Spatial regulation of gene expression during growth of articular cartilage in juvenile mice

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BACKGROUND: In juvenile mammals, the epiphyses of long bones grow by chondrogenesis within the articular cartilage. A better understanding of the molecular mechanisms that regulate the growth of articular cartilage may give insight into the antecedents of joint disease, such as osteoarthritis.

METHODS: We used laser capture microdissection to isolate chondrocytes from the superficial, middle, and deep zones of growing tibial articular cartilage in the 1-wk-old mouse and then investigated expression patterns by microarray. To identify molecular markers for each zone of the growing articular cartilage, we found genes showing zone-specific expression and confirmed by real-time PCR and *in situ* hybridization.

RESULTS: Bioinformatic analyses implicated ephrin receptor signaling, Wht signaling, and bone morphogenetic protein signaling in the spatial regulation of chondrocyte differentiation during growth. Molecular markers were identified for superficial (e.g., Cilp, Prg4), middle (Cxcl14, Tnn), and deep zones (Sfrp5, Frzb). Comparison between juvenile articular and growth plate cartilage revealed that the superficial-to-deep zone transition showed similarity with the hypertrophic-to-resting zone transition.

CONCLUSION: Laser capture microdissection combined with microarray analysis identified novel signaling pathways that are spatially regulated in growing mouse articular cartilage and revealed similarities between the molecular architecture of the growing articular cartilage and that of growth plate cartilage.

Skeletal growth involves endochondral bone formation at the growth plate, which causes elongation of the metaphyses, intramembranous bone formation at the periosteal surface, which causes widening of the diaphysis, and chondrogenesis at the articular surface, which causes expansion of the epiphyses. This last process has received relatively little research attention but has important clinical implications because the articular cartilage is susceptible to degenerative diseases such as osteoarthritis due to its limited capacity for regeneration and repair (1). A better understanding of the molecular mechanisms that regulate the growth and maintenance of the articular cartilage may therefore facilitate the development of new therapeutic measures to prevent destruction and/or to achieve regeneration of articular cartilage.

Articular cartilage, like growth plate cartilage, is composed of chondrocytes surrounded by extracellular matrix and is spatially organized into three histologically distinct zones. The superficial zone (SZ) lies closest to the joint surface and consists of densely packed chondrocytes that are elongated parallel to the articular surface. In the middle zone (MZ), chondrocytes are scattered and more rounded in shape (MZ). In the deep zone (DZ), the chondrocytes are larger in size and arranged in columns perpendicular to the articular surface. These three zones differ not only in cellular morphology but also in their cellular metabolism (2) and extracellular matrix composition (3).

While cell proliferation occurs sparingly in adult articular cartilage, it is much more common in juvenile articular cartilage. Growth of the articular cartilage reportedly occurs appositionally from the articular surface (4), suggesting that the SZ may contain actively dividing chondrocytes, or a progenitor cell population that is capable of generating new chondrocytes (5). The mechanisms responsible for the distinctive properties of SZ are not well understood, but chondrocytes isolated from different zones of the articular cartilage, when cultured in monolayer, show substantially different response to growth factors, including insulin-like growth factor I, fibroblast growth factors (FGF), and transforming growth factor- β (6,7).

Thus, much remains unknown about the fundamental biological differences among the zones of articular cartilage and the mechanisms that initiate and maintain these differences. In the current study, we sought to elucidate the regulation of gene expression across different zones of the articular cartilage. We used laser capture microdissection (LCM) to isolate SZ, MZ, and DZ chondrocytes from juvenile mouse articular cartilage and compared gene expression across the three zones using microarray analysis. We then used bioinformatic approaches to identify spatially regulated signaling pathways that are

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Gene expression in growing cartilage



likely to direct the zonal differences in the articular cartilage. Finally, we compared spatial regulation in articular cartilage and growth plate cartilage, which also contains three zones: resting zone (RZ), proliferative zone (PZ), and hypertrophic zone (HZ).

RESULTS

Spatial Analysis of Articular Cartilage

Articular cartilage from the proximal tibias of 1-wk-old mice was microdissected into the SZ, MZ, and DZ using LCM (**Figure 1a,b**). Microarray analysis was then used to identify spatial differences in gene expression. From SZ to MZ, expression of 47 genes was upregulated and expression of 142 genes was downregulated (\geq twofold; false discovery rate (FDR) < 0.05; **Supplementary Table S1** online). Ingenuity pathway analysis identified multiple signaling pathways that were overrepresented in these gene sets, including ephrin receptor signaling in the upregulated gene set and basal cell carcinoma signaling (which predominantly involves hedgehog signaling) in the downregulated gene set (**Supplementary Table S1** online). This latter finding is consistent with our previous work showing that hedgehog signaling is active primarily on the articular surface (8). From MZ to DZ, 48 genes were upregulated and 161 genes were downregulated (≥twofold; FDR < 0.05; **Supplementary Table S1** online). Pathway analysis showed that insulin-like growth factor I signaling was strongly overrepresented among downregulated genes (**Supplementary Table S1** online). Comparison of SZ and DZ identified substantially more differentially expressed genes: 543 genes upregulated and 1,417 downregulated (**Supplementary Table S1**



Figure 1. Microarray analysis of laser capture microdissected articular cartilage. (**a**) Masson trichrome-stained section of formalin-fixed, decalcified 1-wk-old mouse proximal tibial cartilage showing the location of zones in the articular cartilage that were targeted by laser capture microdissection (LCM). (**b**) Hemotoxylin and eosin-stained frozen sections of 1-wk-old mouse proximal tibial cartilage after LCM. (**c**) Principle component analysis of microarray data from SZ, MZ, and DZ articular cartilage. (**d**) Heat map showing hierarchical clustering of spatially regulated genes (defined by false discovery rate < 0.05 SZ vs. DZ). Red color represents upregulation, and green color represents downregulation. Schematic line graphs on the right represent the typical pattern of changes in gene expression across the three zones, and examples of genes in each cluster are listed. DZ, deep zone; MZ, middle zone; SZ, superficial zone. Bar = 100 µm.

online). Pathway analysis implicated protein kinase A signaling and Wnt signaling among upregulated genes and endocytosis signaling among downregulated genes (**Supplementary Table S1** online).

Principal component analysis suggested that MZ has a transitional expression profile between those of the SZ and DZ (Figure 1c). Similarly, hierarchical clustering of genes that showed a significant change across zones (FDR < 0.05) demonstrated that the majority of genes exhibited a progressive upregulation or downregulation across the three zones (Figure 1d), suggesting that the MZ primarily represents a transition between the SZ and DZ, rather than a functionally distinct region.

Wnt and BMP Signaling in Articular Cartilage

The bioinformatic analysis implicated Wnt signaling in the spatial regulation of articular cartilage (Supplementary Table S1 online). Interestingly, the expression data did not suggest a simple pattern of either increasing or decreasing Wnt signaling across the zones but a more complex pattern (Figure 2 and Supplementary Table S2 online). For example, Wnt5b was upregulated from SZ to DZ, while Wnt9a showed the reverse trend. Wif1, which encodes an inhibitor of Wnt signaling, was downregulated >80-fold from SZ to DZ, while four other soluble modulators/inhibitors of Wnt signaling, Sfrp1, 2, 5, and Frzb, were upregulated in the DZ. Receptors of Wnt signaling appeared to be more abundant on the articular surface, with downregulation of *Fzd4*, 5, and 8 from the SZ to DZ. Expression of Lef1, a transcriptional activator involved in Wnt signaling (9), was upregulated in the DZ. Consistently, β -catenin, which interacts with Lef1 and is critical for canonical Wnt signaling, appeared to be increased by immunohistochemistry at the peripheral part of the DZ (Supplementary Figure S1 online), suggesting cytoplasmic accumulation of β -catenin. However, there was not clear evidence of β -catenin nuclear translocation, which is needed for activation of the canonical Wnt signaling pathway.



Figure 2. Wnt/ β -catenin and TGF- β /Bmp signaling in articular cartilage. Schematic diagram summarizing genes involved in Wnt signaling and BMP signaling that were significantly regulated across different zones of articular cartilage in 1-wk-old mouse. Genes listed in each zone show substantially greater expression in that zone than in other zones. Fold changes and *P* values are listed in **Supplementary Table S3** online. Bmp, bone morphogenetic protein; DZ, deep zone; MZ, middle zone; SZ, superficial zone; TGF, transforming growth factor. In addition, multiple genes involved in bone morphogenetic protein (Bmp) signaling were differentially expressed in different zones of the articular cartilage. For example, several Bmps (*Bmp2*, 4, 5, 6, 7, and *Gdf10*) and their receptors (*Bmpr1a*, *Bmpr2*, and *Acvr1*) were significantly downregulated from SZ to DZ. Consistently, *Grem1*, which encodes a Bmp inhibitor, was upregulated in DZ (**Figure 2** and **Supplementary Table S3 online**). Thus, the findings suggest that there exists a BMP signaling gradient across the articular cartilage, with greater signaling in SZ than that in DZ.

Identification of Spatial Markers for Articular Cartilage

Next, we sought to identify useful molecular markers for each zone of the articular cartilage. We used an empirical formula (10) that gave high rank to genes that were expressed at high levels in a particular zone and at low levels in the other two zones. For SZ, the highest ranking marker was cartilage intermediate layer protein (Cilp, also known as Cilp-1) (expression in SZ 36-fold > MZ 42-fold > vs. DZ; Figure 3a and Supplementary Table S4 online). Other SZ markers included Fgf18, Prg4, Dkk3, Bmp2, and Bmp6. Only two genes, Tnn and Cxcl14, met the criterion for MZ markers, and these showed only moderate zonal specificity (Figure 3a and Supplementary Table S4 online) In DZ, three of the highest ranking markers were previously reported as RZ markers in the growth plate (10), including *Pcp4*, *Efemp1*, and *Sfrp5* (Figure 3a and Supplementary Table S4 online), suggesting that DZ shares molecular similarities with RZ. Sfrp5 and another DZ marker, frizzled-related protein (Frzb), are both secreted frizzled-related proteins that antagonize Wnt signaling (11,12). These findings are in agreement with previous work showing that active Wnt signaling is important for SZ chondrocyte proliferation and Prg4 expression (13).

The expression of specific markers in SZ, MZ, and DZ were evaluated using real-time PCR (**Figure 3b**), and their expression in whole articular cartilage and whole growth plate cartilage was also compared (**Figure 3c**). All six SZ markers showed zone-specific expression not just within the articular cartilage but more generally, in that expression levels were lower in growth plate than that in articular cartilage. In contrast, *Tnn*, *Sfrp5*, and *Frzb* showed similar expression levels in growth plate and articular cartilage (**Figure 3c**).

To further confirm our findings, the localization of several spatial markers in 1-wk-old mouse proximal tibia and distal femur was studied by *in situ* hybridization (Figure 4). *Cilp* expression was present primarily in the one- or two-cell layer closest to the articular surface, whereas signals for *Prg4* were present more broadly in the SZ and part of the immediately adjacent MZ. *Cxcl14* was preferentially detected in the MZ of the articular cartilage and also in HZ of the growth plate. *Frzb* was confirmed to be present specifically in DZ.

Relationship Between Spatial Regulation of Articular Cartilage and Growth Plate Cartilage

We hypothesized that the spatial regulation of gene expression in articular cartilage might be similar to growth plate cartilage, Gene expression in growing cartilage

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Figure 3. Differential gene expression (mean \pm SEM) of potential spatial markers across the tibial articular cartilage of 1-wk-old mice. Different zones were isolated by laser capture microdissection, and expression of potential spatial markers (yellow bars, superficial zone markers; purple bars, middle zone markers; green bars, deep zone markers) were analyzed by (**a**) microarray and (**b**) real-time PCR. *P* values in **b** represent pairwise comparison with Holm–Sidak correction following one-way ANOVA. To compare gene expression between overall articular cartilage and growth plate cartilage, (**c**) manual microdissection of both regions from proximal tibias of 1-wk-old mice was performed followed by real-time PCR. Statistical significance in **c** was calculated by *t*-test, articular cartilage vs. growth plate cartilage. **P* < 0.05; ***P* < 0.001. AC, articular cartilage; D, deep zone; GP, growth plate cartilage; M, middle zone; S, superficial zone.

with the SZ corresponding to the RZ and the DZ corresponding to the HZ. This hypothesis was based on two prior observations. First, SZ and RZ reportedly contain progenitor cell populations (5,14). Second, chondrocytes gradually hypertrophy (enlarge) with the spatial transition from SZ to DZ and from PZ to HZ. To test this hypothesis, we combined the current articular cartilage analysis with our previous spatial analysis in 1-wk-old rat growth plate (10). Gene-set enrichment analysis

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Figure 4. *In situ* hybridization of zonal markers of articular cartilage. Formalin fixed, decalcified sections of 1-wk-old mouse tibial cartilage were hybridized to DIG-labeled riboprobes, producing a purple color and were visualized by scanning the slides with a ScanScope CS digital scanner under bright field microscopy. Left panel, Masson-trichrome–stained sections; middle panel, *in situ* hybridization without counterstaining; right panel, high magnification views taken from within the rectangular area indicated in the corresponding middle panel. DZ, deep zone; HZ, hypertrophic zone; MZ, middle zone; PZ, proliferative zone; RZ, resting zone; SZ, superficial zone. Bar = 100 µm.

(**Supplementary Figure S2** online) showed that the genes most downregulated from SZ to DZ (greatest fold change; FDR < 0.05) are enriched with genes upregulated from RZ to HZ (P = 0.031), and conversely, genes upregulated from SZ to DZ are enriched with genes downregulated from RZ to HZ (P < 0.001). Similarly, among the 200 most strongly upregulated and 200 most strongly downregulated genes from SZ to DZ (greatest fold change; FDR < 0.05), there was a strong negative correlation between changes from SZ to DZ and changes from RZ to HZ (r = -0.4031; P < 0.0001, Pearson's correlation; **Table 1** and **Figure 5a**). This correlation was in large part attributable to the changes from PZ to HZ (r = -0.4595; P < 0.0001). Similarly, Changes from PZ to HZ showed a strong negative correlation with changes from MZ to DZ (r = -0.4567; P < 0.0001; **Table 1**), and, more weakly, with changes from SZ to MZ (r = -0.2489; P = 0.0004; **Table 1**).

Table 1.	orrelation between spatial differences in gene expression			
of articular and growth plate cartilage				

Communication	Pearson's	Quality	
Comparison	<i>r</i> value	<i>P</i> value	
400 genes that differ most from SZ to DZ			
SZ to DZ vs. RZ to PZ	0.0409	0.4149	
SZ to DZ vs. PZ to HZ	-0.4595	< 0.0001	
SZ to DZ vs. RZ to HZ	-0.4031	< 0.0001	
200 genes that differ most from SZ to MZ			
SZ to MZ vs. RZ to PZ	0.0251	0.7242	
SZ to MZ vs. PZ to HZ	-0.2489	0.0004	
SZ to MZ vs. RZ to HZ	-0.2121	0.0026	
200 genes that differ most from MZ to DZ			
MZ to DZ vs. RZ to PZ	-0.0552	0.4376	
MZ to DZ vs. PZ to HZ	-0.4567	<0.0001	
MZ to DZ vs. RZ to HZ	-0.4399	<0.0001	

DZ, deep zone; HZ, hypertrophic zone; MZ, middle zone; PZ, proliferative zone;

RZ, resting zone; SZ, superficial zone.

Additionally, we found a significant overlap between genes that were downregulated from SZ to DZ (\geq twofold; FDR < 0.05) and genes upregulated from RZ to HZ (\geq twofold; FDR < 0.05), and similarly, a significant overlap between genes that were upregulated from SZ to DZ and genes downregulated from RZ to HZ (P < 0.001, Pearson's χ^2 ; **Figure 5b,c**). Thus, multiple analyses suggest that SZ to DZ expression pattern is not, as we had hypothesized, similar to the RZ to HZ pattern. Instead, the correspondence is inverted so that the DZ to SZ expression pattern is similar to that of RZ to HZ.

Ingenuity pathway analysis identified multiple functional pathways that were overrepresented both among genes upregulated in SZ (vs. DZ) and also in HZ (vs. RZ), including ephrin receptor signaling, integrin signaling, and transforming growth factor- β signaling (Figure 5d). Ingenuity pathway analysis also identified functional pathways that were overrepresented both among genes commonly upregulated in the DZ and the RZ, including glutamate receptor signaling and Wnt signaling (Figure 5d).

DISCUSSION

Using LCM and microarray analysis, we analyzed the spatial expression patterns of the articular cartilage in 1-wk-old mice. Bioinformatic analysis of these expression patterns implicated several signaling pathways that were spatially regulated across different zones of the articular cartilage, including Wnt/ β -catenin signaling, which has been previously shown to play an important role in cartilage development (15). Our pathway analysis implicated Wnt/ β -catenin signaling in the spatial regulation of articular cartilage (**Supplementary Table S1** online) with a complex pattern. For example, DZ expressed a high level of Wnt5b, which promotes chondrogenesis and inhibits hypertrophy (16). However, the level of several soluble modulators/inhibitors of Wnt signaling, including *Sfrp1*, *2*, *5*, and *Frzb*, was also higher in DZ. Translocation of β -catenin to the

nucleus, which is indicative of canonical Wnt signaling pathway activation, was not definitively demonstrated in the DZ by immunohistochemistry. Therefore, it remains unclear whether Wnt signaling is more active in the deeper regions of articular cartilage.

Another signaling pathway that is critical for cartilage development and function is transforming growth factor- β /BMPs signaling. Postnatal deletion of *Bmpr1a* (a BMPs receptor) in mice leads to accelerated degeneration of the articular cartilage (17), while a single nucleotide polymorphism (rs12593365) of *GREM1* (a BMPs inhibitor) in humans is associated with 20% reduced risk of radiographic hip osteoarthritis (18), suggesting that BMPs contribute to articular cartilage maintenance. Our microarray analysis showed that *Bmp2*, 4, 6, 7, and *Gdf10* (Bmp3b) were expressed at higher levels in the SZ, while the inhibitor *Grem1* was expressed in the DZ, suggesting greater BMPs signaling at the articular surface, where it may serve to promote expression of Prg4 (19,20).

Our bioinformatic analysis also identified several other pathways that show differential expression patterns in the zones of the articular cartilage including integrin and ephrin receptor signaling. Integrins appear to play an important role in the interaction between cartilage extracellular matrix and the growth and differentiation of articular chondrocytes (21). Ephrin receptor signaling has been reported to be involved in cartilage morphogenesis of the limb bud in the chicken (22), but little is known about its role in articular cartilage.

We also identified mRNA markers for each zone of articular cartilage using microarray, real-time PCR, and *in situ* hybridization. We identified several novel markers for SZ, including *Cytl1, Errfi1*, and *Fgf18*, and confirmed previously reported markers *Prg4* (ref. 23) and *Dkk3* (ref. 24). We also found very specific expression of *Cilp* in the outermost one- to two-cell layers of the SZ. Although *Cilp* is known as cartilage intermediate layer protein and was initially thought to be specifically expressed in MZ (25–27), others have demonstrated *Cilp* primarily in SZ (28). Only two markers were identified for the MZ, both of which showed only moderate specificity, consistent with a transitional nature of the MZ. For the DZ, we identified multiple genetic markers, such as *Sfrp5* and *Efemp1*, which we previously identified as RZ markers in the growth plate (10).

We also compared the spatial regulation between the articular cartilage and growth plate cartilage (10). Contrary to our hypothesis, the analysis suggested that the SZ shows an expression pattern similar to the HZ, and the DZ shows an expression pattern similar to the RZ. Part of this relationship could be attributable to the similarities between the DZ and RZ, which both arise from a common pool of round epiphyseal chondrocytes in the embryo which then become separated by the secondary ossification center. However, there were even more genes commonly expressed in the SZ and HZ (158 genes) than that in RZ and DZ (72 genes). Furthermore, bioinformatic analysis suggested that various signaling pathways, such as ephrin signaling, integrin signaling, and transforming growth factor- β signaling, were implicated by genes commonly upregulated

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Figure 5. Comparison of spatial regulation of gene expression in articular and growth plate cartilage. Our current zonal analysis of 1-wk-old tibial mouse articular cartilage was combined with our previous zonal analysis of 1-wk-old tibial rat growth plate cartilage (10). (a) Heat maps were constructed for the 200 genes that were most strongly upregulated and 200 genes that were most strongly downregulated from SZ to DZ of articular cartilage. Red color represents upregulation, and green color represents downregulation. Genes downregulated from SZ to DZ (green) tended to be upregulated from RZ to HZ (red) and vice versa. (b) Venn diagrams showing the overlap between RZ-to-HZ upregulation and SZ-to-DZ downregulation, and also the overlap between RZ-to-HZ downregulation and SZ-to-DZ upregulation. Numbers on Venn diagram indicate number of genes in each category. Significant regulation was defined as \geq twofold and FDR < 0.05. (c) Observed (black bars) and expected (gray bars) overlap of RZ-to-HZ regulation in growth plate and SZ-to-DZ regulation in articular cartilage. Asterisks indicate observed overlap greater than expected by chance (*P* < 0.001, Pearson's χ^2 test). (d) Ingenuity pathway analysis of the 158 genes that were both significantly upregulated from RZ to HZ and also downregulated from SZ to DZ genes ("SZ/HZ genes") and the 72 genes that were both significantly upregulated from RZ to HZ and also upregulated from SZ to DZ ("DZ/RZ genes"). For SZ/HZ genes, the top 10 biological functions with the lowest *P* value were listed. For DZ/RZ genes, all functions with *P* < 0.05 were listed. DZ, deep zone; FDR, false discovery rate; HZ, hypertrophic zone; MZ, middle zone; PZ, proliferative zone; RZ, resting zone; SZ, superficial zone.

in SZ and HZ (compared with DZ and RZ, respectively). For example, we found evidence for a BMP signaling gradient across the articular cartilage, with greater signaling in the SZ than in DZ, which corresponds to a BMP signaling gradient previously suggested across growth plate cartilage, with greater BMP action in HZ than in RZ. The results of the comparison are consistent with our previous study, which involved a less precise manual dissection of articular cartilage (29).

There have been other studies that used LCM and microarray analysis to investigate the zonal gene expression of the mammalian articular cartilage. For example, Fukui *et al.* (2,30) compared gene expression in the three zones of normal and osteoarthritic human articular cartilage. However, their study involved elderly subjects, whereas our study focused on spatial regulation in growing, early postnatal articular cartilage. Amanatullah *et al.* (31) studied gene expression in 1–3-mo-old bovine articular cartilage and obtained results somewhat similar to ours, including high *Prg4* and *Cilp* expression in SZ. In contrast, the genes that we found to be specifically expressed in the DZ, such as *Frzb*, *Sfrp5*, and *Efemp1*, were reported by Amanatullah *et al.* to be expressed in the MZ, perhaps because of differences in defining the articular zones.

In summary, we used LCM to isolate different zones of the articular cartilage and used microarray to analyze global gene expression. Bioinformatic analysis corroborated previously reported signaling pathways, such as Wnt and Bmp signaling, and implicated novel pathways, such as ephrin and integrin signaling, for spatially associated articular chondrocyte differentiation. In addition, comparison of the spatial regulation of articular and growth plate cartilage revealed interesting similarities between the SZ of the articular cartilage and the HZ of the growth plate.

METHODS

Animal Procedures

All animal procedures were approved by the National Institute of Child Health and Human Development Animal Care and Use Committee. Although our previous spatial analysis of the growth plate cartilage was done in the rat, we chose to use the mouse in our current study of the articular cartilage based on two reasons. First, the mouse array is better annotated (39,000 transcripts) as compared with the rat array (30,000 transcripts). Second, many of the functional studies that give important insight into physiology and pathophysiology of the articular cartilage were done in the mouse using knockout models, and therefore, studying the spatial regulation of gene expression in the mouse articular cartilage will be more relevant to previous studies in the field. One-wk-old C57BL/6 male mice (Charles River Laboratory, Wilmington, MA) were killed by decapitation. Distal femur and proximal tibial epiphyses were excised. For laser capture and manual microdissection, cartilage was embedded in optimum cutting temperature compound (Electron Microscopy Sciences, Hatfield, PA), frozen on dry ice, and stored at -80 °C. For in situ hybridization and histology, cartilage was fixed overnight in 2% (v/v) paraformaldehyde (Electron Microscopy Sciences) and 0.2% (v/v) glutaraldehyde (Electron Microscopy Sciences) at 4 °C and decalcified in 10% (v/v) EDTA and 0.5% (v/v) paraformaldehyde, pH 7.4. Samples were then embedded in paraffin for sectioning.

LCM of Articular Cartilage

Frozen longitudinal cartilage sections (10 μm) were placed on PEN membrane glass slides (MDS Analytical Technologies, Sunnyvale, CA) precoated with 0.01% (w/v) poly-D-lysine (Sigma-Aldrich, St Louis, MO). Slides were thawed and stained as previously described (32) with modifications (70% ethanol, 30 s; deionized water, 15 s; Mayer's hematoxylin (Sigma-Aldrich), 30 s; deionized water, 15 s; Scott's bluing (Sigma-Aldrich), 15 s; 70% ethanol, 15 s; eosin Y (Sigma-Aldrich), 1 s; 95% ethanol, 30 s; 100% ethanol, 30 s; and xylene, 30 s) to shorten staining time and minimize tissue detachment. SZ, MZ, and DZ were microdissected using the cut and capture method in the Arcturus LCM System (Thermo Scientific, Waltham, MA) using CapSure HS Caps (MDS Analytical Technologies) and dissolved in tissue extraction buffer (PicoPure RNA Isolation Kit;

Manual Microdissection of Cartilage

To compare whole articular cartilage and whole growth plate cartilage expression, we performed manual microdissection of 1-wk-old mouse proximal tibial sections (50 μ m thickness) as previously described (33). To minimize cross-contamination, the lowermost articular cartilage DZ and the uppermost growth plate RZ were discarded. Tissue from 20 sections of a single animal was pooled prior to RNA isolation using the PicoPure RNA Isolation Kit. Approximately 1 μ g of total RNA was obtained from the articular or growth plate cartilage of one animal, with a RNA integrity number of 7–8.

Microarray Analysis

For each animal, total RNA from SZ, MZ, or DZ (n = 5 each) was amplified using WT-Ovation Pico RNA Amplification System (NuGEN, San Carlos, CA) following the manufacturer's protocol and hybridized to Mouse Genome 430 2.0 GeneChip Arrays (Affymetrix, Santa Clara, CA). Each array represents a single zone from a single animal, and a total of five animals were used. Affymetrix CEL files were imported into Partek Genomics Suite 6.6 (Partek, St Louis, MO) using the default normalization parameters. Probe-level data were preprocessed, including background correction, normalization, and summarization, using robust multiarray average analysis (34), and log_2 transformed. Bioinformatic analysis was performed using Ingenuity Pathways Analysis Software (Ingenuity Systems, Redwood City, CA). Heat maps were constructed using JMP 9 software (SAS Institute, Cary, NC).

Spatial Markers Identification

To identify useful molecular markers for each zone of the articular cartilage, we employed an empirical formula that was previously used to identify growth plate zonal marker (10). This formula gives a low score to genes that are expressed at high levels in one zone and at low levels in both of the other two zones. For example, the formula for SZ markers was: score = $(M + D + S_A)/S$, where S, M, and D represents expression level of a gene in the SZ, MZ, and DZ, respectively, and S_A represents the geometric mean of expression of all probes on the microarray in the SZ. Genes were then ranked from the lowest to highest score, and an arbitrary cutoff score of 0.5 was used for genes to qualify as good zonal markers.

Quantitative Real-Time PCR

Total RNA, without amplification, from individual animals was reverse transcribed into complementary DNA, and expression was quantified by real-time PCR using an ABI Prism 7900 Fast Sequence Detector (Thermo Scientific), according to the manufacturer's instructions. *Prg4, Cilp, Fgf18, Cytl1, Errf1, Dkk3, Tnn, Cxcl14, Sfrp5,* and *Frzb* mRNA levels were measured by SYBR green–based assays using custom-designed primers (**Supplementary Table S5** online), and the data were normalized to 18S ribosomal RNA measured with a predesigned Taqman assay (Thermo Scientific). Relative expression values were calculated as previously described (35).

In Situ Hybridization

cDNA from 1-wk-old mouse growth plate was amplified using Phusion DNA polymerase (Thermo Scientific) with primers that contained either a T7 promoter (for sense probes) or an SP6 promoter (for antisense probes) (**Supplementary Table S6** online). Single stranded digoxigenin-labeled riboprobes for *in situ* hybridization were transcribed using DIG RNA Labeling Kit (Roche Diagnostics, Indianapolis, IN) following the manufacturer's protocol. Riboprobes were purified by Micro Bio-Spin Columns P-30 Tris RNase free (Bio-Rad, Hercules, CA), followed by alkaline hydrolysis for 30 min as previously described (36). Paraffin-embedded sections of epiphyseal cartilage were hybridized to digoxigenin-labeled riboprobes as described by Bandyopadhyay *et al.* (37) and Murtaugh *et al.* (38).

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with some modifications. Notably, tissues were permeabilized with 300 mg/ml of proteinase K (Thermo Scientific) at 37 °C for 30 min prior to acetylation and prehybridization. For detection, tissue sections were incubated with anti-digoxigenin alkaline phosphatase Fab fragments (Roche Diagnostics) for 2 h at room temperature and then treated with nitro-blue tetrazolium chloride/5-bromo-4-chloro-3'indolyphosphate p-toluidine salt (NBT/BCIP) (Sigma-Aldrich) in the dark until a colorimetric change was detected. The sections were visualized using a ScanScope CS digital scanner (Aperio Technologies, Vista, CA) under bright field microscopy.

Immunostaining

Paraffin-embedded sections of epiphyseal cartilage were permeabilized with 100 mg/ml of proteinase K after rehydration and stained using rabbit anti- β -catenin antibody (C2206; Sigma-Aldrich). Vectastain ABC Kit (Vector Laboratories, Burlingame, CA) was then used for detection of primary antibody, and DAB peroxidase substrate (Vector Laboratories) was used to give a brown color for positive signal. Sections were counterstained with Eosin Y (Sigma) and visualized using a ScanScope CS digital scanner under bright field microscopy.

Statistical Analysis

Data are presented as mean ± SEM. Comparison of expression levels was performed by one-way ANOVA (SigmaPlot 10) using log-transformed relative expression data, followed by pairwise comparisons corrected for multiple comparisons using the Holm-Sidak method. All P values were two tailed, and significance was recognized at P <0.05. For comparison between the spatial regulation of articular cartilage and growth plate cartilage, gene-set enrichment analysis (Broad Institute, Cambridge, MA) was used according to user's manual (39), the probability of overlapping gene sets occurring by chance was determined using Pearson's χ^2 test (SigmaPlot 10).

SUPPLEMENTARY MATERIAL

Supplementary material is linked to the online version of the paper at http:// www.nature.com/pr

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