## Glucocorticoids reduce bone strength through reduction in vascularity and hydration, while concurrent treatment with PTH increases bone mass and preserves angiogenic and nitric oxide gene expression in glucocorticoid-treated mice

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## Introduction

Glaccoatroids (GC) Induce osteonecrosis (ON) and osteoporosis (OP); however, the mechanism is complicated. While GCs may increase the risk of OH by reducing anjegenesis and vasoactivity, the reduction in hone strength that accompanies GC use is greater than can be explained by the loss of hone mass alone. To try to understand this discrepancy, we evaluated GCs effects on novel bone quality measures, including bone bene hydration, bone blood flow, and bone anjegenesis gene experssion. We performed two experiments. The first was to understand the role of GC on bone hydration, bone blood flow, and strength, and whether this is altered by anti-asscular endothelial growth factor (VEGF). In the second study we evaluated GC effects on bone vascularity by evaluating gene expression in bloom, and FPH, a known accolaractive agent, influences

### Methods

- Part 1. NOVEL MEASURES OF BONE STRENGTH: HYDRATION, BLOOD FLOW (SUV), & STRENGTH
- > 9-week-old male BALB/cj mice (n=8 per group) were randomized into groups receiving Vehicle (VEH), CG (4 mg/km/d methylprednisolone) for 120 days), GC for 60 days followed by anti-VEGF for 60 days, or GC for 60 days followed by no treatment for 60 days. Mice were sacrificed on day 60
- Outcome measures: bone strength, PET/CT NaF for blood flow (SUV), bone hydration volume fractions of bound water (BW) using <sup>1</sup>H-NMR relaxometry were measured on the intact right femurs
- > IHC of distal femur blood vessels with endomucin and CD31.
- Part 2. GC EFFECTS ON BONE ANGIOGENESIS GENE EXPRESSION & BONE HEALTH
- > 12-week-old male BALB/cJ mice were randomized into groups receiving VEH, GC (4 mg/kg/d methylprednisolone by pellet), or GC+PTH 40 ug/kg/d for 45 days (n=12-24 per group). Mice were sacrificed on day 45.
- Outcome measures: trabecular bone volume (BV/TV), trabecular thickness (Tb.Th), trabecular number (Tb.N), and structure model index (SMI) of
- lumbar vertebral body (LVB) 5 trabecular bone was determined by MicroCT > RNA was extracted from LVB4 to perform 3'-Tag RNA-Sequencing (RNA-Seq)(ng-4/grn)
- Differentially-expressed genes were determined followed by hierarchical clustering and functional annotation enrichment analyses with the ToppFun tool.

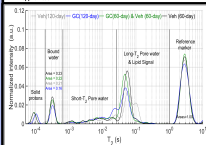


Figure 1. Hydration of bone in GC, GC-vehicle and control groups. Area under the curve for bound water is notably reduced in GC-120 day group compared to vehicle and recovery groups.

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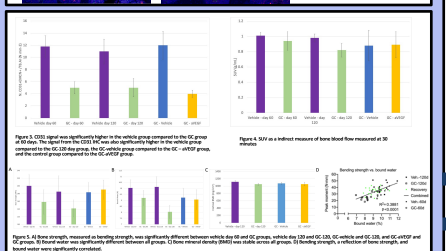




Figure 6. Gene expression associated with the angiogenic and Nitric Oxide (NO) pathways differed between GC-only and VEH mice, and GC-only and GC+PTH treated mice.

Table			
Group	VEH (n=10)	GC only (n=24)	GC+PTH (n=23)
Variable	Mean±SD	Mean±SD	Mean±SD
BV/TV (%)	22.9±2.2	18.8±3.4 <sup>v</sup>	30.8±3.5v8
Tb.Th (μm)	47.9±2.1	42.4±3.4v	57.7±4.3vs
Tb.N (1/mm)	4.80±0.35	4.80±0.36	4.71±0.38
SMI	0.79±0.25	1.23±0.25 <sup>v</sup>	-024±0.41vs
Incidence of Osteonecrosis (%)	0 (0/8)	28 (6/21)	6 (1/17)

vP<0.0005 compared to VEH; 8P<0.0005 compared to GC-only

Table 1. Measurements of bone strength, microarchitecture, and osteonecrosis.

## Summary and Conclusio

GCs reduce bone strength through reduction in bone vascularity and hydration with less change none mass. Interestingly, GCs reduces nitric olde and angiogeneit gene expression while hPTH[1:3] can reverse it. Future studies should address if GC+PTH can prevent GC induced bone fragility through maintenance of vascularity and hydration.

## Acknowledgement

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