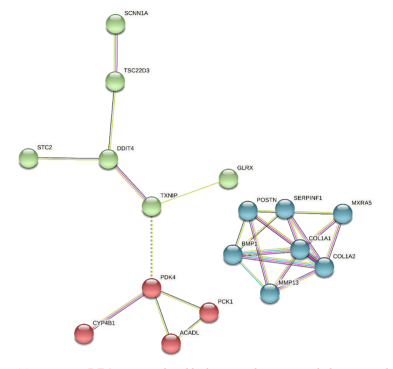


Figure 7 PPI clustering analysis. Twenty-seven DEGs were analyzed by k-means clustering, with disconnected nodes in the network hidden, green clustering (14 DEGs), blue clustering (7 DEGs), and red clustering (6 DEGs). PPI, protein-protein interaction; DEGs, differentially expressed genes.

cytoHubba plug-in according to various calculation methods. Ten candidate hub genes were selected, and 5 hub genes, namely, *COL1A1*, *COL1A2*, *POSTN*, *BMP1*, and *MMP13*, were selected after combining 6 calculation methods (MCC, DMNC, degree, closeness, clustering coefficient, and EPC) (*Table 3*). The 5 genes were analyzed by PPI, with a total of 5 nodes and 10 interaction networks on the left side. We performed heatmap analysis on the 5 hub genes and found significant differences between the OA and normal groups (*Figure 8*). The average degree of the nodes among the networks was 4. The P value was 3.83e-13, which was annotated by GO and KEGG in different colors (*Figure 9*). In the cluster grouping, the genes contained were the hub genes in the blue group (*Table 2*).

Discussion

In 2015, the Osteoarthritis Research Society International (OARSI) proposed a new definition of arthritis (37) to refer to cellular stress and extracellular matrix degeneration caused by microscopic and macroscopic damage. These involve preinflammatory pathways of innate immunity, which is primarily a molecular disorder, followed by anatomical and/or physiological disorders, including cartilage degeneration, bone remodeling, and osteophyte formation. Therefore, it is particularly important to understand the physiological and pathological changes of OA at molecular level.

In this study, we focused on the cartilage of OA, selected the GSE169077 dataset in the GEO database, and obtained 27 DEGs using R software. We used STRING to cluster the 27 DEGs, of which the blue cluster group contained all of the hub genes involved in the relaxin signaling pathway. Relaxin is a peptide hormone discovered by Frederick Hisaw in 1926 in the study of changes in the pelvic canal during pregnancy. It is mainly produced in the ovarian corpus luteum of pregnant mammals and the male prostate. As the research progressed, researchers found relaxin in the heart, liver, lungs, and kidneys was closely related to various functions and diseases, such as regulation of the uterus during pregnancy, lowering blood pressure and vascular resistance, and anti-pulmonary fibrosis and liver fibrosis. Clifton's study of thumb arthritis found that matrix metalloproteinases (MMPs) could act as a potential

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	Table 2 Dices and pathways involved in 5 clusters								
Clusters	Genes	P value	Pathway						
Green	ADM, C10orf10, DDIT4, EPHB2, GAB2, GADD45B, GLRX, GLUL, HILPDA, PDE2A, SCNN1A, STC2, TSC22D3, TXNIP	3.06×10 ⁻⁴	Glucocorticoid receptor pathway						
Blue	BMP1, COL1A1, COL1A2, MMP13, MXRA5, POSTN, SERPINF1	<10 ⁻¹⁶	Relaxin signaling pathway						
Red	ACADL, CYP4B1, HLA-DRA, PCK1, PDK4, TMOD1	2.59×10 ⁻⁵	Estrogen receptor pathway						

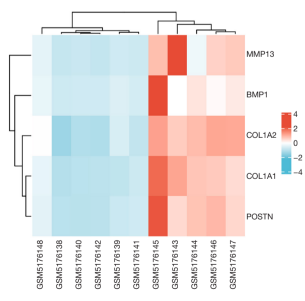
Table 2 DEGs and pathways involved in 3 clusters

DEGs, differentially expressed genes.

Table 3 Comparison of hub genes by using CytoHubba plug-in methods

Rank	MCC	DMNC	Degree	Closeness	Clustering coefficient	EPC
1	COL1A1*	BMP1	COL1A1*	COL1A1*	MMP13	COL1A2*
2	COL1A2*	MMP13	COL1A2*	COL1A2*	MXRA5	COL1A1*
3	POSTN*	MXRA5	POSTN*	POSTN*	SERPINF1	POSTN*
4	BMP1*	SERPINF1	BMP1*	BMP1*	PCK1	BMP1*
5	MMP13*	POSTN	MMP13*	MMP13*	ACADL	MMP13*
6	MXRA5	COL1A1	MXRA5	MXRA5	PDK4	MXRA5
7	SERPINF1	COL1A2	SERPINF1	SERPINF1	BMP1	SERPINF1
8	PCK1	PCK1	PCK1	PCK1	POSTN	PCK1
9	ACADL	ACADL	ACADL	ACADL	COL1A2	ACADL
10	PDK4	PDK4	PDK4	PDK4	COL1A1	PDK4

*, the gene ID was screened for hub genes in this study. MCC, Maximal Clique Centrality; DMNC, Density of Maximum Neighborhood Component; EPC, Edge Percolated Component.



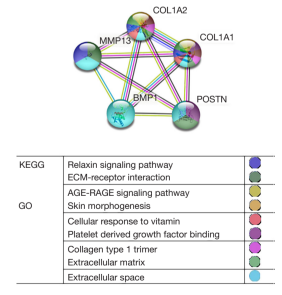


Figure 8 Hub gene heatmap. The heatmap of 5 hub genes showing high expression in the control group and low expression in OA group. OA, osteoarthritis.

Figure 9 PPI analysis of hub genes. KEGG, Kyoto Encyclopedia of Genes and Genomes; GO, Gene Ontology; PPI, protein-protein interaction.

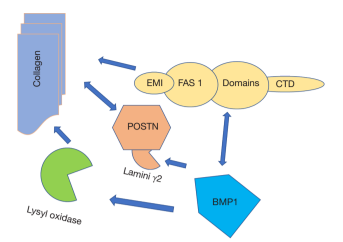


Figure 10 Schematic diagram of POSTN-BMP1-LOX axis. Schematic of the POSTN-BMP1-LOX axis as the basis for the collagen matrix chemical property. EMI, elastin microfibril interfacer; CTD, connect the dots; POSTN, periostin; BMP, bone morphogenetic protein.

target for relaxin, which in turn affects joint stability (38). In addition, Ko *et al.* showed that relaxin antagonizes synovial fibroblasts in OA, thereby preventing the flexion contracture of OA (39). The role of the relaxin signaling pathway in the development of OA cartilage should be further explored.

The upregulated genes involved in BP were mainly concerned with hypoxic response, while the genes involved in CC were correlated with mitochondrial matrix enrichment. The identified DEGs were not concentrated in critical pathways, but GO identified 3 genes involved in the extracellular matrix, while KEGG identified genes involved in protein digestion and absorption. Downregulated DEGs were analyzed by cytoHubba and utilized as the hub genes in this study (*Figures 5,6*).

Among the 5 hub genes, *COL1A1* and *COL1A2* are major collagen producers and important regulatory genes for bone, skin, and other connective tissues. Fang *et al.* suggested that they may be important biomarkers for hydroxymethylation and could be used as target genes for accurate diagnosis and treatment of OA (40). Another study proposed that increased secretion of type I collagen encoded by homologous *COL1A1* and *COL1A2* trimers could lead to subchondral bone stenosis and collagen fiber disorganization in OA (41).

MMPs are zinc-dependent extracellular proteases, the main function of which is to degrade extracellular

interstitial components (42). Recent studies have found that *MMP13* prevents articular cartilage damage through binding to metalloproteinase tissue inhibitor (43), which may be a potential treatment target for OA. In addition, Wang *et al.* found that an *MMP13* inhibitor could reduce the severity of meniscus and ligament injury in an induced OA mice model (44).

Periostin (*POSTN*) is a protein-coding gene functioning as a transforming growth factor β -induced extracellular matrix protein. It is also known as osteoblast-specific factor 2 and is expressed in articular joint components such as cartilage, subchondral bone, meniscus, and ligaments (45). *POSTN* can be crosslinked with other extracellular matrix proteins, such as type I collagen and fibronectin. The study has found that mice with *POSTN* deletion exhibit collagen crosslinking defects and a reduced resistance to mechanical stress (46). In the study by Tajika *et al.*, *POSTN* did not upregulate collagen-related genes such as *COL1* and *COL2*. It is speculated that *POSTN* might indirectly affect collagen-related genes, for example, *COL1* and *COL2*, through regulating the expression of *MMP13* (47).

In this study, bone morphogenetic protein 1 (BMP1) was a pivotal gene, which plays a variety of roles in regulating extracellular matrix formation. BMP1 preprocesses functional structural proteins in the extracellular matrix of active enzymes, binds with collagen I and II, and regulates the size and shape of heteromorphic fibers. Studies have suggested direct interaction between Fas1 domains, and that BMP1 is indirectly related to the EMI domain of POSTN (48,49). Targeted deletion of the POSTN gene in mice led to reduced collagen crosslinking (46,50). Kii et al. reported that the POSTN-BMP 1-LOX axis was the basis of the mechanochemical properties of the collagen matrix (51) (Figure 10). In recent studies, POSTN was found to determine the incidence of OA in women and predicted the occurrence of posttraumatic OA (52). Our interesting finding is that POSTN, BMP1, COL1A1 and COL1A2 are all closely related to collagen synthesis. This suggests that our drug development for osteoarthritis should focus on intervening in the intermediate steps that promote collagen synthesis (Figure 10).

Pain is a common phenotype in osteoarthritis, and extracellular matrix (ECG) receptor interaction is significantly different in the KEGG pathway. The proteoglycan and hyaluronic acid content of the ECM has been found to be elevated in the inflammatory progression of many diseases and, due to its unique reticular fibrous structure, is thought to be a "landing zone" for inflammatory cells. "This "net"

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influences their adhesion, retention, migration and activation. The process of transfer and localisation of leukocytes directly determines the duration of the inflammatory response. Nanus *et al.* (53) studies of synovial tissue from multiple painful knee lesions found that the most differentially expressed RNA was in the inflammatory signalling, which is consistent with our findings in this study.

Our study had some limitations. Firstly, we used only 1 dataset, and there may have been bias in the search for markers for OA. Secondly, the effect of OA on cartilage metabolism is only 1 factor, and systemic inflammatory response and diabetes, among others, can affect cartilage metabolism and integrity and possibly have an effect on results. Finally, we did not carry out experiments to prove the research results, which will be considered in future work.

Conclusions

This study applied bioinformatics to analyze the RNA expression profiles of healthy and OA groups. A total of 27 DEGs were identified, and their biological functions mainly contributed to the structural components and tensile strength of the extracellular matrix. KEGG analysis demonstrated that these DEGs were mainly related to protein digestion and absorption, extracellular matrix receptor interaction, and the PPAR signaling pathway, as well as other signaling pathways. *COL1A1, COL1A2, POSTN, BMP1*, and *MMP13* were identified as possible diagnostic and therapeutic biomarker targets for treating OA.

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Footnote

Reporting Checklist: The authors have completed the STREGA reporting checklist. Available at https://atm. amegroups.com/article/view/10.21037/atm-22-6450/rc

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://atm. amegroups.com/article/view/10.21037/atm-22-6450/coif). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related

to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

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Supplementary

Table S1 Expression of the 27 DEGs

Gene	GSM5176138	GSM5176139	GSM5176140	GSM5176141	GSM5176142	GSM5176143	GSM5176144	GSM5176145	GSM5176146	GSM5176147	GSM5176148
GLUL	4815.89	8961.6	4771.11	3917.58	3414.55	773.357	726.325	1414.48	1135.02	682.857	334.208
TXNIP	26144.7	26927.833	23504.5	19784.6	15690.85	3555.8767	3216.5667	7965.79	3372.6233	2482.59	1769.2533
SERPINF1	70.3827	218.059	132.11	295.933	159.891	1220.89	1865.24	1727.81	1089.75	1576.28	564.427
COL1A1	215.8081	1104.8035	622.21405	2108.323	702.6133	13834.6	11261.306	18343.025	10902.22	9487.358	4533.4605
DDIT4	24361.5	26023.4	25514.7	23399.1	25720.1	2249.78	853.864	2867.36	1730.64	835.125	648.993
ADM	5091.21	20593.8	11575.9	4858.08	10339.9	1771.99	1029.84	1973.33	933.392	837.948	426.589
STC2	2314.27	11905.1	7061.59	4315.5	5252.31	774.364	587.577	2063.06	767.694	591.168	464.117
SCNN1A	2543.33	6019.74	1491.02	1532.27	1401.74	234.623	446.847	394.69	420.38	450.076	219.205
TMOD1	1483.67	2902.04	988.447	1001.83	807.093	60.3715	50.0074	480.516	217.439	32.3687	118.704
GAB2	1315.04	2340.04	923.625	651.105	910.564	156.726	118.464	173.144	283.234	122.875	128.734
PDE2A	291.959	1261.31	305.572	355.433	382.947	61.9933	45.81	146.645	48.6928	43.6849	12.8315
BMP1	32.79	129.568	42.7577	69.7185	45.0776	385.783	564.231	1813.03	425.034	527.358	224.6
MMP13	16.7212	49.7429	31.5722	39.589	10.5638	784.805	154.588	412.165	354.511	380.653	120.794
PDK4	2188.57	8749.01	3061.83	559.65	1145.72	122.264	243.165	413.003	315.515	153.34	155.047
ACADL	311.678	630.498	305.028	189.446	291.387	36.7084	39.5068	35.2594	56.7651	83.6305	44.7138
GLRX	4694.27	13400.2	8032.58	7650.55	8174.08	1355.72	1012.09	1426.9	1734.5	1382.42	743.517
TSC22D3	6437.04	14477.8	5121.73	4893.21	3088.49	464.551	492.259	1014.79	303.818	347.237	417.811
GADD45B	12376.15	25145.3	10491.2	8670.52	10491.58	2524.215	2043.93	3430.605	1664.145	1970.75	1190.6
PCK1	152.723	495.238	398.577	156.349	334.59	14.6904	129.978	30.7288	17.7914	24.164	7.83968
C10orf10	4175.56	9243.23	3308	3679.81	1819.55	562.322	355.981	1468.49	654.828	460.769	290.02
EPHB2	17.6101	56.5981	18.7422	11.4576	24.6747	171.264	203.062	372.663	120.072	130.704	136.957
MXRA5	190.264	1033.79	448.728	774.694	664.567	3583.34	3503.33	9229.21	3438.93	5681.56	2183.4
CYP4B1	406.764	777.005	357.503	123.843	301.312	14.1113	16.1744	23.4173	233.744	43.4878	9.1974
POSTN	396.633	1009.28	336.202	1243.56	360.08	4727.76	5541.58	9553.36	6016.17	4788.15	2013.35
HLA-DRA	18.6538	75.7098	47.0843	31.0232	11.5365	944.596	1047.97	503.655	678.368	153.058	113.394
HILPDA	8427.02	21290.4	17523.1	14021.9	21170.6	2227.95	958.867	2869.76	1288.49	1339.32	474.435
COL1A2	961.867	8876.98	3672.9	5752.245	3398.7525	21020.5	23202.15	26547.35	25810.75	25568.4	14591.795

DEGs, differentially expressed genes.