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Novel Wnt Regulator NEL-Like Molecule-1 Antagonizes Adipogenesis and Augments Osteogenesis Induced by Bone Morphogenetic Protein 2

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From the Divisions of Growth and Development and Section of Orthodontics,* Advanced Prosthodontics,[§] and Diagnostic and Surgical Sciences,[¶] UCLA School of Dentistry, the UCLA Division of Plastic and Reconstructive Surgery,[†] Department of Orthopaedic Surgery and Orthopaedic Hospital Research Center at UCLA, and the Department of Pathology and Laboratory Medicine,[‡] David Geffen School of Medicine at UCLA, University of California, Los Angeles, Los Angeles, California

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Address correspondence to Kang Ting, D.M.D., UCLA School of Dentistry, University of California, Los Angeles, Box 951668, CHS 30-113, Los Angeles, CA 90095-1668; or Chia Soo, M.D., (Correspondent for the Editorial and Production offices), UCLA Division of Plastic and Reconstructive Surgery and Department of Orthopaedic Surgery and the Orthopaedic Hospital Research Center, University of California, Los Angeles, Los Angeles, CA 90095-1579. E-mail: kting@dentistry.ucla.edu or bsoo@ucla.edu. The differentiation factor NEL-like molecule-1 (NELL-1) has been reported as osteoinductive in multiple *in vivo* preclinical models. Bone morphogenetic protein (BMP)-2 is used clinically for skeletal repair, but *in vivo* administration can induce abnormal, adipose-filled, poorquality bone. We demonstrate that NELL-1 combined with BMP2 significantly optimizes osteogenesis in a rodent femoral segmental defect model by minimizing the formation of BMP2-induced adipose-filled cystlike bone. *In vitro* studies using the mouse bone marrow stromal cell line M2-10B4 and human primary bone marrow stromal cells have confirmed that NELL-1 enhances BMP2-induced osteogenesis and inhibits BMP2-induced adipogenesis. Importantly, the ability of NELL-1 to direct BMP2-treated cells toward osteogenesis and away from adipogenesis requires intact canonical Wnt signaling. Overall, these studies establish the feasibility of combining NELL-1 with BMP2 to improve clinical bone regeneration and provide mechanistic insight into canonical Wnt pathway activity during NELL-1 and BMP2 osteogenesis. The novel abilities of NELL-1 to stimulate Wnt signaling and to repress adipogenesis may highlight new treatment approaches for bone loss in osteoporosis. (*Am J Pathol 2016, 186: 419–434; http://dx.doi.org/10.1016/j.ajpath.2015.10.011*)

NEL-like molecule-1 (NELL-1) is an osteoinductive growth factor first identified through its overexpression in pathologically fusing suture specimens from patients with craniosynostosis.^{1,2} Transgenic *Nell1*-overexpressing mice recapitulate craniosynostosis-like phenotypes, exhibiting gross calvarial bone overgrowth and increased osteoblast differentiation.³ Conversely, *Nell1* deficiency severely disrupts bone growth, as mice with nonsense mutations in *Nell1* die perinatally with major skeletal anomalies in the craniofacial complex, spine, and long bones.^{4–6} Highlighting the central role of NELL-1 in skeletal development, NELL-1 mediates key downstream effects of the master osteogenic regulator runt-related transcription factor 2 (RUNX2)⁷ and can partially rescue *RUNX2*

Copyright © 2016 American Society for Investigative Pathology. Published by Elsevier Inc. All rights reserved. http://dx.doi.org/10.1016/j.ajpath.2015.10.011 loss of function.⁸ NELL-1 can also transiently activate mitogen-activated protein kinase signaling to induce RUNX2 phosphorylation and osteogenic differentiation.⁹ Recently, we

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demonstrated that osteoinductive effects of NELL-1 are partially mediated through binding to the intracellular molecule apoptosis related protein 3 and integrin- β_1 .¹⁰ In translational models of bone repair, exogenous NELL-1 induces potent osseous healing of critical-sized rat calvarial defects,¹¹ repair of rat femoral segmental defects (FSDs),¹² as well as successful spinal fusion in rats^{13,14} and sheep.¹⁵ Based on these results demonstrating reproducible preclinical osteoinductivity in multiple small and large animal models, NELL-1 is being developed for therapeutic use in spinal fusion in humans.

Bone morphogenetic protein-2 (BMP2) is a U.S. Food and Drug Administration (FDA)-approved osteoinductive growth factor most commonly used for spinal fusions and treatment of skeletal defects.^{16,17} Notably, BMP2 has species-specific dosing responses in which the required BMP2 concentration for osteogenesis increases with phylogenetic complexity.¹⁸ The BMP2 concentration for consistent bone formation in nonhuman primates is 0.75 to 2.0 mg/mL, but in rodents is only 0.02 to 0.4 mg/mL.¹⁹ Based on data from nonhuman primates, the minimum effective human BMP2 concentration was initially set at 1.5 mg/mL (total dose, 4.2 to 12 mg) in pilot and pivotal trials in humans and is currently the approved concentration for clinical use.¹⁹⁻²¹ Unfortunately, the high BMP2 concentrations required for osteogenesis in humans are associated with significant adverse effects, including an FDA warning of life-threatening cervical swelling,²² ectopic bone formation (U.S. Food and Drug Administration, http://www.fda. gov/MedicalDevices/Safety/AlertsandNotices/PublicHealth Notifications/ucm062000.htm, last accessed September 28, 2011),²³ osteoclastogenesis,²⁴ and inconsistent bone formation.²⁵ Thus, improving the safety and efficacy of BMP2based skeletal repair is crucial. Although NELL-1 is being developed as a standalone osteoinductive therapeutic, we previously performed studies of NELL-1 and BMP2 in animals in an effort to better understand the molecular and cellular bases of NELL-1 and/or BMP2 osteoinductivity.²⁶ Intriguingly, we found that NELL-1 synergistically increases bone formation with BMP2 through, in part, inhibition of BMP2-induced adipogenesis,²⁶ a well-described phenomenon documented in several in vitro and in vivo models.^{11,27,28}

Canonical (β -catenin–dependent) Wnt signaling regulates mesenchymal stem cell maintenance and differentiation during bone development and homeostasis.^{29–32} Generally, increased Wnt signaling promotes osteogenesis over adipogenesis.³³ In mice, targeted disruption of *Axin2*, a negative regulator of Wnt signaling,³⁴ induces craniosynostosis-like skull malformations,³⁵ a phenotype similar to that in *Nell1*–overexpressing mice.³ Because canonical Wnt is a major signal transduction pathway that regulates mesenchymal stem cell commitment to osteogenic over adipogenic lineages, we examined whether NELL-1 may synergize with BMP2 in bone formation through β -catenin–dependent Wnt signaling. Here we show NELL-1 activation of canonical Wnt signaling as a mechanistic basis for the observed NELL-1 inhibition of BMP2-induced adipogenesis and for NELL-1+BMP2 synergy in bone formation. Importantly, we establish NELL-1 as a novel Wnt pathway activator that can be applied, with BMP2 to improving the safety and efficacy of currently available bone-regeneration therapies. Overall, treatment with NELL-1+BMP2 forms bone of better quality than that formed with either NELL-1 or BMP2 alone.

Materials and Methods

Antibodies and Reagents

Human recombinant NELL-1 was purified by Aragen Bioscience (Morgan Hill, CA). Human BMP2 (Infuse Bone Graft) was purchased from Medtronic (Minneapolis, MN). Primary antibodies used in this study were anti-Runx2 (sc-10758; Santa Cruz Biotechnology, Santa Cruz, CA), antiperoxisome proliferator-activated receptor γ (PPAR γ ; sc-7273; Santa Cruz Biotechnology), anti-CCAAT enhancer-binding protein α (C/EBP α) (ab63486-100; Abcam, Cambridge, MA), antiosteocalcin (sc-18322; Santa Cruz Biotechnology), anti-β-catenin (610153; BD Biosciences, Franklin Lakes, NJ), anti-active β-catenin (05-665; Millipore, Billerica, MA) and anti $-\beta$ -actin (sc-1616; Santa Cruz Biotechnology). Recombinant mouse Wnt-3a protein (1324-WN-002) and Dickkopf-related protein 1 (Dkk1) (5897-DK-010) were purchased from R&D Systems (Minneapolis, MN). XAV939 was purchased from Cayman Chemical (Ann Arbor, MI).

Implant Material

Cylindrical poly(lactic-*co*-glycolic acid) (PLGA) scaffold implants were fabricated as described previously.^{12,36} Briefly, 85/15 D,L-PLGA (inherent viscosity, 0.61 dL/g; Absorbable Polymers, Pelham, AL)/chloroform solutions were mixed with 200 to 300 μ m—diameter sucrose to obtain 94% porosity and were compressed in a polytetrafluoroethylene mold to form scaffolds measuring 6 mm in length and 4 mm in diameter. After freeze-drying overnight, scaffolds were immersed in three changes of distilled deionized water (ddH₂O) to dissolve the sucrose. All scaffolds were disinfected by immersion in 70% ethanol for 30 minutes, followed by three rinses with ddH₂O.

Scaffolds were then coated with hydroxyapatite to ensure efficient NELL-1 and BMP2 loading and to enhance overall osteoconductivity of the hydrophobic PLGA scaffold. Briefly, two sterile, supersaturated solutions containing ion concentrations fivefold greater than those in human plasma [simulated body fluid (SBF) 1 and 2] were sequentially applied to the scaffolds. The ionic concentrations and preparation of SBF 1 and SBF 2 were previously published.³⁶ Immediately before the coating process, dried PLGA scaffolds were subjected to glow discharge argon

plasma etching (Harrick Scientific, Ossining, NY) to improve wetting and coating uniformity. Etched PLGA scaffolds were then incubated in SBF 1 for 12 hours and changed to SBF 2 for another 12 hours at 37° C inside of a water-jacketed incubator. Coated PLGA scaffolds were rinsed gently with sterile ddH₂O to wash away excess sodium chloride solution and were dried in the laminar flow hood. The thickness of the hydroxyapatite coating was 40 to 100 nm. Apatite-coated PLGA scaffolds did not exhibit a reduction in porosity and maintained the same pore interconnectivity and pore size ranging between 200 and 300 µm.

Before implantation, scaffolds were impregnated with recombinant human BMP2 and/or recombinant human NELL-1. Doses of each growth factor (Table 1) were chosen based on the ability of BMP2 to consistently induce adipocytic bone cysts¹² and NELL-1 to significantly improve bone regeneration in an FSD model in our previous studies.³⁷ Each growth factor was diluted to the appropriate concentration in phosphate-buffered saline (PBS), added dropwise uniformly to the scaffolds for 20 minutes, and lyophilized in a freeze-drier overnight at -20° C. Implants for PBS controls were generated in the same way, except that no growth factors were added.

Femoral Segmental Defect Animal Model

All animal procedures were conducted in accordance with the Guide for the Care and Use of Laboratory Animals of the University of California (Los Angeles, CA) were and approved by the Chancellor's Animal Research Committee. Twenty-seven male Lewis rats, aged 3 months, were purchased from Taconic Farms (Hudson, NY) and assigned to experimental groups as shown in Table 1. The FSD model was performed as previously described.¹² Briefly, a 25- to 30-mm longitudinal incision was aseptically made over the anterolateral aspect of the femur, and the overlying muscles were separated to expose the femoral shaft. For fixation, a polyethylene plate (length, 23 mm; width, 4 mm; height, 4

 Table 1
 Experimental Groups for FSD Surgeries

mm) was placed on the anterolateral surface of the femur and secured with four 0.9 mm-diameter threaded Kirschner wires drilled through the plate and both cortices of the femur. Two 26-gauge stainless steel cerclage wires were tightened around the plate and bone for additional stability. With a small, oscillating saw blade (Stryker, Kalamazoo, MI), a 6-mm, critical-sized, mid-diaphyseal defect was generated. The volume of the defect was approximately 75 mL. An implant was placed into the defect, and then the overlying muscle, fascia, and skin were closed with 4-0Vicryl absorbable suture (Ethicon, Somerville, NJ). Animals were sacrificed at 2 and 8 weeks after surgery.

3D Microcomputer Tomography Evaluation

After sacrifice, rat femurs were scanned by high-resolution three-dimensional (3D) microcomputer tomography (microCT) (SkyScan 1172F; Bruker microCT N.V., Kontich, Belgium) at an image resolution of 27.4 µm (55 kV and 181 μÅ radiation source, using a 0.5-mm aluminum filter) and analyzed using DataViewer, Recon, CTAn, and CTVol software version 1.15.7.0+ provided by the manufacturer (Bruker microCT N.V.). All quantitative and structural morphometric data use nomenclature described by the Nomenclature Committee of the American Society for Bone and Mineral Research.³⁸ The global thresholding principle was applied for a threshold of 100 to all samples to isolate mineralized bone for bone volume and tissue volume measurements and calculations. Two-dimensional (2D) X-ray images were obtained by the creation of an axial cut through the center of the defect area. A volume of interest of 500 slices (approximately 10 mm) centered on the middle of the defect was selected for sagittal sections and 3D reconstruction. Sagittal sectional images were generated through the mediolateral center of the femur. A volume of interest of 50 slices (approximately 1 mm) centered on the middle of the defect was reconstructed for axial sectional images. Bone volume and tissue volume measurements were calculated from analyses performed for a volume of 300 slices (approximately 6 mm).

	Number of animals	Scaffold volume, μL	Total dose, μg	
Experimental group, concentration			NELL-1	BMP-2
8 weeks				
PBS only (control)	3	75	0	0
300 μg/mL BMP2	3	75	0	22.5
600 μg/mL BMP2	3	75	0	45
600 μg/mL NELL-1 + 300 μg/mL BMP2	3	75	45	22.5
600 μg/mL NELL-1 + 600 μg/mL BMP2	3	75	45	45
2 weeks				
PBS only (control)	3	75	0	0
600 μg/mL BMP2	3	75	0	45
600 µg/mL NELL-1	3	75	45	0
600 μ g/mL NELL-1 + 600 μ g/mL BMP2	3	75	45	45

BMP2, bone morphogenetic protein 2; FSD, femoral segmental defects; NELL-1, NEL-like molecule-1; PBS, phosphate-buffered saline.

Biochemical Analysis

Finite element (biomechanical) analysis was performed using microCT images converted to DICOM files using SkyScan DICOM converter software (DicomCT application, SkyScan 1172F) according to our prior methods.³⁹ Tetrahedral 3D mesh models were generated using a random volume of interest within the defect site (size, 0.5 mm³) using ScanIP software version 7 (Simpleware Ltd., Exeter, UK). Finite element analyses were performed using the ABAQUS software version 6.12 (Dassault Systèmes, Forest Hill, MD) with boundary conditions set as encastre, constrained in all directions. Next, we applied a uniform compressive pressure of 0.5 MPa on the superior surface of the volume of interest. The von Mises stress in the samples was recorded.

Histological, Immunohistochemical, and Histomorphometric Analysis

Hematoxylin and eosin (H&E), Masson's trichrome, and immunohistochemical staining were performed as previously described.²⁶ Briefly, tissues were embedded in paraffin, sectioned into 5-µm slices, and deparaffinized and rehydrated before antibody addition. All primary antibodies were used at a dilution of 1:100. Appropriate secondary antibodies (Dako North America, Inc, Carpinteria, CA) were used at a dilution of 1:200. Photomicrographs were acquired using Olympus BX51 (×40, ×200, and ×400 magnification lens, UPLanFL; Olympus, Center Valley, PA) and SZX12 microscopes (×10 magnification lens, DF PLAPO 1.2× pf; Olympus) and a MicroFire digital microscope camera with ImageFrame software version 2.1 (Optronics, Goleta, CA). Staining intensity was quantified using ImagePro Plus (Media Cybernetics, Rockville, MD). Results are reported as the intensity measurements from eight random fields at $\times 200$ magnification.

Oil red O staining was performed on 10- μ m frozen sections using a 0.3% oil red O solution in 60% isopropyl alcohol/40% ddH₂O for 15 minutes. Slides were counterstained with hematoxylin for 15 seconds to visualize nuclei. The intensity of oil red O-positive staining was quantified using ImagePro Plus in eight separate random fields at \times 200 magnification.

Histomorphometric analyses were performed on H&E– stained sections. Only regions within the defect region containing newly formed trabecular bone were analyzed. The histomorphometric parameters of bone area, tissue area, trabecular number, trabecular perimeter, trabecular width, and trabecular spacing were measured at $\times 100$ magnification and analyzed using Photoshop CS5 (Adobe Systems, San Jose, CA) as previously described.⁴⁰ Five sections per specimen were analyzed. All histomorphometric measurements in each specimen were assessed blindly (X.Z., J.S., and K.S.L.), and all nomenclature, symbols, and units are used per established American Society for Bone and Mineral Research conventions.⁴¹

Cell Culture

The M2-10B4 cell line, a clone derived from bone marrow stromal cells (BMSCs) from a (C57BL/6J × C3H/HeJ)F1 mouse,⁴² was purchased from ATCC (Manassas, VA). Cells were maintained in growth medium [RPMI 1640 (Invitrogen, Carlsbad, CA) supplemented with 10% heat-inactivated fetal bovine serum, 1 mmol/L sodium pyruvate, and 100 U/mL penicillin/streptomycin]. Osteogenic differentiation medium consisted of RPMI 1640, 10% FBS, 50 µg/mL ascorbic acid, and 3 mmol/L β-glycerophosphate. Growth medium only was used for adipogenic differentiation studies as previously reported.⁴³

Primary human BMSCs were purchased from PromoCell (Heidelberg, Germany). Cells were cultured using Mesenchymal Stem Cell Growth Medium (C-28010; PromoCell). *In vitro* osteogenesis and adipogenesis were induced by mesenchymal stem cell osteogenic differentiation medium (Mesenchymal Stem Cell Growth Medium plus 50 µg/mL ascorbic acid and 3 mmol/L β -glycerophosphate)⁴⁴ and Mesenchymal Stem Cell Adipogenic Differentiation Medium (C-28011; PromoCell).

For osteogenic or adipogenic differentiation, M2-10B4 cells or human BMSCs were seeded at 1×10^5 cells per well or 8×10^4 cells per well, respectively, in 24-well plates for 24 hours, and then cultured with differentiation medium with NELL-1 and/or BMP2.

RNA Extraction, Real-Time PCR, and Immunoblot Analysis

Total RNA extraction and real-time PCR were performed as previously described.⁴⁵ SYBR green real-time PCR primer sequences for mouse M2-10B4 cells are given in Table 2. For human BMSCs, TaqMan primer-probe sets HS99999905_m1 (glyceraldehyde-3-phosphate dehydrogenase), Hs01115513-m1 (PPAR γ), Hs02915002-s1 (C/EBP α), Hs00609452-g1 [osteocalcin (OCN)], and Hs00959010-m1 [osteopontin (OPN)] were used (Applied Biosystems, Carlsbad, CA). All fold-change values from three independent experiments are reported.

Nuclear and cytoplasmic proteins were isolated using an NE-PER Nuclear and Cytoplasmic Extraction Kit (Thermo Scientific, Rockford, IL). Western blot analysis was performed as previously described²⁶ using antibodies against PPAR γ , C/EBP α , and β -catenin at a dilution of 1:1000.

In Vitro Staining and Alkaline Phosphatase Activity Assay

For oil red O staining of cultured cells, monolayers were rinsed with PBS, fixed with 10% formalin for 1 hour at room temperature, and stained with 3% oil red O solution (see *Histological, Immunohistochemical, and Histomorphometric Analysis*) for 10 minutes. Adipocytes were quantified through counting of oil red O—positive cells in four wells per experimental condition. Results are reported from 16 random fields at ×200 magnification. Alizarin red staining and alkaline

Gene	Forward primer	Reverse primer	
Gapdh	5'-TGCACCACCAACTGCTTAGC-3'	5'-CCACCACCTGTTGCTGTAG-3'	
Pparg	5'-ggaaagacaacggacaaatca-3'	5'-tacggatcgaaactggcaa-3'	
Cebpa	5'-tgaacaagaacagcaacgag-3'	5'-TCACTGGTCACCTCCAGCAC-3'	
Alp	5'-tgccactgtgagaagacctg-3'	5'-TGCACAGGAAGTGAGTCTGG-3'	
Runx2	5'-ccgcacgacaaccgcaccat-3'	5'-CGCTCCGGCCCACAAATCTC-3'	
0cn	5'-gcaataaggtagtgaacagactcc-3'	5'-AGCAGGGTTAAGCTCACACTG-3'	
Opn	5'-TGATGCCACAGATGAGGACCT-3'	5'-cagagggcatgctcagaagc-3'	

Table 2 Mouse Primer Sequences for Real-Time PCR

phosphatase activity assays were performed as previously described.²⁶

Immunocytochemistry

M2-10B4 cells were seeded on Millicell EX Slides (PEZGS0816, Millipore) at 5×10^4 cells per well in RPMI 1640 + 10% FBS for 24 hours and serum starved in RPMI 1640 + 1% FBS overnight. After 2 hours of treatment, cells were fixed using ice-cold acetone for 10 minutes. Anti–active β -catenin antibody (Millipore) was applied at a dilution of 1:200. ABC complex (Vector Laboratories, Burlingame, CA) was applied to the sections after incubation with biotinylated secondary antibody (Dako). AEC Substrate (Dako) was used as a chromogen, and the sections were lightly counterstained with hematoxylin. Photomicrographs were acquired using an Olympus BX51 device (×200 UPLanFL magnification lens).

Luciferase Reporter Assay

M2-10B4 cells were transfected with 20 µg of Super(16×) TOPFLASH (TCF/LEF reporter plasmid) and 1 µg of *Renilla* luciferase plasmids for 24 hours and then seeded at 4.0×10^3 cells per well in 96-well plates. Cells were starved in RPMI 1640 + 1% FBS overnight and then treated with BMP2 and/ or NELL-1. Luciferase activity was measured 48 hours after treatment using Dual Luciferase Reporter Assay System (Promega, Madison, WI) per the manufacturer's instructions.

Preparation of Lentiviral Vectors for Runx2 Reporter Assays

The transduction plasmid for preparing lentiviral vector carrying the *Runx2* P1-enhanced green fluorescent protein (EGFP) expression cassette (*Runx2*-EGFP reporter) was prepared by substituting the cytomegalovirus promoter in the pRRL-cPPT-CMV-X-PRE-SIN plasmid⁴⁶ with the mouse *Runx2* P1 promoter. The mouse *Runx2* P1 promoter was obtained by PCR of mouse genomic DNA (forward primer, 5'-GCGAATTACTCGAGAGCAGCAGCACTGTTGC-TCAGAA-3'; reverse primer, 5'-GCGAATGCCCGGG-TCACACAATCCAAAAAAGC-3'). 293T cells were cotransfected with the transduction plasmids, the package plasmid pCMV-dR8.2-vprX, and the envelope plasmid

pCMV-VSVG.⁴⁷ The viral vectors were collected at 2 to 4 days after transfection and were filtered, and concentrated. The concentrations of viral vector were quantified by a count of the core protein p24 by enzyme-linked immuno-sorbent assay as previously described.⁴⁷

Cell Infection

M2-10B4 cells were seeded in 24-well cell culture plates at 2×10^4 cells per well 16 hours before infection. Viral vectors with p24 counts of 0.4 µg were added to each well in 24-well plates. Three hours after infection, the viral vectors were washed away, and fresh medium was added to the cultures. At 24 hours after infection, BMP2 and/or NELL-1 were added into the culture media.

Flow Cytometric Assay of GFP-Positive Cells

Cells were trypsinized and collected 3 days after infection. Flow cytometry was performed to quantify GFP expression in the collected cells using a Cytomics FC500 cell sorter (Beckman Coulter, Brea, CA). Cells infected with mock vector were used as the negative control to establish gates. The percentages of GFP-positive cells were counted to quantify the expression of GFP in the infected cultures.

Statistical Analysis

Means \pm SDs were calculated from numerical data. To allow for multiple comparisons between groups, statistical analyses were performed using one-way analysis of variance for singlevariable comparisons and two-way analysis of variance for dual-variable comparisons (ie, experiments involving both Dkk or XAV939 and BMP2 and/or NELL-1). This procedure was followed by a *post hoc* Tukey range test to directly compare the data between the two groups. P < 0.05 was considered to be significant for all statistical tests used.

Results

NELL-1 Inhibits BMP2-Induced Cystlike Bone Formation *in Vivo*

Although the 300 and 600 $\mu g/mL$ BMP2 concentrations used here are less than the 1.5 mg/mL concentration used

clinically, these concentrations are sufficient for reproducible induction of abnormal bone formation in a rat FSD model, as we have previously described.¹² In addition, we used NELL-1 at 600 μ g/mL because we previously showed that this NELL-1 concentration in a demineralized bone matrix carrier was able to induce fusion in an FSD model.³⁷ PLGA scaffolds containing PBS, BMP2, or NELL-1+BMP2 combined, were implanted into rat FSDs (Figure 1A).

We performed high-resolution 2D radiography and 3D microCT reconstructions of the defect sites 8 weeks after surgery. Control defects treated with PBS exhibited definitive non-union. In contrast, BMP2 treatment alone resulted in bone fusion, but the newly formed bone was characterized

peripherally by a thin, continuous cortical shell that extended well beyond the original bony defect margins and internally by sparse and thin trabeculae formation. Strikingly, NELL-1+BMP2 cotreatment produced bone fusion, with newly formed bone confined mostly to the defect sites with considerably thicker cortical architecture and a much denser, native bonelike, trabecular structure compared with the BMP2-only group (Figure 1B). Notably, NELL-1 on PLGA scaffolds increased bone regeneration compared with that in controls, but did not induce bony fusion at up to 12 weeks after surgery (Supplemental Figure S1), unlike NELL-1 on demineralized bone matrix scaffolds.³⁷

On quantitative microCT analysis of bone volume and tissue volume (Figure 1, C and D, respectively), all



Figure 1 Microcomputer tomography (microCT) and biomechanical evaluation of femoral segmental defects (FSDs) at 8 weeks after surgery reveals induction of cystlike bone formation with bone morphogenetic protein 2 (BMP2) alone and suppression of this effect by addition of NEL-like molecule-1 (NELL-1). BMP2 dose: 300 or 600 μ g/mL. NELL-1 dose: 600 μ g/mL. **A: Left panel**, schematic of FSD surgical procedure; **middle panel**, hydroxyapatite-coated PLGA scaffold implanted into defect; **right panel**, intraoperative photograph after scaffold implantation. **B**–**D:** From top to bottom, two-dimensional (2D) radiography, sagittal section, axial section, and three-dimensional (3D)-reconstructed microCT images (**B**). **Green boxes** indicate the region of interest selected for microCT reconstruction below. BMP2 treatment results in formation of abnormal bone extending beyond the cortical margins, which is not observed with NELL-1+BMP2 co-treatment. **Red boxes** indicate the region of interest selected for quantification of bone volume (**C**) and tissue volume (**D**). **E:** Finite element analysis and quantification of von Mises stress. Data are expressed as means \pm SD. **P* < 0.05, ***P* < 0.01 versus control group; [†]*P* < 0.05 versus same dose of BMP2. *n* = 3 (**C** and **D**). Scale bar = 5 mm (**B**).

treatments significantly increased bone volume compared with no treatment; however, no statistically significant difference in bone volume between the BMP2-only and NELL-1+BMP2 cotreatment groups was observed. On the other hand, NELL-1+BMP2 cotreatment significantly reduced tissue volume compared with BMP2 treatment alone. Moreover, computer-simulated biomechanical (finite element analysis) testing demonstrated increased bone strength with BMP2+NELL-1 cotreatment compared with BMP2 only (Figure 1E). Taken together, these measurements indicate that similar amounts of bone were induced by either BMP2 treatment alone or NELL-1+BMP2 cotreatment, but the bone was distributed throughout a larger volume of tissue with BMP2-only treatment compared with NELL-1+BMP2 cotreatment.

We next analyzed the bone quality of FSDs treated with PBS only (control), BMP2 alone, or NELL-1+BMP2 by histological examination of specimens collected at 8 weeks



Figure 2 Histological evaluation at 8 weeks after surgery reveals increased osteogenesis with NEL-like molecule-1 (NELL-1) + bone morphogenetic protein 2 (BMP2) cotreatment and reduction of adipogenesis induced by BMP2. BMP2 dose, $600 \mu g/mL$; NELL-1 dose, $600 \mu g/mL$. **A:** Hematoxylin and eosin (H&E) staining reveals a large extension of bone surrounded by thin cortical bone in BMP2-only samples, which is not observed in phosphate-buffered saline control or NELL-1+BMP2 co-treated samples. The **boxed areas** indicate the regions magnified in **B**. Boundaries of defect are as indicated. Histomorphometric analyses are based on H&E staining. **B:** Masson's trichrome staining reveals fibrous tissue (light blue) in control samples, abundant adipocytes (large white droplets) in BMP2-only samples, and osteoid matrix (dark blue) ossifying into mature trabecular bone (red) in NELL-1+BMP2 co-treated samples. **C:** Osteocalcin (Ocn) immunohistochemical analysis. **Arrowheads** indicate positive Ocn staining and its quantification. **D:** Oil red O staining and its quantification. Data are expressed as means \pm SD. **P* < 0.05, ***P* < 0.01 versus control group; ^{††}*P* < 0.01 versus same dose of BMP2. *n* = 3 (**B** and **D**). Scale bars: 4 mm (**A**), 1 mm (**B**), 0.2 mm (**C**), 0.4 mm (**D**). Def, extent of defect area; Tb. N, trabecular number; Tb. Pm., trabecular perimeter; Tb.Sp, trabecular spacing; Tb.Wi, trabecular width.

after surgery (Figure 2). The control defects exhibited definitive fibrous non-union of the femoral cortices, with no trabecular bone formation at the defect centers. BMP2 treatment alone induced fracture union, but the newly formed bone contained exuberant adipose tissue interspersed with sparse trabecular bone that was surrounded by a thin cortical bone shell extending up to approximately 4 mm beyond the original bone defect margins. In contrast, NELL-1+BMP2 cotreatment generated tightly woven, trabecular bone largely confined to the defect sites. Histomorphometric analyses on serial H&E-stained sections showed increased bone area/tissue area ratio, trabecular number, perimeter, and width, and reduced trabecular spacing in NELL-1+BMP2 treated samples. Masson's trichrome staining and immunostaining for osteocalcin (Ocn), a marker of bone matrix and osteoblastic differentiation, confirmed that NELL-1+BMP2 cotreated defects exhibited increased bone formation with denser trabeculations than with BMP2 treatment alone. To specifically assess and quantify adipogenic differentiation observed by H&E and Masson's trichrome histological examination, we performed oil red O staining on cryosectioned specimens. Consistent with our previous histological findings,¹² BMP2 treatment alone significantly increased the intensity of oil red O staining, an effect notably absent with NELL-1+BMP2 cotreatment.

To gain better insight on the early cellular effects of NELL-1 and BMP2 on defect healing, we evaluated NELL-1, BMP2, or NELL-1+BMP2 treated FSDs at 2 weeks after surgery (Figure 3). Our published findings show that Nell-1 does not form ectopic bone,²⁶ but that Nell-1 does form robust bone growth when in the

presence of an osteogenic cell source such as BMSCs.¹¹ In agreement with our previous studies, NELL-1 treatment alone did not result in inappropriate expansion of bone tissue into the muscle compartment.¹¹ Instead, new bony trabeculations arising from the cut femoral bone ends were observed growing into the defect. In contrast, BMP2-only samples exhibited a massive reactive tissue zone characterized by increased cellularity in the muscle and bone defect compartments that extended well beyond the original margins of the bone defect site. Meanwhile, NELL-1+BMP2 cotreatment induced a much smaller reactive tissue zone, with increased trabecular bone largely confined to the original bone defect area, and with minimal tissue reactivity in the muscle compartment. Together, our microCT and histological data indicate that NELL-1 reduces BMP2-induced exuberant bone formation and adipogenesis to improve BMP2-mediated osteogenesis.

NELL-1 Reduces BMP2-Induced Adipogenesis

We previously observed that goat BMSCs transduced with adenoviral *BMP2* and injected into nude mouse muscle pouch models form large bone voids filled with fatty tissue.¹¹ This phenomenon was also observed with BMP2 use in dog models and in humans.^{27,28} Furthermore, forced *Nell1* and *Bmp2* overexpression by adenoviral treatment produces more bone than either adenoviral *Nell1* or *Bmp2* alone in a nude mouse muscle pouch model.²⁶

To further analyze the effects of NELL-1 on BMP2induced adipogenesis, we focused on the expression of Ppar γ and C/Ebp α , two important transcriptional regulators



Figure 3 Histological evaluation at 2 weeks after surgery. **A:** Hematoxylin and eosin (H&E) staining demonstrates increased cellularity and aberrant expansion of bone tissue into the muscle compartment with bone morphogenetic protein 2 (BMP2) treatment alone as early as 2 weeks, but not with NEL-like molecule-1 (NELL-1) treatment alone or NELL-1+BMP2 cotreatment. The **boxed areas** indicate the regions magnified in **B. Arrowheads** indicate the lateral limit of new tissue formation. BMP2 dose: 600 μ g/mL; NELL-1 dose, 600 μ g/mL. **B:** Trichrome staining for mature mineralized bone (red) and osteoid matrix (blue). Please note that the implanted scaffold also appears red. In the defect site, NELL-1+BMP2 cotreatment induces more trabecular bone than does BMP2 treatment alone, but BMP2 alone also increases cellularity and induces a tissue reaction containing poorly trabeculated bone and fibrous tissue outside of the cortical margin of the defect. The same location in the other groups contains primarily muscle tissue only. n = 3 (**B**). Scale bars: 4 mm (**A**), 1 mm (**B**). Def, extent of defect area.

of adipogenesis^{48,49} (Figure 4). PPAR γ immunohistochemistry of the increased adipogenesis observed by histological examination was used for analyzing FSD specimens collected 2 weeks after surgery. Samples treated with BMP2 alone exhibited strong Ppary staining in the cells surrounding and between the trabeculae at the defect center, consistent with increased adipogenesis. Importantly, NELL-1 significantly repressed BMP2-induced Ppary expression in NELL-1+BMP2 cotreated animals. We next sought to replicate these findings in vitro using the mouse BMSC line M2-10B4 under adipogenic conditions, as previously described.⁴³ At 9 days, BMP2-induced lipid accumulation was increased more than fivefold, as assessed by oil red O staining. However, the addition of NELL-1 antagonized the proadipogenic effects of BMP2 and significantly reduced both Ppary and C/Ebpa expression. Because the effects of BMP2 on BMSCs may be species dependent,⁵⁰ and because cell lines may respond differently from primary cells, we also confirmed the inhibitory effects of NELL-1 on BMP2-induced adipogenesis using primary rat BMSCs (Supplemental Figure S2A) and primary human BMSCs.

These data confirm that NELL-1 inhibits BMP2-stimulated adipogenesis *in vitro* in BMSCs from multiple species.

NELL-1 Enhances BMP2-Induced Osteogenesis

To determine whether the suppression of adipogenesis corresponded with an increase in osteogenesis, we analyzed Runx2 expression in the centers of FSDs collected 2 weeks after surgery (Figure 5A). Runx2 expression was signifiwith NELL-1+BMP2 cantly increased cotreatment compared with BMP2 treatment alone. To model this effect in vitro, M2-10B4 cells were cultured under osteogenic conditions (Figure 5, B and C). As expected, NELL-1 and BMP2 alone increased the expression of markers of early (Runx2 expression), intermediate (alkaline phosphatase and osteopontin), and terminal osteogenic differentiation (osteocalcin expression and alizarin red-positive bone nodules) over those in controls. However, when added together, NELL-1+BMP2 significantly increased all osteogenic markers relative to NELL-1 or BMP2 alone, consistent with our 8-week imaging and histological findings of



Figure 4 Addition of NEL-like molecule-1 (NELL-1) reduces bone morphogenetic protein 2 (BMP2)-induced adipogenesis. **A:** Peroxisome proliferatoractivated receptor γ (PPAR γ) immunohistochemical analysis of femoral segmental defects (FSDs; within the scaffold implant) at 2 weeks after surgery reveals increased expression induced by BMP2 treatment alone but not NELL-1+BMP2 cotreatment. BMP2 dose, 600 µg/mL; NELL-1 dose, 600 µg/mL. Quantitation of PPAR γ staining is show in graph to the right. **B**–**D**: M2-10B4 cells were treated with phosphate-buffered saline (PBS) only (control), 800 ng/ mL NELL-1, and/or 300 ng/mL BMP2 under adipogenic differentiation. The expression of PPAR γ and CCAAT enhancer—binding protein α (C/EBP α) was determined on day 3 by real-time PCR (**B**) and Western blot analysis with quantification of PPAR γ and C/EBP α (**C**). **D:** Lipid accumulation was visualized by oil red 0 staining at 9 days and quantified by light microscopy from eight random areas. **E:** Real-time PCR analysis of *PPARG* and *CEBPA* expression in human primary bone marrow stromal cells (BMSCs) treated with PBS, 800 ng/mL NELL-1, and/or 300 ng/mL BMP2 for 2 days. All fold-changes are reported relative to control group. Data are expressed as means \pm SD. **P* < 0.05, ***P* < 0.01 versus control group; [†]*P* < 0.05, ^{††}*P* < 0.01 versus same dose of BMP2. *n* = 3 (**A**, **C**, and **E**), *n* = 4 (**D**). Scale bars = 0.2 mm (**A** and **D**).



Figure 5 Addition of NEL-like molecule-1 (NELL-1) enhances bone morphogenetic protein 2 (BMP2)-induced osteogenesis. **A:** Expression of runt-related transcription factor 2 (Runx2) in NELL-1 and BMP2-treated samples of femoral segmental defect (FSD; within the scaffold implant) at 2 weeks after surgery. BMP2 dose, 600 μ g/mL; NELL-1 dose, 600 μ g/mL. Runx2-positive cells appear primarily adjacent to trabecular bone. Quantification of Runx2 staining is shown in graph to the right. **B** and **C:** M2-10B4 cells were treated with phosphate-buffered saline, 800 ng/mL NELL-1, and/or 100 ng/mL BMP2 under osteogenic differentiation conditions. **B:** Real-time PCR analysis of osteogenic differentiation marker gene. *Runx2* expression was assessed at 1, 3, and 6 days; *ALP* expression was assessed at 3, 6, and 9 days; *Ocn* and *Opn* expressions were analyzed at 6, 9, and 12 days. **C:** Mineralization was visualized by Alizarin red staining at 12 days and quantified by light microscopy from eight random areas. **D:** Real-time PCR analysis of *OCN* and *OPN* expressions in human primary bone marrow stromal cells (BMSCs) treated with NELL-1 and/or BMP2 for 6 days. All fold-changes are reported relative to control group. Data are expressed as means \pm SD. **P* < 0.05, ***P* < 0.01 versus control group; †*P* < 0.05, ††*P* < 0.01 versus same dose of BMP2. *n* = 3 (**A**, **B**, and **D**), *n* = 4 (**C**). Scale bars = 0.2 mm (**A** and **C**).

high-quality bone formation. Repeating these experiments in primary human BMSCs (Figure 5D) showed that NELL-1 significantly increased OCN and OPN expression in human BMSCs in combination with BMP2.

NELL-1 Induces Canonical Wnt Signaling

Because NELL-1 was previously found to activate canonical Wnt signaling in both MSC and osteoclastic cell types,³⁹ we hypothesized that NELL-1-mediated effects may involve canonical Wnt activation. We first examined *in vitro* activation of canonical Wnt signaling by immunocytochemical staining of M2-10B4 cells using an antibody against active β -catenin (Figure 6A). BMP2 did not induce β -catenin nuclear accumulation at either 100 ng/mL or 300 ng/mL, whereas NELL-1+BMP2 cotreatment stimulated more β -catenin nuclear accumulation compared with BMP2 alone. Next, Western blot analysis of M2-10B4 cytoplasmic and nuclear fractions (Figure 6B) showed significantly increased nuclear β -catenin levels after treatment with NELL-1+BMP2 compared with BMP2 only. Similarly, the TOPFlash canonical Wnt reporter assay demonstrated no luciferase activity induced by BMP2, and restored luciferase activity induced by NELL-1+BMP2 (Figure 6C). Next, we examined *in vivo* activation of Wnt signaling by immunohistochemical staining for β -catenin (Figure 6D) in the same FSD specimens used for Ppar γ (Figure 4A) and Runx2 (Figure 5A) staining. Increased nuclear β -catenin localization was observed only in NELL-1+BMP2 cotreatment groups. The BMP2-alone group revealed primarily cytoplasmic β -catenin staining. Taken together, these *in vitro* and *in vivo* results demonstrate a novel NELL-1 function to induce canonical Wnt activity and suggest that NELL-1 regulation of BMP2-induced osteogenesis and adipogenesis may occur through activation of canonical Wnt signaling.

Canonical Wnt Signaling Is Required for NELL-1 Effects on BMP2-Induced Lineage Differentiation

To determine whether NELL-1 repression of BMP2-induced *PPARG* and *CEBPA* expression requires Wnt activity, we treated M2-10B4 cells with two specific inhibitors of canonical Wnt signaling: recombinant mouse Dkk1and the

small molecule XAV939 (Figure 6E). Dkk1 functions as an antagonist of canonical Wnt signaling by binding to lowdensity lipoprotein receptor—related protein (Lrp)5/6 and preventing its interaction with Wnt-Frizzled complexes.⁵¹ In contrast, XAV939 antagonizes Wnt signaling by stabilizing Axin, an inhibitory Wnt pathway component.⁵² By itself, each Wnt inhibitor increased expression of *Pparg* and *Cebpa* when added BMP2 alone. Importantly, NELL-1 suppression of BMP2-induced *Pparg or Cebpa* expression was largely reversed by Wnt signaling inhibition.

To determine whether Wnt signaling is required for NELL-1 augmentation of BMP2-mediated osteogenesis, we added DKK1 to NELL-1+BMP2-treated M2-10B4 cells. DKK1 eliminated the synergistic increase in Runx2 expression induced by NELL-1 addition to BMP2 (Figure 6F). M2-10B4 cells were also transduced with a lentiviral vector containing the Runx2 P1 promoter driving Gfp expression. Treatment of the transduced cells with BMP2 or BMP2+NELL-1 increased Runx2 P1-driven Gfp expression. Moreover, when XAV939 was added, significant inhibition of Runx2 P1 expression was found in NELL-1+BMP2 co-treated cells (Figure 6F). Collectively, these data indicate that NELL-1 regulation of Runx2, PPARy, and C/EBPa, key transcription factors induced by BMP2 that dictate osteogenic and adipogenic lineage specification, is dependent on intact canonical Wnt signaling.

Discussion

In this study, we identified NELL-1 as a novel Wnt regulator that can suppress adipogenesis and promote osteogenesis in the context of the high BMP2 doses used clinically. We also demonstrated the ready feasibility of combining NELL-1 with BMP2 to significantly improve the clinical safety and efficacy of BMP2-based bone regeneration.

Proposed Molecular Mechanisms of NELL-1 and BMP2 Effects on Osteogenesis

The ability of NELL-1 to improve BMP2-induced bone quality can be explained in terms of their mechanistic effects on the master osteogenic and adipogenic regulators Runx2 and PPAR γ , respectively. BMP2 induces osteogenic differentiation of mesenchymal progenitor cells by activating Runx2 expression through the action of the Smad family of transcription factors.⁵³ This agrees with the well-known ability of BMP2 to induce ectopic and orthotopic bone.⁵⁴ However, high-dose BMP2 also activates PPAR γ and C/EBP α via Smad signaling^{55,56} and promotes adipogenesis *in vitro* and *in vivo*.^{12,57} Importantly, increased PPAR γ expression antagonizes Runx2, which in turn inhibits the osteogenic differentiation of mesenchymal progenitor cells, resulting in inconsistent bone formation.^{31,57,58}

NELL-1 can function as a downstream mediator of Runx2 because *NELL1* expression is directly regulated by

Runx2 binding to its promoter region⁵⁹ and *Nell1* overexpression can partially rescue *Runx2* haploinsufficiency.⁸ In addition, NELL-1 regulates Runx2 bioactivity by enhancing its phosphorylation.⁹ Co-immunoprecipitation and surface plasmon resonance experiments indicate that NELL-1 does not physically interact with BMP2 (data not shown), and we and others have established that NELL-1 does not participate in the Smad1/5/8 pathways (Supplemental Figure S2B),⁹ suggesting that NELL-1 activates Runx2 via a mechanism independent of the BMP/ Smad pathways.

In the present study, we describe NELL-1 stimulation of Runx2 expression and activity via canonical Wnt signaling, which is essential for differentiation of progenitor cells into osteoblasts and normal fetal bone development.^{60,61} Endogenous Wnt signaling is also increased in areas of bone healing, and Wnt-filled liposomes can accelerate bone healing.²⁹ In osteoblast precursor cells, canonical Wnt signaling directly stimulates the Runx2 P1 promoter to drive osteoblast differentiation.⁶⁰ Because we showed that NELL-1 significantly increased the nuclear accumulation of β catenin, it is possible that Runx2 expression may be likewise stimulated to induce osteoblastogenesis. Thus, NELL-1 is not only regulated by Runx2 and acts as its downstream mediator, but NELL-1 may also regulate Runx2 via canonical Wnt signaling. In addition to transcriptional regulation by BMP2, Runx2 physically interacts with Smad proteins to form active transcriptional complexes for osteogenic gene expression, resulting in the synergy between Runx2 and BMP2.^{62,63} Therefore, we hypothesize that NELL-1 may augment BMP2/Smad osteogenesis by stimulating canonical Wnt signaling and promoting Runx2.

Proposed Molecular Mechanisms of NELL-1 and BMP2 Effects on Adipogenesis

Besides up-regulating osteogenesis, canonical Wnt signaling is widely reported to inhibit adipogenesis in diverse cell types.^{64,65} NELL-1 reduction of PPAR γ expression was blocked by canonical Wnt inhibitors, indicating that NELL-1 down-regulation of adipogenesis was dependent on intact Wnt signaling. It was recently reported that BMP2 represses Wnt signaling during bone repair in a 1-mm in vivo monocortical trephine defect model.⁶⁶ In contrast, our data show lack of BMP2 induction, rather than repression per se, of canonical Wnt signaling. This discrepancy may be due to differences in defect size (ie, 6mm full-thickness FSD versus 1-mm monocortical trephine defect) and origin of bone-forming progenitor cells (ie, bone marrow, periosteum, or surrounding soft tissues). Irrespectively, the up-regulation of Wnt by NELL-1 can explain the ability of NELL-1+BMP2 cotreatment to increase osteogenesis while simultaneously decreasing adipogenesis. Collectively, NELL-1 promotes BMP2 osteogenesis by both up-regulating Runx2 and down-regulating PPARy via canonical Wnt signaling.



NEL-like molecule-1 (NELL-1) induces canonical Wnt pathway activity in vivo and in vitro. A: M2-10B4 cells were treated with indicated doses Figure 6 of bone morphogenetic protein 2 (BMP2) or BMP2+NELL-1 in RPMI 1640+1% fetal bovine serum for 2 hours and stained for active β -catenin. Recombinant Wnt-3a (100 ng/mL) was used as a positive control. B: Western blot analysis of nuclear and cytoplasmic fractions of β-catenin 2 hours after treatment under the same conditions as A. Quantitation of nuclear and cytoplasmic β-catenin shown in graphs to the middle and right. C: M2-10B4 cells were transfected with Super (16×) TOPFlash and Renilla luciferase plasmids. Cells were treated with indicated doses of BMP2 or BMP2+NELL-1 for 48 hours. The luciferase activity was normalized to control. **D:** Immunohistochemical analysis for β -catenin expression in BMP2 or BMP2+NELL-1 treated samples of femoral segmental defect (within the scaffold implant) at 2 weeks after surgery. BMP2 dose, 600 µg/mL; NELL-1 dose, 600 µg/mL. Arrowheads highlight positive nuclear staining for β -catenin. Quantitation of β -catenin staining is show in graph to the right. Insets and boxed areas indicate areas of magnification. E: Intact canonical Wnt signaling is required for anti-adipogenic and pro-osteogenic effects of NELL-1. M2-10B4 cells were seeded at the density of 1×10^5 cells per well in 24-well plates for 24 hours and treated with phosphate-buffered saline, 300 ng/mL BMP2, or 300 ng/mL BMP2 + 800 ng/mL NELL-1 under adipogenic differentiation conditions with or without 100 ng/mL Dickkopf-related protein 1 (DKK1) or with or without 1 µmol/L XAV-939 for 3 days. Peroxisome proliferator-activated receptor γ (PPAR γ) and CCAAT enhancer-binding protein α (C/EBP α) expression was measured by realtime PCR. F, top graph: M2-10B4 cells were seeded at the density of 1×10^5 cells per well in 24-well plates for 24 hours and treated with PBS, 300 ng/mL BMP2, or 300 ng/mL BMP2 + 800 ng/mL NELL-1 under osteogenic differentiation conditions with or without DKK1 for 3 days. Runx2 expression was measured by real-time PCR. Bottom graph: M2-10B4 cells were transduced with Runx2-enhanced green fluorescent protein reporter lentivirus and treated with PBS, 300 ng/mL BMP2, or 300 ng/mL BMP2 + 800 ng/mL NELL-1 with or without 1 µmol/L XAV939 24 hours after transduction. Runx2 reporter assay was performed on transduced cells after 3 days. All fold changes are reported relative to control group. Data are expressed as means \pm SD. *P < 0.05, **P < 0.01 versus control group; $^{\dagger}P < 0.05$, $^{\dagger\dagger}P < 0.01$ versus same dose of BMP2; $^{\ddagger}P < 0.05$, $^{\ddagger}P < 0.01$ versus no inhibitor treatment. n = 3 (A, B, D, and F), n = 4 (C). Scale bars = 0.1 mm (A; D, insets); 0.04 mm (D, boxed area).

Previously we identified NELL-1–integrin- β_1 binding as essential for the NELL-1–mediated activation of Wnt signaling.⁶⁷ In this study, we showed that Dkk1 blocked NELL-1 regulation of BMP2–induced osteogenesis and adipogenesis. Because DKK1 antagonizes canonical Wnt signaling by binding to LRP5/6, preventing its interaction with the Wnt-Frizzled complex,⁵¹ our data suggest that NELL-1 stimulates canonical Wnt by extracellular mechanisms. The small molecule XAV939, which blocks Wnt signaling by intracellular mechanisms, also abrogated the canonical Wnt response to NELL-1, further verifying the function of NELL-1 in this pathway. Our ongoing study of the mechanisms of NELL-1 activation of Wnt signaling, via integrin- β_1 or other cell surface receptors, will greatly improve our mechanistic understanding of NELL-1 functions.

Cellular Effects of NELL-1 and BMP2 in Femoral Segmental Defect Healing

Controlling the proper differentiation of local progenitor cells in the defect space is a highly regulated process crucial for normal bone development, maintenance, and healing after injury.⁶⁸ At the cellular level, BMP2 may expand the pool of mesenchymal progenitors by inducing local cell proliferation,⁶⁹ reducing apoptosis of local cells,⁷⁰ or chemotactically recruiting mesenchymal progenitors from the bone marrow or neighboring compartments such as the periosteum and the surrounding muscle and soft tissues.⁷¹ These mechanisms all increase the number of cells at the defect site and may contribute to the reactive tissue zone induced by clinically relevant, high-dose BMP2¹² (Figure 1). Indeed, we found increased proliferation and decreased apoptosis on BMP2 treatment (Supplemental Figure S3), correlating with the increased cellularity induced by BMP2 (Figure 3). Another interesting aspect is the precise cellular constituent that was responsive to NELL-1 and BMP2 in our model. A femoral bone defect environment will expose multiple, distinct cell types to exogenous BMP2 and NELL-1 protein, including BMSCs, endosteal/periosteal progenitor cells, and cortical osteoblasts. Prior studies from our research group have confirmed that multiple cell types are NELL-1 responsive. For example, NELL-1 induction of osteogenesis is best studied in BMSCs³⁹ but also has been reported in committed osteoblasts,⁶⁷ adipose-derived stem cells,⁷² and perivascular stem cells.^{73,74} Although complex, Understanding the cellspecific responses to NELL-1 and BMP2 treatment may allow for future refinements in protein-based methods of tissue engineering.

Here, NELL-1 alone, delivered from a synthetic PLGA carrier, resulted in limited osteogenesis at the defect margins (Supplemental Figure S1), a location adjacent to the periosteum and bone marrow, which are sources of BMPs and BMSCs.^{75,76} In contrast, we have shown that NELL-1 treatment alone can induce complete defect healing using a demineralized bone matrix carrier, which contains residual BMPs.^{37,77} Finally, using a femoral onlay model,¹² we found

that NELL-1 does not mediate inflammatory cell infiltration, in contrast to BMP2.⁷⁸ Based on these observations, we believe that NELL-1 is not directly chemotactic, but instead promotes a more directed lineage specification of BMP2induced or recruited progenitors toward osteogenesis, while at the same time inhibiting adipogenic lineage commitment. Indeed, we observed reduced cellularity (Supplemental Figure S3) and adipogenesis (Figures 2 and 3) with NELL-1+BMP2 cotreatment compared with BMP2 treatment alone. Just as BMPs have differing effects depending on cell subpopulation,⁶⁶ an important future direction will be to identify the precise cell populations targeted by NELL-1 *in vivo* to further improve its osteoinductive potential.

Potential Applications for NELL-1 in the Clinic

Clinically, osteoinduction in humans requires high-dose BMP2, which elicits significant adverse effects (http://www. fda.gov/MedicalDevices/Safety/AlertsandNotices/PublicHealth Notifications/ucm062000.htm).²²⁻²⁵ The BMP2 doses used in this study were based on those used clinically¹⁷; lesser BMP2 doses are not osteoinductive in humans.⁷⁹ Based on our previous data,¹² low BMP2 doses (<150 µg/mL) induce osteogenesis and bone healing in rat FSDs but do not elicit adipogenic effects. We did find that treatment groups receiving lesser doses of BMP2 supplemented with NELL-1 achieved radiographic scores greater than those from groups receiving BMP2 alone (data not shown). Because NELL-1 can improve bone formation even if lesser BMP2 doses are used, an interesting future direction would be to test whether the combination of NELL-1 with BMP2 can reduce the currently required clinical dose of BMP2, to potentially reduce the adverse effects of high-dose BMP2.

Because NELL-1 alone significantly inhibits PPARy expression (Figure 4), it may also have an inherent suppressive effect on adipogenesis independent of BMP2-induced adipogenesis.⁸⁰ Because canonical Wnt inhibitors did not completely eliminate the antiadipogenic effect of NELL-1 on BMP2-induced adipogenesis, we hypothesize that other pathways, such as Hedgehog signaling, may be involved in adipogenic regulation.⁸¹ The ability of NELL-1 to activate Wnt signaling suggests potential utility in conditions such as osteoporosis, in which the balance between osteogenesis versus adipogenesis and the balance between bone deposition versus resorption are perturbed to favor bone loss.^{31,82–85} In fact, in recognition of the importance of Wnt signaling in bone disease in humans, antibodies against two endogenous Wnt pathway inhibitors, sclerostin and DKK-1, are currently in Phase 2 clinical trials in osteoporosis.⁸⁵ Additionally, ligand activators of PPAR γ , such as the antidiabetic drug class thiazolidinediones, induce adipogenesis and osteoclastogenesis,⁸⁶ whereas *PPARG* haploinsufficiency increases osteoblastogenesis over adipogenesis.⁸⁷ Meanwhile, Wnt signaling promotion can decrease osteoclastogenesis.⁸⁸ Thus, NELL-1 stimulation of Wnt signaling may not only repress PPARy to increase osteoblastogenesis but also reduce osteoclastogenesis and associated bone loss.^{24,89} In support of this concept, mice homozygous for *Nell1* deficiency exhibit reduced expression of the osteoclastogenesis inhibitor osteoprotegerin,⁶ and the heterozygous *Nell1*-deficient mice that survive to adulthood manifest an osteoporotic phenotype.³⁹ Interestingly, BMP2 and Wnt signaling synergistically increase osteoprotegerin expression,⁹⁰ and ongoing studies will determine whether BMP2+NELL-1 can similarly induce osteoprotegerin and/or inhibit osteoclastogenesis.

Conclusions

Taken together, the findings from our previous^{12–14,26,37} and present studies demonstrate a strong mechanistic rationale for combining NELL-1 with BMP2 to significantly improve the safety and efficacy of the currently clinically available bone-regeneration capabilities. The combination treatment of NELL-1 with BMP2 may be particularly valuable in clinical scenarios in which bone regeneration is impaired, such as in patients undergoing corticosteroid treatment or having osteoporosis. In addition, the novel ability of NELL-1 to stimulate Wnt signaling and to repress adipogenesis may highlight new treatment approaches for osteoporotic bone loss that aim to both increase osteoblastogenesis and decrease osteoclastogenesis.

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Supplemental Data

Supplemental material for this article can be found at *http://dx.doi.org/10.1016/j.ajpath.2015.10.011*.

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