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men are vitamin D3 deficient compared to ~20% of EA men. The level of skin pigmentation is correlated with the extent of							
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### Introduction

African American (AA) men are disproportionally affected by prostate cancer (PCa). AA men are not only at increased risk of PCa compared to American men of European descent (EA), but also are at the highest risk of aggressive PCa and death from PCa. Vitamin D3 deficiency increases PCa mortality, highlighting the importance of maintaining adequate vitamin D3 status for prostate health. Vitamin D3 is acquired in the diet or via UVB/sunlight-initiated synthesis in the skin. Cutaneous melanin absorbs UVB radiation, which leads to reduced vitamin D3 status compared to ~20% of EA men. The level of skin pigmentation is correlated with the extent of African ancestry and serum vitamin D3 status. Besides vitamin D3 status, the activity of vitamin D3 is mediated by the vitamin D receptor (VDR) and determined by several cytochrome P450 metabolism enzymes that bioactivate/inactivate the active form of the hormone, 1,25-dihydroxyvitamin D3 (1,25D).

We hypothesized that the high prevalence of vitamin D3 deficiency in AA men is associated with reduced prostatic concentrations of vitamin D3, which leads to lower expression of vitamin D pathway genes and suppress pro-differentiating actions of vitamin D3 in the prostate; this ultimately abrogates the chemoprotective effects of this natural hormone and raises the susceptibility of AA men to aggressive PCa.

#### **Keywords**

Vitamin D, prostate cancer, African-American

#### **Overall Project Summary**

This research proposal brings together two well known health disparities that affect African-American men. The first is that African-American men are disproportionally affected by prostate cancer in that African-American men are not only at increased risk of prostate cancer compared to American men of European descent, but also are at the highest risk of aggressive prostate cancer and death from prostate cancer . The second disparity is the rampant vitamin D3 deficiency in the African-American population. There is a biological component to this deficiency because sun-induced vitamin D3 synthesis in the skin is significantly reduced in melanin-rich pigmented skin. Consequently, about two thirds of African-American men are vitamin D3 deficient compared to about 20% of men of European descent. It is important to maintain a healthy vitamin D3 status because vitamin D3 deficiency increases the risk of prostate cancer mortality.

Our study will directly examine the amount of vitamin D3 in the prostate tissue of a racially diverse group of patients to discern differences in African-American men. Secondly, we will investigate several mediators of vitamin D3 activity in the prostate tissue and in a novel cell culture model to identify innate molecular differences that may be present between men of European descent and in African-American men that increase susceptibility to aggressive prostate cancer in the latter group.

#### **Key Research Accomplishments**

In year one of this award we have made significant progress in both the patient sample analyses and the *in vitro* assays.

With the patient samples, in Y1 we completed African ancestry estimation and measurement of serum vitamin D metabolites in all of the patients for these study (Figure 1). Of note, we changed cohorts as the patient specimens we planned to use (N=50 from)collaborator Vince Freemen) became unavailable due to a freezer meltdown. We were able to acquire the necessary samples of frozen prostate tissue, serum and whole blood from 50 patients (25 AA and 25 EA) by collaborating with Dr. Peter Gann (here at UIC) and via purchase from the Cooperative Human Tissue Network (CHTN). In our cohort, the percentage of African Ancestry in men ranged from 2-95% (Figure **1A**), which demonstrates the diverse ethnic background of self-declared black men and underscores the value of this additional analysis in interpreting our final data sets.

Serum measurement of 25-hydroxyvitamin D (25D) is used to determine vitamin D status and the AA patients had significantly lower 25D (**Figure 1B**). The levels of 1,25-dihydroxyvitamin D (1,25D), the active hormone, were also measured and not significantly different (**Figure 1C**).

In Y2 we have completed the extraction and measurement of the vitamin D metabolites has been fully optimized in the prostate tissue. The vitamin D measurement was done in collaboration with Heartland Assays, a lab that only measures vitamin D metabolites and was started by the world renowned vitamin D expert Bruce Hollis. Heartland Labs measured all of our serum samples and in the frozen human prostate tissues. The





tissue findings were quite surprising and show that the AA prostate has *higher* 1,25D compared to the EA, despite having lower 25D (**Figure 2A-B**). The levels of the vitamin D metabolism and response genes will be measured in Y3 and may explain this unexpected finding.

In Y2 the laser-capture microdissection (LCM) on the frozen prostate tissues has been completed. There is sufficient RNA from all patients to complete gene expression analysis by whole transcriptome amplification followed by PCR (**Table 1**).

Since moving to the University of Arizona in September of 2014, Dr. Kittles remains an active participant in the project. He has completed SNP analysis of all of the patients for the ancestry estimation as well as for 27 SNPs in key vitamin D response/metabolism genes (VDR, VDBP, CYP24A1, CYP27B1, etc).



dihydroxyvitamin D levels in the patient cohort. Concentrations of 25-D and 1,25D in serum by UHPLC-MS-MS. Mean and 95%CI shown

#### Conclusion

To date, we have completely analyzed prostate and serum vitamin D metabolites in the cohort. SNPs for African ancestry and 27 SNPs related for vitamin D have also been completed in the in a cohort of 50 diverse prostate cancer patients. Similar with other reports, vitamin D status in the AA patients is lower than the EA as measured by serum 25D. Unexpectedly, the tissue levels of 1,25D were higher in the AA patients compared to EA. Molecular analysis of the patient samples has begun and may reveal gene expression changes in the vitamin D metabolism genes that explain the tissue 1,25D status. We are on schedule to complete Aims 1 and 3 of the project. Aim 2, which is in vitro, is also on schedule. An updated proposed timeline for our

#### Tissue ID Race RNA (ng/uL) Total RNA (ng) 260/280 4 ΕA 8.59 214.75 2.09 9 AA 7.15 178.75 1.84 10 AA 7.35 183.75 1.48 11 AA 10.24 256 1.95 12 AA 21.96 549 1.73 AA 416.5 14 16.66 3.39 15 AA 4.98 124.5 1.58 24 AA 6.57 164.25 1.8 AA 530.5 36 21.22 1.69 38 ΕA 11.87 296.75 1.68 27.39 39 AA 684.75 1.74 AA 12.51 312.75 2.79 40 43 ΕA 7.4 185 2.1 45 AA 10.65 266.25 1.99 57 AA 12.35 308.75 1.89 58 ΕA 11.26 281.5 2.16 59 AA 9.39 234.75 2.21 AA 62 14.72 368 2.5168 AA 9.9 247.5 1.92 74 AA 17.83 445.75 2.23 78 ΕA 14.24 356 2.88 80 AA 15.57 389.25 2.12 82 AA 10.05 251.25 2.49 83 AA 2.09 12.23 305.75 86 ΕA 9.96 2.2 249 AA 87 10.31 257.75 1.91 88 EA 7.51 187.75 1.87 89 EA 17.48 437 1.76 90 AA 11.14 278.5 1.75 WD-20645 AA 30.14 753.5 2.63 WD-20669 AA 10.49 262.25 1.81 WD-20675 AA 20.14 503.5 1.85 WD-20693 ΕA 7.93 198.25 2.66 1.91 WD-20696 ΕA 6.59 164.75 703 1.5 WD-25289 AA 28.12 WD-25292a ΕA 6.16 154 1.72 WD-25295 AA 13.98 349.5 1.61 WD-25298 EΑ 6.98 174.5 1.66 WD-25301 EΑ 9.51 237.75 1.58 WD-25304 ΕA 6.98 174.5 1.91 WD-25307 AA 8.65 216.25 1.93 WD-25313 EΑ 7.9 197.5 1.91

WD-25316

EΑ

6.05

151.25

1.63

current status and remainder of the project is shown in Figure 3.

Figure 3. Updated Project timeline							
Y1	Y2		Y3				
Serum 25D and 1,25D measurement	Tissue 25D and 1,25D measurement		Combined analysis of all patient data <i>and</i> Manuscript preparation				
Patient ancestry SNP analysis	Patient vitamin D SNP analysis						
Tissue LC	RT-qPCR for vitamin D-related genes						
Cell culture ancest analysis	ry SNP	Cell culture differentiation by vitamin D					

# Table 1. RNA from LCM-collected benign prostate epithelium

## **Publications, Abstracts, and Presentations**

# **Publications:**

Thesis: Farhat, Rachael S. Title: Examination of Vitamin D Disparities in African American and Caucasian Prostate Cancer Patients and Cells. URL http://hdl.handle.net/10027/19512 Publication Date 2015

Abstract and Invited Presentation: Dec 2013 American Association for Cancer Research (AACR)- The Science of Cancer Health Disparities in Racial/Ethnic Minorities and the Medically Underserved, Atlanta, GA."Vitamin D and prostate Cancer in African-American Men"

Abstract and Poster presentation: November 2015 Steroid Research Congress Chicago. "Vitamin D metabolites in prostate tissue and serum from African-American Men". Richards, Z, Farhat R, Kittle R, Nonn L.

**Inventions, Patents and Licenses** none

Reportable Outcomes none

Other Achievements none

References none

Appendices none