Suppression of Sost/Sclerostin and Dickkopf-1 Augment Intervertebral Disc Structure in Mice

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ABSTRACT
Intervertebral disc (IVD) degeneration is a leading cause of low back pain, characterized by accelerated extracellular matrix breakdown and IVD height loss, but there is no approved pharmacological therapeutic. Deletion of Wnt ligand competitor Lrp5 induces IVD degeneration, suggesting that Wnt signaling is essential for IVD homeostasis. Therefore, the IVD may respond to neutralization of Wnt ligand competitors sost(gene)/sclerostin(protein) and/or dkk1. Anti-sclerostin antibody (scl-Ab) is an FDA-approved bone therapeutic that activates Wnt signaling. We aimed to (i) determine if pharmacological neutralization of sclerostin, dkk1, or their combination would stimulate Wnt signaling and augment IVD structure and (ii) determine the prolonged adaptation of the IVD to global, persistent deletion of sost. Nine-week-old C57Bl/6J female mice (n = 6–7/group) were subcutaneously injected 2 × week for 5.5 weeks with scl-Ab (25 mg/kg), dkk1-Ab (25 mg/kg), 3:1 scl-Ab/dkk1-Ab (18.75:6.25 mg/kg), or vehicle (veh). Separately, IVD of sost KO and wild-type (WT) mice (n = 8/group) were harvested at 16 weeks of age. First, compared with vehicle, injection of scl-Ab, dkk1-Ab, and 3:1 scl-Ab/dkk1-Ab similarly increased lumbar IVD height and β-catenin gene expression. Despite these similarities, only injection of scl-Ab alone strengthened IVD mechanical properties and decreased heat shock protein gene expressions. Genetically and compared with WT, sost KO enlarged IVD height, increased proteoglycan staining, and imbibed IVD hydration. Notably, persistent deletion of sost was compensated by upregulation of dkk1, which consequently reduced the cell nuclear fraction for Wnt signaling co-transcription factor β-catenin in the IVD. Lastly, RNA-sequencing pathway analysis confirmed the compensatory suppression of Wnt signaling and revealed a reduction of cellular stress-related pathways. Together, suppression of sost/sclerostin or dkk1 each augmented IVD structure by stimulating Wnt signaling, but scl-Ab outperformed dkk1-Ab in strengthening the IVD. Ultimately, postmenopausal women prescribed scl-Ab injections to prevent vertebral fracture may also benefit from a restoration of IVD height and health. © 2022 The Authors. Journal of Bone and Mineral Research published by Wiley Periodicals LLC on behalf of American Society for Bone and Mineral Research (ASBMR).

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Introduction
There are no FDA-approved pharmacological treatments for intervertebral disc (IVD) degeneration, a major contributing factor of low back pain. Osteoporosis may contribute to IVD degeneration and some pharmacological treatments for bone maintenance target anabolic pathways innate to the IVD. Anti-sclerostin antibody (scl-Ab) treatment is an FDA-approved bone anabolic for postmenopausal women at high risk of vertebral fracture. Sclerostin and dkk1 are inhibitors of the Wnt/β-catenin signaling pathway and global suppression of sclerostin by systemic injection of scl-Ab or genetic ablation of its precursor SOST(human)/sost(mouse) promotes bone formation and mildly attenuates bone resorption. Individuals administered scl-Ab...
do not report an altered incidence of back pain than control subjects, suggesting it might not be harmful to the IVD. Although osteocytes in bone are the major source of sclerostin and dickkopf-1 (dkk1), IVD cells also express sclerostin and dkk1 but the impact of regulating sost/sclerostin or dkk1 on the IVD has yet to be determined.

Sclerostin and dkk1 are inhibitors of the canonical Wnt signaling pathway but differ in some notable ways. Both dkk1 and sclerostin interact with LRPS/6 to competitively prevent various Wnt ligands from binding to initiate the canonical Wnt signaling pathway. In the Wnt signaling pathway, where activation of this pathway is composed of β-catenin translocation to the cell nucleus, association with co-transcription factors T-cell factor (TCF) and lymphoid enhancer factor (LEF), and transcription of target genes. Wnt/β-catenin signaling regulates cell fate and extracellular matrix (ECM) anabolism in a range of musculoskeletal tissues. For instance, inactivation of Wnt signaling shifts differentiation of mesenchymal stem cells from osteoblastogenesis to chondrogenesis and activation in early chondrocytes triggers hyperchondrocyte maturation. Sclerostin and dkk1 can both bind to the first β-propeller of LRPS/6, but dkk1 can also bind to the second, third, and fourth β-propellers of LRPS/6. A pathway-related distinction between dkk1 and sclerostin is that dkk1 is a direct target of Wnt/β-catenin signaling pathway.

In the spine, the IVD development requires Wnt signaling and loss of Wnt signaling by aging and/or injury plumbs ECM anabolism. The nucleus pulposus serves as the hydration core of the IVD and houses notochordal cells that require Wnt signaling to maintain their cellular phenotype. Age- and injury-related reduction of Wnt signaling trigger the replacement of notochordal cells by more mature nucleus pulposus cells that are less equipped to produce ECM. Contrary, genetic stabilization of β-catenin in the nucleus pulposus increases notochordal cell expression and ECM anabolism and can promote ECM-related transcription during IVD injury. Lastly, in vivo deletion of LRPS in IVD cells reduces Wnt signaling and suggests that the IVD may be sensitive to Wnt ligand competitors that bind LRPS/6.

Therefore, we hypothesized that (i) neutralization of sclerostin and/or dkk1 and (ii) deletion of gene precursor to sclerostin sost would stimulate ECM anabolism in the IVD by increasing canonical Wnt signaling. Neutralization of sclerostin, dkk1, and in combination similarly increased Wnt signaling and IVD height. Next, using histology, MRI, qPCR, and RNA sequencing, global genetic deletion of sost increased the water content of the IVD, proteoglycan staining, IVD height, and decreased cellular stress mechanisms related to protein folding, but these changes were accompanied by gene and protein expression changes consistent with mature cell phenotypes by compensation of Wnt signaling. Overall, suppression of sost/sclerostin and/or dkk1 augment the structure of the IVD.

Materials and Methods

Mice

Nine-week-old C57Bl/6J female mice (n = 6–7/group) were injected with 25 mg/kg of either anti-sclerostin antibody (scl-Ab), anti-dkk1 antibody (dkk1-Ab), a 3:1 ratio of the two antibodies (18.75 mg/kg of scl-Ab and 6.25 mg/kg of dkk1-Ab), a 1:1 ratio (12.5 mg/kg of scl-Ab and dkk1-Ab), a 1:3 ratio (6.25 mg/kg of scl-Ab and 18.75 mg/kg of dkk1-Ab), or the buffer in which the antibodies were made (veh) for 5.5 weeks, twice per week (all previously described). Table 1. L3–S1 of these mice were used for another study and not analyzed here. Sost KO mice and their wild-type (WT) littermates (n = 8/group) total on a C57Bl/6 background have been previously described. Mice were housed in a 12-hour light/dark cycle, fed standard chow, and all experiments were performed with prior IACUC approval. Lumbar and caudal (tail) spinal sections were harvested. Mice were euthanized by hypoxia as a primary means and cervical dislocation as a secondary means. Spinal levels were further divided up for specific testing (Table 2).

Histology and immunohistochemistry

IVD for each stain were run in a single batch. L1–3 and CC10–11 were fixed in 15 mL of 10% formalin on a rocker for 24 hours, submerged in 70% ethanol, embedded in paraffin, and sectioned (5 μm). Safranin O/fast green counterstain images were analyzed by four independent observers for an average IVD degeneration score. In short, the nucleus pulposus (NP), annulus fibrosus (AF), and boundary between the two structures were scored based on structural properties and added for a total IVD score between 0 and 14, with increasing scores denoting greater IVD degeneration. Qualitatively, proteoglycan in the NP was estimated as the amount of intensity staining per area using ImageJ. Alicant blue staining was the counterstain for the IHC staining of sclerostin (BAF 1589, R&D Systems, Minneapolis, MN, USA), osterix (#22552, Abcam, Cambridge, MA, USA), and collagen 2 (II-11683-c, DSHB, Iowa City, IA, USA). For both osterix and collagen 2 quantification in the NP, positive cells were counted and compared with the total number of cells that were stained brown for the protein and blue for the cell nuclei. The percent of the area-stained brown for osterix was quantified and compared for the AF. Samples with sectioning anomalies and statistical outliers (see Statistics) were excluded from analysis. Total sample size per group was noted in each figure description.

Mechanical testing and analysis

After extraction of CC6–CC7 motion segments, the IVD was prepared, mechanically tested, and analyzed based on a previously published technique. The IVD was isolated from vertebrae CC6 and CC7 by dissection at the growth plate with the assistance of a microscope (M400 Photomakroscop; Wild, Heerbrugg, Switzerland). Growth plate-IVD-growth plate segments were imaged by an X-ray machine (Bruker, Kontich, Belgium) to determine IVD height using ImageJ (NIH). Next, a petri dish was glued to each IVD using cyanoacrylate and filled with phosphate buffered saline (PBS, pH 7.2) to maintain an osmotic environment. IVD were tested using a microindentation system (BioDent;
Table 2. Functional Spinal Unit or Bone for Each Outcome

| Outcome       | Tail       | Lumbar     |
|---------------|------------|------------|
| MRI           | CC6–7      | N/A        |
| Mechanics     | CC6–7      | N/A        |
| Western blot  | CC6–8, CC10–12 | N/A |
| qPCR          | CC8–10, CC4–5, and CC14–15 | L3–5 |
| Histology     | CC11–13    | L1–3       |

N/A = not applicable. CC4–5 and CC14–15 segments used for the injection-related groups.

Active Life Scientific, Santa Barbara, CA, USA) for quasi-static and dynamic mechanical properties. The testing sample was aligned with a 2.39 mm probe that fully covered the entire IVD and loaded sinusoidally (1 Hz) for 20 cycles under a preload of 20 grams at a compressive strain of 5%. Compressive strains were determined using X-ray-derived IVD heights. TriPLICATE trials were performed at each strain, with at least 10 minutes of rest time between each trial.

MATLAB code was used to remove the noise generated from the first and last loading cycle in the force-distance curve for each trial. The mechanical property outcomes from a semi-automated software program included relative maximum force (N), displacement (μm), loading stiffness (N/μm), unloading stiffness (N/μm), energy dissipation (N⋅μm = μJ), and loss tangent from the force-distance curve. Relative maximum force and displacement were calculated by considering the difference between the maximum and minimum values for these outcomes. Each outcome was computed as the average of the two trials with the lowest standard deviation out of three trials.

Magnetic resonance imaging (MRI)

Motion segment CC6–7 was submerged and wrapped in 1 × PBS-soaked gauze overnight until imaged. Imaging was completed on the Bruker BioSpin 9.4 T MRI, using a 400-mm-slice thickness for 2D imaging. The motion segment was imaged in a sagittal orientation using a 0.052 × 0.052 mm voxel resolution in the x-y direction and a 0.4 mm voxel resolution in the z direction taking 16 averages/slice. Two samples were stacked in a glass tube to remain upright, and two glass tubes were placed, separated by foam composite, inside of a 15 mL tube to ensure samples would not move while being imaged. Images were analyzed for quantification by ImageJ (NIH). Area and intensity of the IVD were determined and multiplied to estimate the hydration content of the IVD.

Micro-computed tomography

Motion segments L6–S1 and CC6–7 were harvested and submerged in 1 × PBS before imaging. Specimens were imaged using the Bruker SkyScan 1272 Micro-CT at a resolution of 8 μm. Vertebral bone was contoured at the periosteum and endosteum for trabecular and cortical analysis. For the trabecular analysis, the growth plate was used as a landmark and trabecular bone analysis consisted of the next 30 consecutive images (slices). For cortical analysis, the longitudinal center of the bone was identified and 15 images above and below were analyzed using the Bruker CTan64 MicroCT software. Parameters measured included bone volume/tissue volume (BV/TV), trabecular number (Tb.N), trabecular thickness (Tb.Th) for trabecular bone, and cross-sectional thickness for cortical bone, using a lower threshold of 60 and upper threshold of 225 for analysis.

QPCR

L3–5 and CC8–10 IVDs were harvested, frozen in liquid nitrogen, pulverized, and suspended in TRIzol (Ambion, Austin, TX, USA) until further processing. RNA isolation and purification steps were followed (RNeasy Mini Kit, Qiagen, Valencia, CA, USA) and RNA concentration was quantified (NanoDrop, Thermo Fisher Scientific, Waltham, MA, USA). CDNA was synthesized (script, Bio-Rad, Hercules, CA, USA) from 400 ng of total RNA for the following Taqman probes (Life Technologies, Carlsbad, CA, USA): aggrecan (Mm00565794_m1), keratin8 (Mm04209403_g1), dmp1 (Mm01208363_m1), sost (Mm00470497_m1), adams5 (Mm00478620_m1), collagen1 (Mm00801666_g1), encollagen2 (Mm01309565_m1), osterix (Mm04209856_m1), β-catenin (Mm01350387_g1), serpina1a (Mm02748447_g1), serpina1c (Mm04207703_m1), serpina1d (Mm00842095_m1), sostdc1 (Mm03024258_s1), foxa2 (Mm01976556_s1), axin2 (Mm00443610_m1), sfrp4 (Mm00840104_m1), gdf5 (Mm00433564_m), hspa1b (Mm03038954_s1), ccl5 (Mm00434946_m1), il1b (Mm00434228_m1), wnt16 (Mm00446420_m1), dkk1 (Mm00438422_m1), Wnt3a (Mm00437337_m1). Relative gene expression was normalized to IPOP (Mm01255158_m1) for each group and then experimental values (sost KO) were normalized to the average of the WT values (2ΔΔCt).

Western blotting

WT and sost KO IVDs between CC6–8 were isolated for whole-cell lysate and cytoplasmic and nuclear separation Western blots. IVDs were minced in ice-cold PBS (Thermo Fisher) containing 2% fetal bovine serum (FBS; Atlanta Biologicals, Flowery Branch, GA, USA) and protease inhibitor PMSF (Sigma, St. Louis, MO, USA). Two-tail IVDs from a single animal, per isolation method, were homogenized using a Tissue Tearor (BioSpec Products, Bartlesville, OK, USA). Whole-cell lysate was generated using diluted 1 × cell lysis buffer (Cell Signaling, Danvers, MA, USA) supplemented with PMSF. Fractionation of the nuclear protein was performed according to the Pierce cytoplasmic and nuclear extraction kit instructions (Sigma). Samples were run on an SDS-Page gel (Bio-Rad) and transferred to PVDF membrane (Bio-Rad). Blots were probed for anti-β-catenin (unphosphorylated; Cell Signaling) and subsequently an HRP-tagged anti-rabbit secondary antibody (#7074S, Cell Signaling). Whole-cell lysates were normalized to HRP-tagged actin antibody (#A3854, Sigma-Aldrich) and nuclear fractions to HRP-tagged histone H3 (#12648, Cell Signaling). All blots were developed using Immobilon Luminata Forte (Sigma), and the images were digitally collected with the Amersham imager 600 (GE Healthcare, Madison, WI, USA). Densitometry quantification was conducted on ImageJ to enumerate relative protein values between groups.

Bioinformatics

Gene set enrichment analysis (GSEA) software was used to generate the top down- and upregulated pathways by deletion of sost in the IVD. The input included gene name and raw counts for each sample. WEB-based Gene SeT AnaLysis Toolkit was used...
to generate the top up- and downregulated pathways between the WT and sost KO group. Using GSEA method of interest and geneontology for the functional database, the input included gene symbols and associated fold change for each gene. R Studio was used to generate plots such as PCA plot, volcano plot, and heat maps.

Fig. 1. Quantitative and qualitative intervertebral disc (IVD) structural properties. (A) Magnified images (5×) of Safranin O and fast green counterstained histological sections of the lumbar IVD for vehicle (veh, n = 5 mice), 25 mg/kg anti-sclerostin-antibody injection (scl-Ab, n = 5), and 25 mg/kg injection of anti-dkk1-antibody (dkk1-Ab, n = 5) (top row, left to right) and combination injection of 18.75 mg/kg anti-sclerostin-antibody and 6.25 mg/kg anti-dkk1-antibody (3:1, n = 5), combination of 12.5 mg/kg each of anti-sclerostin- and anti-dkk1-antibody (1:1, n = 7), and a combination injection of 6.25 mg/kg of anti-sclerostin-antibody and 18.75 mg/kg of anti-dkk1-antibody (1:3, n = 6) (bottom row, left to right). (B) Quantitative measurement of lumbar IVD height of all 6 groups. (C) Quantitative measurement of proteoglycan intensity staining in percentage of the lumbar IVD. Red staining for proteoglycan. Scale bar = 100 μm. Injection groups were compared with the vehicle group using a Dunnett’s test, with significance noted below p < 0.05.

Fig. 2. Injection of Scl-Ab improved intervertebral disc (IVD) mechanical properties. C57Bl6 mice were injected with vehicle (veh, n = 7 mice), 25 mg/kg anti-sclerostin-antibody injection (scl-Ab, n = 6), 25 mg/kg injection of anti-dkk1-antibody (dkk1-Ab, n = 7), combination injection of 18.75 mg/kg anti-sclerostin-antibody and 6.25 mg/kg anti-dkk1-antibody (3:1, n = 6), combination of 12.5 mg/kg each of anti-sclerostin- and anti-dkk1-antibody (1:1, n = 7), or a combination injection of 6.25 mg/kg of anti-sclerostin-antibody and 18.75 mg/kg of anti-dkk1-antibody (1:3, n = 7). (A) Loading stiffness, (B) unloading stiffness, (C) ultimate force, (D) energy dissipation, (E) displacement, and (F) loss tangent of the IVD. Injection groups were compared with the vehicle group using a Dunnett’s test, with significance noted below p < 0.05.
Statistics

Post ANOVA, a Dunnett’s test was used for the injection-related studies, with vehicles as the main comparison. A one-sample t test was used to test for outliers beyond 3 SD. A Student’s t test was used between WT and sost KO IVD and between veh and Scl-Ab. Box plots were used as the main graphical output. A box plot displays a five-number summary of the data. The top and bottom lines of the box plot represent the first and third quartile, while the center line in the box is the median. The top and bottom points represent the minimum and maximum of the data. Each point on the graph represents one sample. Total number of samples for each analysis are included in the figure legend. False discovery rate (FDR) was computed from p values using the Benjamini-Hochberg procedure. A p value or FDR value (where applicable) < 0.05 was considered significant.

Results

Injection of neutralization for sclerostin, dkk1, and in 3:1 combination increased IVD height

Preliminary data suggested that the IVD gene expression of sost relative to dkk1 is ~3:1 (data not shown). Therefore, in addition to scl-Ab or dkk1-Ab, we injected mice at 3:1, 1:1, and 1:3 scl-Ab:dkk1-Ab for consistency. Systemic administration of scl-Ab, dkk1-Ab, and the 3:1 combination of scl-Ab and dkk1-Ab increased lumbar IVD height (F = 7.74 and p < 0.001) by 22%, 26%, and 32% (p = 0.02, 0.02, 0.005), respectively (Fig. 1A, B) but the 1:1 and 1:3 combination injections did not significantly increase IVD height. Neither IVD degeneration score nor proteoglycan staining intensity were altered by any injections. In tail IVD, no variety of injection impacted IVD height (F = 0.73 and p = 0.60), proteoglycan staining intensity (F = 3.88 and p = 0.01), or IVD degeneration score (F = 1.25 and p = 0.31) (Supplemental Fig. S1).

Injection of sclerostin-neutralization antibody improved IVD mechanics

Despite no change in tail IVD height by injection of any Wnt signaling inhibitor neutralization combination compared with vehicle injection, injection of scl-Ab alone most improved the

Fig. 3. Injection of scl-Ab and/or dkk1-Ab increased pro-Wnt signaling gene expression. (A) Gene expression related to Wnt signaling in tail intervertebral disc (IVD): β-catenin, sost, and dkk1. Gene expression of β-catenin in the 3:1 combination injection was trending toward significance (p = 0.061). (B) Heat shock protein hspa1b, inflammatory marker serpinA1a, and extracellular matrix (ECM) marker aggrecan expression in the IVD. Scl-Ab injection trended to decrease gene expression of hspa1b (p = 0.069). Injection groups (veh: n = 4; scl-Ab: n = 5; dkk1-Ab: n = 6; and 3:1: n = 5) were compared with the vehicle group using a Dunnett’s test, with significance noted below p < 0.05.

Fig. 4. Scl-Ab injection reduced extracellular matrix (ECM), heat shock protein, and osterix gene expression in the intervertebral disc (IVD). Gene expression of scl-Ab injection (n = 5) compared with the vehicle-treated tail IVD (n = 4) for (A) col1a1, (B) col2a1, (C) hspa1a, and (D) osx. Student’s t tests compared groups, with significance noted below p < 0.05.
mechanical properties of tail IVD. Namely, injection of scl-Ab doubled ($p = 0.006$, $p = 0.008$, respectively) the loading ($F = 2.52$ and $p = 0.05$) and unloading stiffness ($F = 2.76$ and $p = 0.04$) of the IVD (Fig. 2). A similar trend appeared with ultimate force, but no changes were statistically significant ($F = 1.35$ and $p = 0.27$). Neither energy dissipation, displacement, nor loss tangent was affected by any injection group ($F = 3.44, 1.62, 1.59$, and $p = 0.01, 0.18, 0.19$).

Injection of sclerostin-neutralization-, dkk1-, and 3:1 combinatorial-antibody stimulated Wnt signaling

Quantitative PCR was determined in the injection groups that increased IVD height relative to vehicle: scl-Ab, dkk1-Ab, and 3:1 scl-Ab:dkk1-Ab. Scl-Ab and dkk1-Ab upregulated transcription of canonical Wnt signaling co-factor β-catenin ($F = 4.346$ and $p = 0.01$) by 1.7-fold ($p = 0.04$) and 1.2-fold ($p = 0.02$), respectively. Scl-Ab injection promoted β-catenin protein expression in the nucleus pulposus (NP). (A) Images of β-catenin protein expression in the NP of the vehicle ($n = 7$) and anti-sclerostin antibody group ($n = 6$). (A') Quantitative measurements of the fraction of β-catenin protein-expressing cells to the total number of cells in the NP. (B) Images of col2a1 protein expression in the NP of the vehicle ($n = 6$) and sclerostin antibody group ($n = 6$). (B') Quantitative measurements of the fraction of col2a1 protein-expressing cells to the total number of cells in the NP. Scale bar = 100 μm. Student’s t tests compared groups, with significance noted below $p < 0.05$.

Fig. 6. Sost deletion reduced sost gene and protein expression in the intervertebral disc (IVD). Sost gene expression from qPCR and qualitative images of sclerostin staining of the WT and sost KO IVD ($n = 5$) was effectively deleted from the sost KO group. (A) qPCR of sost from WT IVD (CT: 34.09–34.87, 25–75%). Sost gene expression was not detected in sost KO IVD. (B) Magnified images (40×) of the nucleus pulposus (NP) of the WT IVD (top) and sost KO IVD (bottom) using Alcian blue counterstain to show sclerostin protein expression in the cells of the IVD. The WT expressed defined dark brown staining (black arrows), whereas the sost KO cells expressed less, more diffuse staining, indicating less sclerostin expression in the sost KO. (C) Magnified image (40×) of the annulus fibrosus (AF). Scale bar = 100 μm. Dashed box represents the highlighted NP portion (C). Solid box represents the highlighted portion of the AF (D).
respectively (Fig. 3A). Injection of the 3:1 combination trended to upregulate β-catenin expression. Corroboratively, scl-Ab reduced sost gene expression by 85% \((p = 0.0004)\) and 3:1 combination reduced sost gene expression by 61% \((p = 0.006\) versus veh, \(F = 3.452, p = 0.023\)). Highly variable dkk1 expression in the vehicle group obfuscated reduction by each injection group. In addition, we determined the potential regulation of dkk1 by scl-Ab and vice versa because dkk1 and sclerostin share the capacity to bind the first propeller of LRP5/6 or may compensatorily upregulate the other. Neither injection of antibody significantly regulated the transcription of the other antibody’s precursor in the IVD. Heat shock protein a1b \((hspa1b)\) gene expression in the IVD was not different between groups, but scl-Ab trended to reduce hspa1b \((p = 0.069)\). Hspa1b and serpinA1a were similarly regulated by each antibody injection \((R = 0.83, p < 0.05)\), but neither hspa1b \((F = 2.43\) and \(p = 0.11\)), serpina1a \((F = 1.58\) and \(p = 0.23\)), nor aggrecan \((F = 1.70\) and \(p = 0.21)\) expression were changed by any injection compared with vehicle (Fig. 3B). Inflammation-related markers cxcl9 and il-1β were not detected in any IVD.

Anti-sclerostin-antibody injection increased β-catenin protein and reduced mature NP gene expression in the IVD

Based on the mechanical property improvements by scl-Ab, low variability of upregulation of β-catenin by scl-Ab, and trending downregulation of hspa1b, we further characterized ECM transcription and cell phenotype in the IVD by scl-Ab. Injection of scl-Ab decreased the gene expression of col1a1 by 57% \((p = 0.002)\), col2a1 by 54% \((p = 0.02)\), and oste in by 35% \((p = 0.049)\) (Fig. 4A, B, and D). Further, scl-Ab increased the number of cells expressing β-catenin protein in the nucleus pulposus by 81% \((p = 0.004)\) and reduced the number of nucleus pulposus cells that expressed col2a1 by 46% \((p = 0.01)\) (Fig. 5B). Similar to hspa1b, scl-Ab decreased the gene expression of hspa1a, a gene associated with cellular stress, \((31)\) by 95% \((p = 0.02)\) (Fig. 5C).
Systemic deletion of *sost* reduced sclerostin protein expression in the IVD

Quantitatively, *sost* gene expression was consistently detectable in the WT IVD, but *sost* was not detected in any of the *sost* KO IVD (Fig. 6A). Qualitatively, sclerostin protein expression (brown staining, black arrow) appeared in the NP cells of WT IVD (Fig. 6B). By contrast, sclerostin staining in the *sost* KO was minimally expressed in NP cell nuclei and relegated to the cell membrane and extracellular matrix (Fig. 6C). The AF minimally expressed sclerostin (Fig. 6D).

Deletion of *sost* in the IVD increased IVD height, proteoglycan staining, and hydration

Proteoglycans are hydrophilic proteins, and loss of proteoglycans is a common feature of IVD degeneration.\(^{32,33}\) Compared with WT IVD, deletion of *sost* KO increased lumbar IVD height by 21% (\(p = 0.007\)) and tail proteoglycan staining by 150% (\(p = 0.02\)) in the NP (Fig. 7A–C). However, the accumulation of proteoglycan staining, while potentially beneficial for hydration, led to slight disorganization of the band of cells in the NP, statistically insignificantly increasing the histological IVD degeneration score (Supplemental Fig. S2). Next, we determined the

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**Table 3. Area, Intensity, and Water Content Determined by MRI in WT and *sost* KO Intervertebral Disc**

| Genotype | Area (mm\(^2\)) | Intensity | Water content |
|----------|-----------------|-----------|---------------|
| WT 1     | 0.32            | 956       | 301.12        |
| WT 2     | 0.32            | 1235      | 395.20        |
| WT 3     | 0.32            | 1004      | 391.56        |
| WT AVG   | \(0.34 \pm 0.04\) | \(1065 \pm 149\) | \(362.63 \pm 52.29\) |
| SOST KO 1| 0.65            | 1145      | 739.67        |
| SOST KO 2| 0.64            | 1155      | 733.43        |
| SOST KO 3| 0.34            | 1225      | 416.50        |
| SOST KO 4| 0.40            | 1501      | 600.40        |
| SOST KO AVG | \(0.51 \pm 0.16\) | \(1257 \pm 167\) | \(622.50 \pm 151.61^*\) |

*Student’s t tests compared groups, with significance noted \(p < 0.05\) (*). Values in bold represent the mean.
hydration of the IVD by MRI, a standard noninvasive clinical imaging technique for determining the morphology and hydration of the IVD.\textsuperscript{34,35} WT IVD had a patchy distribution of water, whereas the IVD from the AF was clearer and brighter in the sost KO IVD (Fig. 8). Relative to the WT IVD, sost KO increased the estimated water content, i.e., hydration, by 72% ($p = 0.04$), but there was no statistical difference in the total area of the IVD or intensity of the water between groups (Fig. 8; Table 3 rpinA1d, sostdc1, keratin8, wnt16). These data show that deletion of sost mimicked the IVD.

Sost deletion induced compensatory normalization of Wnt signaling by upregulation of Wnt signaling inhibitors

Canonical Wnt signaling requires translocation of $\beta$-catenin to the cell nucleus. Therefore, we determined the nuclear and cytoplasmic fraction of $\beta$-catenin protein. Sost KO IVD had less active $\beta$-catenin protein in the cell nuclei than WT IVD (Fig. 9A), suggesting less Wnt signaling. The cytoplasmic fraction was not statistically different between groups (Supplemental Fig. S3). In addition, dkk1 is a target of Wnt/$\beta$-catenin signaling, so we also determined the gene expression of Wnt inhibitors. Sost KO IVD expressed greater dkk1 by 10-fold ($p = 0.04$) and sfrp4 by fourfold ($p = 0.001$). Lack of differences in gene expression of co-transcription factor $\beta$-catenin between WT and sost KO IVD suggested compensation of Wnt signaling after deletion of sost.

Compensation of Wnt signaling from sost deletion triggered extracellular matrix degradation and the expression of a mature NP cell phenotype in the IVD

Sost KO regulated gene and protein expression of extracellular matrix metabolism toward anti-anabolism, catabolism, and IVD cell phenotypes toward chondrogenesis. Specifically, deletion of sost downregulated gene expression of aggrecan by 52% ($p = 0.03$) (Table 4A). Common markers of notochordal cells and mature NP cells were determined, respectively. Sost KO IVDs expressed less transcription of notochordal marker foxA2\textsuperscript{360} by 77% ($p = 0.01$) (Fig. 10B), less transcription of early NP marker col2a1 by 67% ($p = 0.05$) (Fig. 10A), and fewer cells in the NP-expressed col2a1 (Fig. 11A, A'). By contrast, accrual of mature NP can be marked by increased chondrogenic marker\textsuperscript{377} and osteoblast transcription factor sp7 (Oxs). Compared to WT IVD, sost KO IVD expressed increased osterix gene expression by 4.4-fold ($p = 0.03$) and protein expression (Figs. 10B and 11B, B'). Sost deletion similarly altered gene expression in lumbar IVD from rma-seq with upregulation of ECM degradation, downregulation of notochordal marker gdf5\textsuperscript{380} downregulation of early mature NP markers (ColX\textsuperscript{399}), and upregulation of mature NP cell marker bglap (Supplemental Fig. S4). Gene expression that was not statistically different between WT and sost KO IVD included serpinA1a-c (serpinA1c+, $p = 0.06$), serpinA1d, sostdc1, keratin8, wnt16, Water Content Determined by axin2, serpinA1a, cxcl9, il-1$\beta$, and wnt3a (Table 4).

Using whole transcriptomic analysis, sost deletion downregulated pathways related to protein folding and upregulated pathways related to immune response in the IVD

In an overall sense, based on the PCA plot (Fig. 12A), sost KO IVDs are not transcriptomically distinct from WT IVDs. Nevertheless, 35% of the top 20 downregulated pathways were related to cellular response to external stimuli (e.g., protein folding, FDR < 0.05), 15% to metabolism, 15% to extracellular matrix organization, 10% to gene expression, and 5% to developmental biology, signaling transduction, metabolism of proteins, transport of small molecules, and other pathways (Fig. 12B), whereas of the top 20 upregulated pathways, 85% were related

### Table 4. Genes Not Significantly Regulated by sost Deletion or Expressed in the Intervertebral Disc

| Gene             | WT Value | Sost KO Value | p Value |
|------------------|----------|---------------|---------|
| SerpinA1C        | 1 ± 0.18 | 0.72 ± 0.09   | 0.06    |
| SerpinA1D        | 1 ± 0.56 | 0.52 ± 0.19   | 0.11    |
| Sostdc1          | 1 ± 0.35 | 1.86 ± 0.71   | 0.11    |
| Keratin8         | 1 ± 0.83 | 0.50 ± 0.50   | 0.26    |
| Wnt 16           | 1 ± 0.69 | 1.79 ± 1.46   | 0.36    |
| Axin 2           | 1 ± 0.53 | 1.38 ± 0.75   | 0.38    |
| SerpinA1A        | 1 ± 0.37 | 0.52 ± 0.19   | 0.80    |
| Cxcl9            | Undetermined | Undetermined | N/A      |
| IL-1β            | Undetermined | Undetermined | N/A      |
| Wnt 3a           | Undetermined | Undetermined | N/A      |

$\text{N/A} = \text{not applicable. Student's t tests compared groups, with significance noted p < 0.05.}$

![Fig. 10. Global deletion of sost induced the gene expression of a mature cell type in the intervertebral disc (IVD). Gene expression of (A) extracellular matrix (ECM) and (B) transcription factors of cell type markers in WT and sost KO IVD (n = 5, group). Student’s t tests compared groups, with significance noted below p < 0.05.](image-url)
to immune pathways (FDR < 0.05) and 5% each to cell cycle (FDR < 0.05), signaling transduction (FDR < 0.05), and hemostasis (FDR < 0.05). Specifically, the heat map for the top downregulated pathway was “protein folding,” which included heat shock proteins, and for the top upregulated pathway was immune response (Fig. 12C). More specifically, the volcano plot highlights in red the genes that were most significantly differentially regulated (Fig. 12D). A list of the top 20 differentially downregulated and top 20 upregulated were included (Tables 5 and 6). Lastly, GSEA analysis for the sost KO IVDs corroborated that Wnt signaling was reduced based on the normalized enrichment score (−1.53, FDR < 0.01; Tables 7 and 8; Supplemental Fig. S5).

Sost KO and scl-Ab increased vertebral bone structure

Osteocytes and osteoblasts are the predominate sclerostin-expressing cells.\(^{(15,40)}\) Here, WT tail vertebra expressed sclerostin staining in both osteocytes and osteoblasts, whereas global deletion of sost blunted its expression in the vertebra (Supplemental Fig. S6) and, consequently, increased trabecular and cortical bone structure (Table 8). Scl-Ab may have induced a compensatory upregulation of sclerostin protein (Supplemental Fig. S7). Separately, scl-Ab increased the number of osteocytes expressing substance P because of the need for innervation during bone formation.\(^{(41)}\)

**Discussion**

FDA-approved anti-sclerostin antibody (scl-Ab) augmented the structure and strength of the IVD. We hypothesized that increasing Wnt signaling by suppression of Wnt competitors by pharmacological or genetic approaches would augment the structure of the IVD. Sclerostin and dkk1 are known inhibitors of the canonical Wnt signaling pathway and work in a similar fashion.\(^{(62)}\) Scl-Ab, dkk1-Ab, and the 3:1 combinatorial injection of sclerostin and dkk1 increased lumbar IVD height via strong upregulation of Wnt/\(\beta\)-catenin signaling, but injection of a 1:1 ratio of sclerostin to dkk1 antibody or a 1:3 ratio were not as beneficial. By contrast, only scl-Ab injection improved the mechanical properties of the IVD. Prolonged suppression of sclerostin by global deletion of sost increased IVD height, water content, and proteoglycan staining but induced a significant normalization of Wnt signaling. Overall, these data show that systemic administration of scl-Ab promotes major features of the IVD that are lost with IVD degeneration.

Sclerosteosis patients with a sost gene mutation are associated with greater bone mass and stature.\(^{(43)}\) Although mice with injection of scl-Ab,\(^{(44)}\) conditional deletion of sost in bone cells, or global KO of sost all have greater bone mass, vertebral bone length does not change.\(^{(45)}\) Therefore, neutralization of sost/sclerostin may extend IVD height by direct stimulation of Wnt signaling and/or adaptation to greater bone mass.

IVD cells express Wnt signaling inhibitors sost/sclerostin and dkk1,\(^{(45)}\) but their role in the IVD was undetermined. Nucleus pulposus-specific upregulation and downregulation of Wnt signaling co-transcription factor \(\beta\)-catenin is anabolic and catabolic to the ECM/stiffness of the IVD, respectively.\(^{(12)}\) Here, injections of scl-Ab, dkk1-Ab, and in combination increased IVD height and \(\beta\)-catenin gene expression. We also administered combinatorial injections of scl-Ab and dkk1-Ab (3:1, 1:1, and 1:3), where 3:1 injection outperformed 1:1 and 1:3 injections in increasing IVD height. Therefore, we excluded 1:1 and 1:3 injections from further examination. The 3:1 injection most closely approximated the WT gene expression of sost relative to dkk1, which could have most effectively neutralized sclerostin and dkk1 to upregulate Wnt/\(\beta\)-catenin signaling in the IVD.

Our data and others have noted that sclerostin and dkk1 share a mutual compensatory regulation of the Wnt/\(\beta\)-catenin signaling pathway,\(^{(11,12,29)}\) which may impact IVD cell differentiation. We found that global sost deletion induced a strong
upregulation of Wnt signaling inhibitors that consequently normalized β-catenin protein in IVD cell nuclei and Wnt signaling. Normalization of Wnt signaling may have attenuated the initial benefits of activation and could explain the subdued difference between sost KO and WT IVD. Similarly, we noted compensation of sclerostin in osteocytes of sclerostin-antibody-treated mice. Next, we previously found that similar to ECM anabolism, regulation of Wnt signaling in the IVD was associated with differential regulation of cell type expression.

| Table 5. Top 20 Genes Upregulated by Sost KO |
| Genes | Fold change | p Value | False discovery rate |
|-------|-------------|---------|---------------------|
| Ntn   | 3.34        | <0.001  | <0.001              |
| Tmem181b-ps | 2.66       | <0.001  | <0.001              |
| Sfrp4 | 3.97        | <0.001  | <0.001              |
| Gm1821 | 6.58       | <0.001  | 0.002               |
| H2-Q10 | 5.36       | <0.001  | 0.007               |
| Wdfy1 | 1.90        | <0.001  | 0.091               |
| Mepe  | 3.66        | <0.001  | 0.149               |
| Clec10a | 3.49       | <0.001  | 0.233               |
| Sp6   | 2.85        | <0.001  | 0.268               |
| Cdl1d1 | 2.53       | <0.001  | 0.340               |
| Fasn  | 2.91        | <0.001  | 0.340               |
| Tnfrsf19 | 1.79      | <0.001  | 0.359               |
| Sic1  | 3.43        | <0.001  | 0.364               |
| Dgat2 | 2.27        | <0.001  | 0.364               |
| Serping1 | 1.93     | <0.001  | 0.371               |
| Aggpat2 | 1.94       | <0.001  | 0.421               |
| Cxcl9 | 3.35        | 0.001   | 0.512               |
| Cebpα | 1.95        | 0.001   | 0.513               |
| Efcb7 | 2.21        | 0.001   | 0.513               |
| Myh7  | 33.95       | 0.001   | 0.513               |

| Table 6. Top 20 Genes Downregulated by Sost KO |
| Genes | Fold change | p Value | False discovery rate |
|-------|-------------|---------|---------------------|
| Gm14305 | −109.99   | <0.001  | <0.001              |
| 1700030C10Rik | −153.71 | <0.001  | <0.001              |
| Hspa1b | −37.64     | <0.001  | <0.001              |
| Hist1h1b | −6.72    | <0.001  | 0.051               |
| Hsp1  | −7.38      | <0.001  | 0.254               |
| Rec8  | −2.34      | <0.001  | 0.286               |
| Eps8l1 | −8.73     | <0.001  | 0.286               |
| Rn45s | −3.69      | <0.001  | 0.286               |
| Hspa1a | −4.57     | <0.001  | 0.286               |
| Aldh3a1 | −4.52   | 0.001   | 0.513               |
| Serpine1 | −2.97    | 0.001   | 0.513               |
| Slc15a2 | −3.13     | 0.001   | 0.513               |
| Nfatc2 | −9.53     | 0.002   | 0.565               |
| Erdr1 | −1.93      | 0.002   | 0.656               |
| Foxa2 | −3.21      | 0.002   | 0.656               |
| Gm7120 | −2.34      | 0.003   | 0.656               |
| Egf1  | −2.08      | 0.003   | 0.776               |
| Hsp90aa1 | −1.75   | 0.003   | 0.788               |
| Gdf5  | −1.96      | 0.005   | 1.000               |
Table 7. Top 20 Downregulated Pathways by Sost Deletion

| Name                        | Normalized enrichment score | Nominal p value | False discovery rate |
|-----------------------------|-----------------------------|-----------------|----------------------|
| Interferon α-response       | -1.57                       | <0.001          | 0.007                |
| Bile acid metabolism        | -1.56                       | <0.001          | 0.006                |
| Notch signaling             | -1.56                       | 0.010           | 0.006                |
| Wnt β-catenin signaling     | -1.53                       | 0.011           | 0.008                |
| Peroxisome                  | -1.52                       | 0.001           | 0.009                |
| Allograft rejection          | -1.51                       | 0.001           | 0.010                |
| Interferon γ-response       | -1.47                       | 0.000           | 0.016                |
| UV response DN              | -1.39                       | 0.006           | 0.036                |
| Xenobiotic metabolism       | -1.36                       | 0.006           | 0.047                |
| Fatty acid metabolism       | -1.32                       | 0.013           | 0.072                |
| Kras signaling up           | -1.30                       | 0.006           | 0.082                |
| Adipogenesis                | -1.29                       | 0.004           | 0.089                |
| Coagulation                 | -1.26                       | 0.059           | 0.118                |
| IL6 JAK STAT3 signaling     | -1.23                       | 0.112           | 0.161                |
| Inflammatory response       | -1.16                       | 0.133           | 0.317                |
| Angiogenesis                | -1.15                       | 0.263           | 0.311                |
| Hedgehog signaling          | -1.11                       | 0.326           | 0.441                |
| Oxidative phosphorylation   | -1.09                       | 0.226           | 0.479                |
| Apical junction             | -1.09                       | 0.243           | 0.478                |
| Complement                  | -1.09                       | 0.262           | 0.462                |

Table 8. Sost KO Vertebral Bone Structural Properties

| Bone parameter | Tail                      | Lumbar            |
|----------------|---------------------------|-------------------|
|                | WT                        | Sost KO           | WT                | Sost KO         |
| Tb.BV/TV (%)   | 49.4 ± 3.1                | 53.5 ± 4.6        | 25.3% ± 4.8       | 45.7 ± 5.1*     |
| Tb.N (1/mm)    | 5.52 ± 0.30               | 5.84 ± 0.14       | 6.11 ± 0.50       | 6.45 ± 0.76     |
| Tb.Th (mm)     | 0.09 ± 0.01               | 0.09 ± 0.01       | 0.07 ± 0.01       | 0.07 ± 0.01     |

Tb.BV/TV = trabecular bone volume; Tb.N = trabecular number; Tb.Th = trabecular thickness. *: p < 0.05
dkk1-Ab similarly increased IVD height as each alone, when the ratio of the administration reflected in vivo ratios, but scl-Ab alone most strengthened the IVD. Persistent genetic deletion of sost increased proteoglycan staining and IVD hydration but also induced a compensatory upregulation of Wnt signaling inhibitors that consequently normalized Wnt signaling. Together, these data show that the musculoskeletal benefits of scl-Ab romosozumab (Evenity), which is commercially available, may extend beyond bone to improve key features of the IVD and could potentially be used as a therapeutic for IVD degeneration.

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All authors state that they have no conflicts of interest.

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**Author Contributions**

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**Data Availability Statement**

The data that support the findings of this study are openly available in biorxiv at [http://doi.org/10.1101/2021.07.01.449486](http://doi.org/10.1101/2021.07.01.449486) (2021), reference number [2021.2007.2001.449486]
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