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#### Identification of Gene Signature Associated with Elevated Bone Formation Rate in Aging Mice

Krista Jackson Marian University - Indianapolis, kjackson466@marian.edu

Aaron M. Hudnall Marian University - Indianapolis, ahudnall110@marian.edu

Jonathan W. Lowery Ph.D. Marian University - Indianapolis, jlowery@marian.edu

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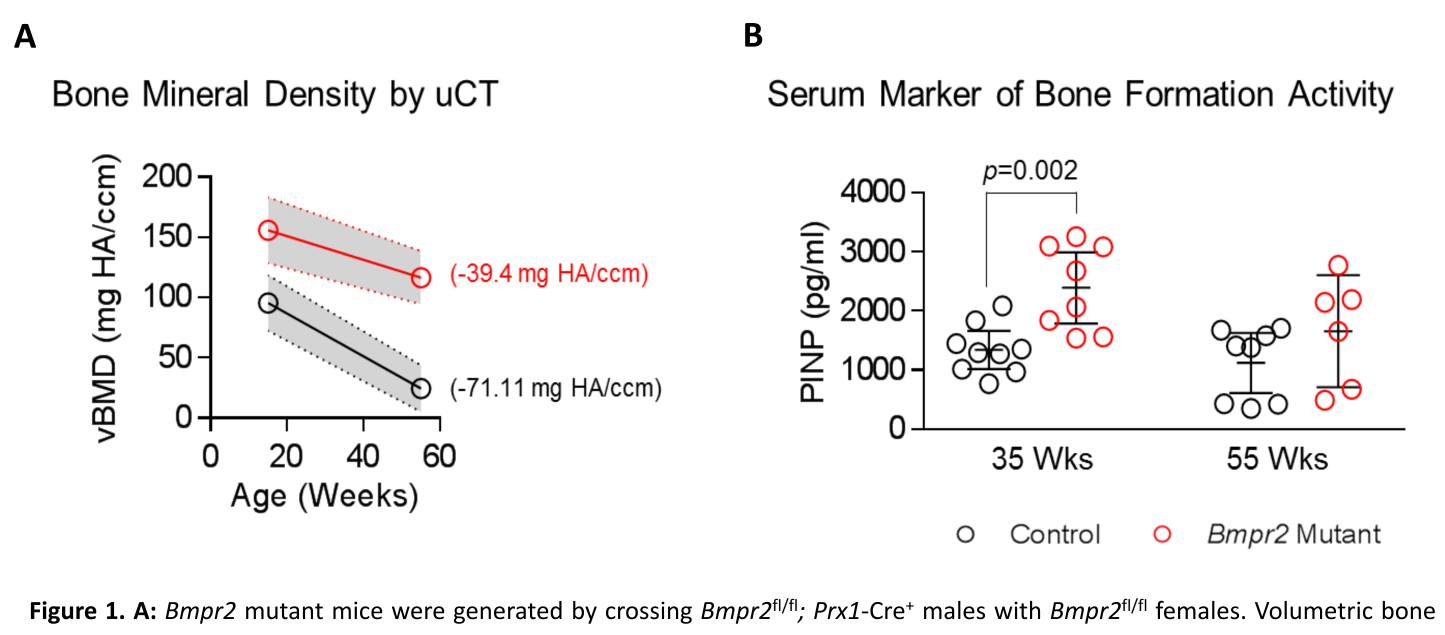
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# Identification of a gene signature associated with elevated bone formation rate in aging mice

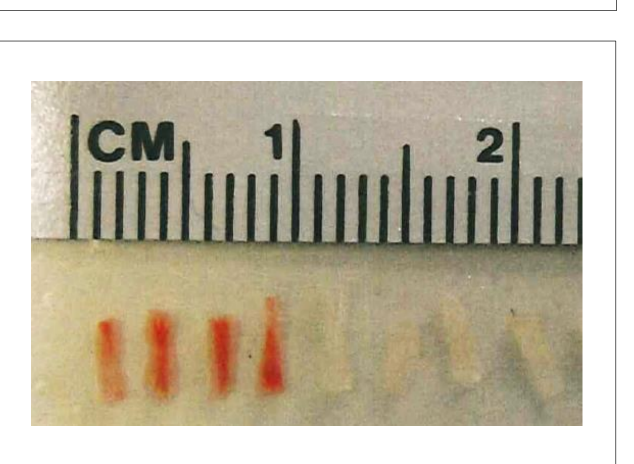
Osteoporosis, a disease of low bone mass that results from bone resorption exceeding bone formation, places individuals at enhanced risk for fracture, disability, and death. There is an urgent and unmet need for novel targets in treating osteoporosis, requiring a better understanding of the endogenous mechanisms regulating bone formation. We reported that deletion of the *Bmpr2* gene in skeletal progenitor cells of mice causes substantially elevated bone mass in young adulthood due to increased bone formation rate (Lowery et al, 2015). As yet unpublished work indicates the age-related decline in bone mass of Bmpr2 mutant mice is reduced approximately three-fold compared to control mice; quantification of serum bone turnover markers indicates this is caused by a sustained increase in bone formation rate to at least 35 weeks of age with no alteration in bone resorption. Here, we determine the gene signature associated with elevated bone formation rate using genome-wide transcriptome profiling in bones of 35-week-old control and *Bmpr2* mutant mice. Applying stringent criteria comparing the expression data to eight well-accepted housekeeping genes (Ppib, Gapdh, Hprt, Tbp, Ppia, GusB, Prkg1, and Ywhaz), we found that, out of 24,980 exon-containing transcripts detected in both genotypes, 334 genes were up-regulated and 310 were down-regulated at least two-fold compared to controls. An additional 704 genes were detected in only one genotype. We refined this putative signature by performing transcriptome profiling in these animals at 55 weeks of age when bone formation rate is no longer elevated. This revealed that, of those genes altered at 35 weeks of age, 461 (71.5%) were either no longer up-regulated or down-regulated in *Bmpr2* mutant mice by 55 weeks of age. Bioinformatic analyses on this refined gene set indicates that elevated bone formation rate in *Bmpr2* mutant mice correlates with enrichment for genes containing binding sites for transcription factors associated with skeletal homeostasis, including FOXP1, SOX2, EGR1, E2F1, KLF4, CNOT3, STAT4, and FOXA1. Further, several genes corresponding with osteoblast differentiation and activity, such as *Pak4* and Pla2g4a, the latter of which encodes cytosolic phospholipase A2 and whose deletion causes osteopenia, are up-regulated in *Bmpr2* mutant mice. Collectively, our findings provide insight into the mechanisms regulating age-related bone loss and highlight potential targets for therapeutic modulation of bone mass.

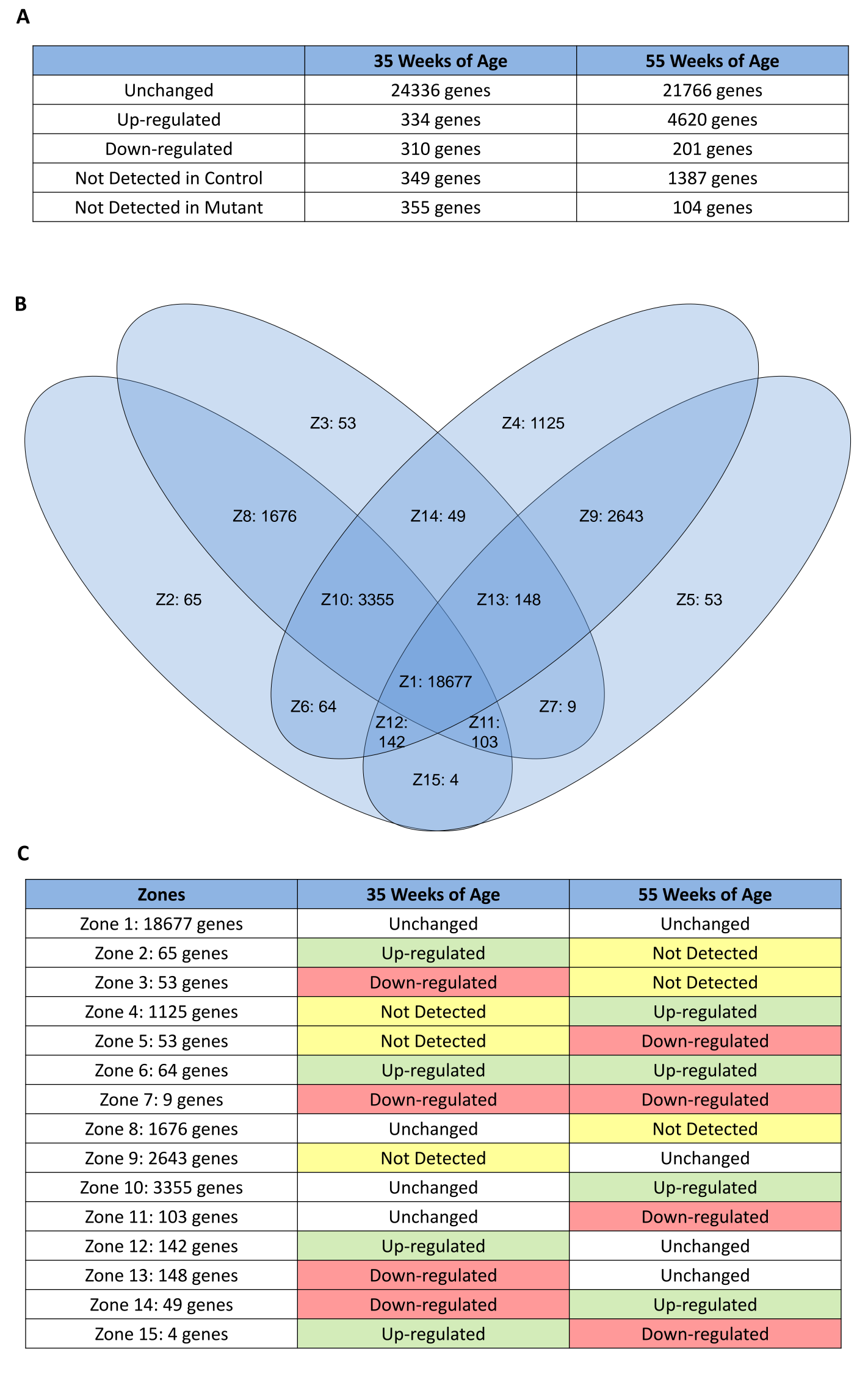


mineral density (vBMD) was quantified by micro-CT in females at 15 and 55 weeks of age. Mean decline in mg hydroxyapatite per cubic centimeter for each genotype between 15 and 55 week old cohorts is indicated (mg HA/ccm); gray bars denote 95% confidence intervals. B: Quantification of the bone formation marker PINP in sera of control and Bmpr2 mutant mice using ELISA. Individual samples are represented by circles and group mean by horizontal lines ± SEM; p values determined by unpaired t test.

#### **RNA-Seq Workflow:**

- 1) Humerii obtained from four each control and Bmpr2 mutant mice at 35 weeks and 55 weeks of age
- 2) Marrow removed by gentle centrifugation
- 3) Bones homogenized and total RNA collected
- 4) Each genotype pooled at equal RNA amounts per mouse 5) Pooled RNA samples shipped to GENEWIZ; quality control
- performed 6) rRNA depletion and library synthesized then sequenced
- 7) Bioinformatics analysis using ENRICHR





**Figure 3: A:** Results of RNA-Seq analyses at 35 and 55 weeks of age; expressed relative to control. **B-C:** Comparison of *Bmpr2* mutant results relative to control at 35 and 55 weeks of age represented in Venn diagram (B) and tabular (C) forms.

# Krista Jackson, Aaron Hudnall, Jonathan W. Lowery Division of Biomedical Science, Marian University College of Osteopathic Medicine, Indianapolis, Indiana

## **ChEA 2016**

SMAD1_18555785_ChIP-Seq_MESCs_Mouse
DACH1_20351289_ChIP-Seq_MDA-MB-231_Human
FOXP1_22492998_ChIP-Seq_STRATIUM_Mouse
SOX2_19030024_ChIP-ChIP_MESCs_Mouse
EGR1_23403033_ChIP-Seq_LIVER_Mouse
TBP_23326641_ChIP-Seq_C3H10T1-2_Mouse
SOX2_18692474_ChIP-Seq_MESCs_Mouse
E2F1_21310950_ChIP-Seq_MCF-7_Human
RNF2_16625203_ChIP-ChIP_MESCs_Mouse
EKLF_21900194_ChIP-Seq_ERYTHROCYTE_Mouse

### ENCODE and ChEA Consensus TFs from ChIP-X

CEBPD_ENCODE	
STAT5A_ENCODE	
NELFE_ENCODE	
PBX3_ENCODE	
NFIC_ENCODE	
PML_ENCODE	
FOS_ENCODE	
TAF7_ENCODE	
SP1_ENCODE	
E2F4_ENCODE	

Figure 4: A: Example ENRICHR analysis on zone 15 genes.

#### **Conclusions, Significance & Future Directions:**

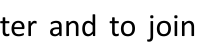
- *Bmpr2* mutant mice display high bone mass in young adulthood and reduced agerelated bone loss.
- Genome-wide transcriptome profiling of *Bmpr2* mutant bones identified 461 differentially expressed genes associated with increased osteoblast activity. The differential gene signature is enriched for genes containing binding sites for corresponding with osteoblast differentiation and activity are up-regulated in *Bmpr2* mutant mice.
- Collectively, our findings provide insight into the mechanisms regulating agerelated bone loss and highlight potential targets for therapeutic modulation of bone mass.
- Future studies will involve functional studies to narrow the gene signature to those that regulate osteoblast function.

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transcription factors associated with skeletal homeostasis. Several genes

