Name of Subcontractor: Norman J. Karin, Ph.D.

Title: Assistant Professor/Integrative Biology

Institution: The University of Texas Health Science Center at Houston

Name of Project: Applications of Anabolic Vitamin D Analogs as Countermeasures to Bone Loss

Amount of Grant: 

* Amount Spent, if Different from Amount Granted:

Date Project Was Completed: June 30, 1998

Grants Officer: K. Robert Opperman

Title: Director, Contracts and Grants Management

Phone: 500-4946

Fax: 500-4955
Applications of Anabolic Vitamin D Analogs as Countermeasures to Bone Loss

The experiments in Round 2 were designed to extend the results of our efforts in Round 1 which led us to hypothesize that the seco-steroid, 1,25-dihydroxyvitamin D₃ [1,25(OH)₂D₃], acts in synergy with parathyroid hormone (PTH) to regulate bone calcium homeostasis¹. Our work centered on one particular target of 1,25(OH)₂D₃ action, the voltage-sensitive calcium channels (VSCC’s), which are activated acutely by this steroid within milliseconds of exposure².

We found that the vitamin D analog, AT, which cannot occupy nuclear vitamin D receptors, was also a potent agonist of PTH priming. In these two months we have learned that two other steroids, estrogen and progesterone, are ineffective as priming agents [although estrogen at high (nonphysiological) concentrations will prime bone cells’ response to PTH]. Interestingly, our data indicated that the 24,25-dihydroxy metabolite of vitamin D₃ can neither cause intracellular Ca²⁺ transients nor modify the response of bone cells to PTH. This exquisite specificity of 1,25(OH)₂D₃ suggests an important new role for this hormone in the regulation of bone mass.

We also determined that Ca²⁺ channel expression (mRNA level) is reduced in cultured rat calvarial cells treated with the synthetic steroid, dexamethasone, compared to untreated control cells. When cells treated with dexamethasone were loaded for 1, 3, and 5 weeks, respectively, there was up regulation of the mRNA that was dependent on the duration of loading. This is further evidence of steroid-linked regulation of VSCC expression in bone cells, and supports our interpretation of the results of our experiments with vitamin D₃.

A second area of research focused on the effects of mechanical strain on VSCC expression in bone. These experiments were performed in collaboration with Dr. Steven Goldstein (Univ. Michigan), who generously provided RNA extracted from dog bones that had been exposed to mechanical strain in vivo. Our results suggest that mechanical loading elevated VSCC expression in the long bones from 3 of the 6 animals tested.

To confirm the identity of the VSCC isoform expressed in dog bone, we used a set of PCR primers designed for rat calcium channel on dog RNA samples. The primers amplified 246 base pair (bp) fragment from cDNA synthesized using as a template mRNA from ROS cells, primary rat calvarial cells and rat bone marrow cell culture. Similarly, this set of primers also amplified a 246bp DNA from dog bone cDNA samples. In addition to the 246bp PCR product, we consistently see another 600bp product in dog samples. These data suggest a unique form of Ca²⁺ channel in dog bone; we hypothesize that this form may be present in human bone as well.

A second line of experimentation, carried out in collaboration with Dr. Randall Duncan, a NASA-funded investigator in Indianapolis, centered on RT-PCR analysis of effects of mechanical strain on Ca²⁺ channel expression in cultured bone cells. Compared to unstrained controls, the expression of vitamin-D-sensitive Ca²⁺ channels is elevated 3- to 5-fold over a 24 hr period.

One presentation of data generated by Round 2 research was made³, and two abstracts have been submitted.
References:

