

AWARD NUMBER: W81XWH-15-1-0099

TITLE: DNA Damage and Oxidative Stress in Dyskeratosis Congenita: Analysis of Pathways and Therapeutic Strategies Using CRISPR and iPSC Model Systems

PRINCIPAL INVESTIGATOR: Frederick Goldman

CONTRACTING ORGANIZATION: University of Alabama at Birmingham
Birmingham, AL 35233

REPORT DATE: June 2018

TYPE OF REPORT: Annual

PREPARED FOR: U.S. Army Medical Research and Materiel Command
Fort Detrick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for Public Release;
Distribution Unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.

REPORT DOCUMENTATION PAGE

Form Approved
OMB No. 0704-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing this collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to Department of Defense, Washington Headquarters Services, Directorate for Information Operations and Reports (0704-0188), 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302. Respondents should be aware that notwithstanding any other provision of law, no person shall be subject to any penalty for failing to comply with a collection of information if it does not display a currently valid OMB control number. **PLEASE DO NOT RETURN YOUR FORM TO THE ABOVE ADDRESS.**

1. REPORT DATE June 2018		2. REPORT TYPE Annual		3. DATES COVERED 1 Jun 2017 - 31 May 2018	
4. TITLE AND SUBTITLE DNA Damage and Oxidative Stress in Dyskeratosis Congenita: Analysis of Pathways and Therapeutic Strategies Using CRISPR and iPSC Model Systems				5a. CONTRACT NUMBER	
				5b. GRANT NUMBER W81XWH-15-1-0099	
				5c. PROGRAM ELEMENT NUMBER	
6. AUTHOR(S) Frederick Goldman, MD Erik Westin, PhD E-Mail: fgoldman@peds.uab.edu				5d. PROJECT NUMBER	
				5e. TASK NUMBER	
				5f. WORK UNIT NUMBER	
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) University of Alabama at Birmingham 1600 7th Avenue south, ACC 512 Birmingham, AL 35233-0111				8. PERFORMING ORGANIZATION REPORT NUMBER	
9. SPONSORING / MONITORING AGENCY NAME(S) AND ADDRESS(ES) U.S. Army Medical Research and Materiel Command Fort Detrick, Maryland 21702-5012				10. SPONSOR/MONITOR'S ACRONYM(S)	
				11. SPONSOR/MONITOR'S REPORT NUMBER(S)	
12. DISTRIBUTION / AVAILABILITY STATEMENT Approved for Public Release; Distribution Unlimited					
13. SUPPLEMENTARY NOTES					
14. ABSTRACT Dyskeratosis congenita is a disorder that arises due to prematurely shortened telomeres and characterized by a classical clinical triad of leukoplakia, skin dyspigmentation and nail dystrophy with concomitant marrow failure. DC symptomology, to a degree, corresponds to critically shortened telomeres that limits cellular replicative potential and thus prematurely exhausts stem cell pools. Our previous findings support a hypothesis whereby shortened telomeres increase DNA damage responses within the cell leading to heightened reactive oxygen species (ROS). Recent work supported under this research grant have uncovered a suppressed steady-state RNA expression and protein levels of the NRF2 and NRF3 pathways within DC skin fibroblasts. This suppression is also found in cells exposed to an environment with experimentally-elevated oxidative stress. Furthermore, preliminary RNA-Seq experiments suggest that senescence associated secretory pathway (SASP) is altered in DC cells and suppressed upon mobilization of telomerase. Finally, decreased oxidative stress levels could be found in DC cells upon forced expression of an NRF2 heterodimerizing protein, MAFG, suggesting that suppression of this pathway yields to an unmitigated increase in ROS. Together, these findings provide support for a mechanism whereby shortened telomeres mobilize a DNA damage response that in turn suppresses antioxidant proteins leading to an increase in oxidative stress.					
15. SUBJECT TERMS None provided					
16. SECURITY CLASSIFICATION OF:			17. LIMITATION OF ABSTRACT	18. NUMBER OF PAGES	19a. NAME OF RESPONSIBLE PERSON
a. REPORT	b. ABSTRACT	c. THIS PAGE			USAMRMC
Unclassified	Unclassified	Unclassified	Unclassified	24	19b. TELEPHONE NUMBER (include area code)

Table of Contents

	<u>Page</u>
1. Introduction.....	4
2. Keywords.....	4
3. Accomplishments.....	4
4. Impact.....	20
5. Changes/Problems.....	20
6. Products.....	20
7. Participants & Other Collaborating Organizations.....	21
8. Special Reporting Requirements.....	21
9. Appendices.....	21

Introduction

Most cells rely on telomere attrition to undergo a limited number of cell divisions as a means to circumvent uncontrolled cell growth. Telomere attrition is one well-characterized means by which replication potential is limited, as illustrated by the dramatic phenotype presented by patients with Dyskeratosis Congenita (DC). DC arises due to mutations found in telomere/telomerase related genes. The classical clinical triad of DC is leukoplakia, skin dyspigmentation and nail dystrophy with concomitant bone marrow failure (BMF). DC symptomology, to a degree, corresponds to critically shortened telomeres that limits cellular replicative potential and thus prematurely exhausts stem cell pools. We have previously characterized DC cells and demonstrated inherent defects in multiple cell types and have uncovered evidence that elevated reactive oxygen species (ROS), due to shortened telomeres, may be causative in limiting cellular potential. Other labs have indicated that hematopoietic stem cells (HSC) are acutely sensitive to oxidative stress, which could be a contributing factor to BMF in DC. This increase in ROS appears to be related to a signal transduction pathway initiated by shortened telomeres and transduced by the DNA damage response (DDR) via p53. **Together, this evidence suggests that ROS may be causative towards establishing a senescent state in a short-telomere background.** Little is known about relevant factors lying downstream of p53 that ultimately lead to telomere-dependent elevated ROS. Preliminary evidence generated by our lab has found expression of the antioxidant-related genes *NRF2* and *TXN* are repressed in DC fibroblasts and rescued upon exogenous expression of *TERT*. DC fibroblasts expressing *NRF2* and *NRF3* also acquire a phenotype similar to normal cells while *NRF2* shRNA expressing controls appear senescent. Furthermore, individual overexpression of SOD genes indicated *EcSOD* expression significantly decreased ROS. Data gathered as part of this research grant continues to support the outlined hypothesis. This includes further verification of *NRF2* and *NRF3* suppression at the RNA and protein levels. Potential deregulation of *NRF2* heterodimerizing partners like MAFG could also be implicated by data gathered here. Finally, evidence of an altered senescence-associated secretory phenotype has been found in DC fibroblasts. Together, our accumulating data continue to provide evidence how telomere attrition facilitates entry into senescence.

Keywords (limit to 20 words)

Dyskeratosis congenita, bone marrow failure, aging, telomere, oxidative stress, reactive oxygen species, p53, stem cells

Accomplishments

Accomplishments: Major Goals

Our timeline suggested the following approach to this study:

- Year 1 – Make iPS cells, evaluate antioxidants, assess pathways
- Year 2 – Study HSCs and modulation of ROS
- Year 3 – Fully evaluate antioxidants on multiple cell types/genotypes

The following Table was submitted as part of our Statement of Work and has been modified slightly to highlight and update the work completed within each Aim. Text that has been highlighted in red indicates adjustments to the Table based on changes from the previous Progress Report in addition to changes made based on results and/or hurdles encountered during the course of research.

Specific Aim 1: Mechanism of oxidative stress	Proposed Timeline	% Completion
Major Task 1 Overexpress/knockdown genes of interest	Months	
Vector construction	3-6	Finished
Update: addition of new candidate genes	1	Finished
Cell infections	6-12	Finished
Update: infection of new candidate genes	1	Finished*
Milestone(s) Achieved: Assessment of effect	2	Finished*
Local IRB/IACUC Approval	3	Finished
Milestone Achieved: HRPO/ACURO Approval	6	Finished
Major Task 2 CRISPR knockout/RNAseq		
Viral infection/prep	3-6	CRISPR KO virus library prep: finished; RNA-Seq: ~Finished
Cell manipulation	3-6	CRISPR KO virus infection: Finished; Single cDNA infections: Finished
Bioinformatics	1	CRISPR KO library: 25%; RNA-Seq: 100%
Milestone(s) Achieved: Assessment of effect	4	N/A

Specific Aim 2: Anti-oxidant screening	Proposed Timeline	% Completion
Major Task 1 Library/drug acquisition		
Modify/optimize treatment conditions	1	5% (began candidate drug screen; have not begun large library screen)
DHE analyses in fibroblasts first, then other cells	5	5% (began candidate drug screen; have not begun large library screen)
Milestone(s) Achieved: Create pool of molecules that decrease DHE and increase cell proliferation	9	N/A

Specific Aim 3: Production of HSC from iPSC cells	Proposed Timeline	% Completion
Major Task 1 Create iPSC	Months	
Production of iPSC clones from DC fibroblasts	1-3	Finished
Clone characterization	1-3	Finished
Update: production of iPSC clones harboring DC mutations generated by CRISPR: Design	1	Finished
Update: production of iPSC clones harboring DC mutations generated by CRISPR: Production and evaluation	3	10%; Finished
Milestone(s) Achieved Acquisition of stable pluripotent cells	3-6	Finished
Major Task 2.1 Create HSC		
HSC characterization/differentiation	3-6	10%
Milestone(s) Achieved: Achieve true HSC	12-18	0%
Update: Major Task 2.2 Knockout Telomere Genes in CD34 ⁺ Cells		
Update: knockout key telomere-related genes using CRISPR/Cas9 protein component	1	75%
Major Task 3.1 Evaluate ROS in HSC (iPSCs)		
Subset analysis of HSC derived from iPSCs	3-6	10%
Examine pathways	3-6	0%
Milestone(s) Achieved: Determine factors effecting ROS in HSC	9	0%
Update: Major Task 3.2 Evaluate ROS in CRISPR KO CD34 ⁺ cells		
Update: CRISPR KO design	1	100%
Update: CRISPR KO verification	1	100%
Update: CRISPR KO analyses	3	5%
Update: Subset analysis of HSC	3-6	0%
Examine pathways	3-6	0%
Milestone(s) Achieved: Determine factors effecting ROS in HSC	9	0%

Progress report June 2018 (2017 & 2016 Progress Reports provided below)

Due to the delayed start of our grant in 2015, we were approved to receive a no cost extension (MOD P00002 for W81XWH-15-1-099) to allow us three full calendar years to complete our research. **We will therefore submit a final report in the late fall of 2018 to provide an update on the final three months of research.**

2018 Accomplishments: Major activities:

The past year has focused on a couple major goals:

1. Finish overexpression and knockdown of candidate genes culled from RNA-Seq experiments to assess effect on oxidative stress in DC cells (**Aim 1**)
2. Finish CRISPR studies utilizing knockout and overexpression libraries (**Aim 1**)
3. Finish targeting of CD34+ cells for knockout of genes related to DC (**Aim 3**)

In regards to **Aim 1**, we have overexpressed all key candidate genes (~30) derived from RNA-Seq focused analyses (2018 Figure 1) in multiple cell types acquired from DC patients (lung fibroblasts, skin fibroblasts) and interrogated each gene subset for changes in oxidative stress. A number of genes significantly decreased oxidative stress in DC cells with no negative effect on log-phase growth (2018 Figure 2; 2018 Table 1). However, a number of genes slowed cell growth. We hypothesized that overexpression of some putative antioxidant genes may in fact alter steady-state levels of oxidative stress required for log-phase growth yet may be effective at decreasing oxidative stress when the cells are contacted-inhibited (grown to confluency). To circumvent this particular problem, we came up with an alternative strategy that grows the cells to confluency and examines the cells after a month of confluency to assess whether the transgene had an effect on oxidative stress. Using this particular methodology led to uncovering a more genes that decreased oxidative stress with one particular gene decreasing oxidative stress by over two-fold (data not shown). With these candidate genes in hand, we are currently performing a number of functional assays to assess whether DC cells are afforded a proliferative benefit by the expression of these genes. Further, we are also assessing whether these exogenous genes alter telomerase or p53 activity which may, in part, explain the resulting decreases in oxidative stress. QRT-PCR expression assays and western blots will be used to better understand the effects of these exogenous genes.

Further related to **Aim 1**, we have optimized our CRISPR library screens in a number of ways to extract as much meaningful information from each library infection. Once DC fibroblasts (mid- to late-passage cells) have been infected with an empty vector control and have undergone antibiotic selection, very few DC cells are able to create colonies of cells greater than 8-16 cells (3-4 population doublings). However, with the knockout library (and SAM gene activation library) we routinely find ~100 colonies per 100,000-500,000 cells infected. This allows us to perform two sets of analyses. The first is the ability to ring-clone individual colonies and isolate clones isolate from a single infection. This permits us to isolate DNA from this single clone, PCR the gRNA and submit for Sanger sequencing. This allows us to use the gRNA as a barcode for the particular gene that was knocked out in the clone and then ensure that the gene has undergone non-homologous end joining scarring that would knockout the gene. Using this methodology, we have uncovered a number of candidate genes that are currently being re-targeted by single CRISPR knockout vectors to verify the initial clonal results. The second set of analysis that we can perform using the library is by creating large pools of cells that have been infected by the library and sort cells based on their oxidative stress profile. We can sort the low oxidative stress cells (lowest ~15%; 2018 Figure 3) and either 1. isolate DNA or 2. replate for a subsequent second sort to better concentrate the cells for consistency regarding low oxidative stress profiles. Once cells have been isolated, we PCR the gRNAs with indexed primers to permit deep sequencing of gRNAs within subsets of cells. Once clones are sequenced, they are then analyzed for enrichment of gRNAs overrepresented in these populations.

Related to **Aim 3**, we have targeted DC related genes isolated from non-DC cord blood samples (CD34+ cells) and are continuing to evaluate these cells for functional changes related to colony forming assays and gene expression changes. However, the main focus of research for the final three months using CD34+ progenitor cells is to better understand the relationship among RNA-Seq analyses, CRISPR knockout/activation and amelioration of CD34+ cells. Targeted analyses of CD34+ cells knocked out for DC-related genes will primarily focus on whether genes uncovered by RNA-Seq/CRISPR screens can alter deficits found in CD34+ knockouts. The primary data point of interest will be assessing levels of oxidative stress but will also likely include colony forming assays (compared to controls) and potentially western blotting for p53 activation.

2018 Accomplishments: Specific Objectives

Our objectives for Year 3 were to try and complete experiments related to all Aims. Leading up to Year 3, preparation had been completed in regards to all Aims however Aims 1 and 3 quickly became the major focal points based on considerations related to time and effort. Aims 1 and 3 will likely be finished in the remaining three months of research (no cost extension). In regards to Aim 2, further analyses will be performed, time permitting.

2018 Accomplishments: Significant results or key outcomes

We have performed a number of RNA-Seq analyses and have found many novel candidate genes that are differentially expressed in DC cells compared to their control and/or rescued counterparts. Given our pipeline to test genes related to the NRF2 and antioxidant pathway we can easily test these genes as well. Many of these new genes from these analyses were expected based on previous literature investigating senescence that serve as internal controls (e.g. cell cycle related genes, miRNAs, MMPs, cytokines). Furthermore, the use of CRISPR a novel means by which to uncover genes that are capable of rescuing DC cells from proliferative defects found in cell culture. A significant result from these assays is the acquisition of genes capable of decreasing ROS within these cells and/or extending the lifespan. A number of these genes have been characterized as interacting with oncogenes/tumor suppressors. However, the most promising findings lie in uncovering genes that have no know interaction with oncogenes/tumor suppressors that may afford proliferative benefits without altering key cellular processes that could be regarded as proto-oncogenic events. Furthermore, use of CRISPR is of utmost importance for use in CD34⁺ cells as an alternative means to evaluate this difficult-to-study cell type from DC patients. We have found that we are able to successfully knockout DC-related genes at a high rate and believe that we can improve upon this percentage. Cells generated from these experiments will serve as an important source to evaluate the effects of telomere-related ROS in hematopoietic progenitors.

(Progress report June 2017)

2017 Accomplishments: Major activities:

The second year of our research focused primarily on five goals:

1. Infect DC and control fibroblasts with cDNA and shRNA to more closely examine the effects of candidate genes and ROS (Aim 1)
2. Examine expression levels in a number of primary cells and their derivative cell lines (12 total; e.g. cell rescue and ROS diminution via overexpression of TERT and shRNA targeting p53) (Aim 1)
3. Begin gene discovery approach using CRISPR knockout and activation libraries (Aim 1)
4. Build upon previous experiments that manipulate antioxidant pathways using drugs (Aim 2)
5. Differentiate DC iPSCs to generate hematopoietic progenitors (Aim 3)

In regards to **Aim 1**, all antioxidant-related genes of interest (e.g. NRF2-related pathway) have been cloned into viral vectors and have been used to infect DC and controls fibroblasts. In parallel, a number of RNA-Seq experiments were also finished to identify the expression of genes that were elevated in DC cells (cells carrying TERT, TINF2 or DKC1 mutations; fibroblasts and T cells) and control cells (normal fibroblasts and T cells) or rescued cells (DC fibroblasts expressing TERT or p53 shRNA; [2017 Figure 1](#), below). After bioinformatics analysis, we found genes that are commonly suppressed in DC cells but elevated in control/rescued cells ([2017 Figure 2](#)) and identified approximately 20 more novel genes consisting of transcription factors, genes related to oxidative stress, and genes related to cell survival. These genes' cDNAs have been cloned into viral vectors and have been used to infect DC and control fibroblasts for experiments that will coincide with NRF2-related antioxidant gene expression studies. Together, the RNA-Seq candidate genes and the antioxidant candidate genes will be used to test ROS levels (DHE/DCF), proliferation, senescence (SA-B-gal) in the final year of our studies. In addition, pilot experiments using CRISPR knockout and activation libraries began in Year 2 and have generated sufficient data to scale-up experiments. We have successfully titered virus so that cells receive less than one virus per infection (MOI ~0.15-0.30) and supported by experiments sequencing clones isolated from infections. These clonal isolation experiments are executed by isolating proliferative clones of cells within a population of infected cells, post antibiotic selection. These clones are, in essence, rescued by the underlying unique CRISPR gene knockout within each clonal population of cells and thus supporting these proof-of-principal concept. At this time, 20 clones have been isolated from previous infections. Each clone has had the gRNA sequence PCR amplified and sequenced (the gRNA acts as a barcode for the gene target) and have found single gRNA populations within these cells suggesting a single infection ([2017 Figure 3](#)). Future work will entail sequencing the likely target site to ensure gene targeting and verification by overexpressing individual gRNAs to recapitulate initial findings and examine oxidative stress. Furthermore, we will sort cells and capture cells with decreased ROS compared to controls and perform next generation sequencing of the gRNA pool to examine which gRNAs are over-represented in this subset of cells. Verification will entail cloning candidate gRNAs into individual vectors to replicate initial experiments.

Aim 2 is focused on the use of candidate drugs and drug libraries to examine the effect of

decreasing ROS on cell health. Initial experiments began in Year 1 and have continued into Year 2 however much of Year 2 has been dedicated to Aim 1 and Aim 2. Year 3 will execute experiments designed to test these drugs and interrogate oxidative stress levels and subsequent proliferative capacity in drugs capable of decreasing ROS. Given the size of this experiment and the large number of cells required to finish it, we have massively expanded and frozen mid-passage DC and control cells to sufficient numbers to finish this Aim in Year 3.

Experiments designed as a part of **Aim 3** are tailored to assess one key cell type that is rare to study in DC patients due to inherent bone marrow failure: the hematopoietic progenitor (CD34⁺ cells). Our experiments are designed to assess whether elevated ROS in CD34⁺ DC cells plays a causative role in bone marrow failure found in these patients. To this end, we have proposed creating iPSCs from DC patients and coax these cells to differentiate towards CD34⁺ hematopoietic cells using protocols commonly used in our lab. We have succeeded in creating iPSCs from DC cells carrying TERT and DKC1 mutations yet we have found these cells are difficult to maintain in a pluripotent state and furthermore exceedingly difficult to generate sufficient CD34⁺ cells to examine. We have enacted a number of alternative plans to make up for this issue if not improve on the initial plan of action:

1. We are repeating the production of iPSC clones to assess whether different clones may be more permissive for hematopoiesis
2. We are using CRISPR/Cas9 to knockout DC-related genes and alternatively induce DC-centric mutations into normal iPSCs that have been found to be permissive in regards to generating CD34⁺ cells
3. Circumvent iPSCs altogether and use CRISPR/Cas9 to directly knockout or mutate target DC-related genes in normal CD34⁺ cells

In regards to the use CRISPR/Cas9 to modify normal cells to recapitulate the effects of DC mutations, our lab utilizes a proteinaceous Cas9 component to nucleofect CD34⁺ cells and edit the underlying genome. We have tested multiple gRNAs to evaluate whether we can knockout DKC1 and TINF2 in this manner. As indicated in [2017 Figures 3 and 4](#), we have successfully targeted and performed gene knockout via frameshift mutations in a large percentage of these genes in CD34⁺ cells (~42-43%) as well as an even greater percentage in iPSCs (data not shown). Based on our previous analytical experience using this method to calculate modified cells (TIDE: Tracking of Indels by Decomposition), we believe that this percentage of modified cells is an underestimation and will analyze this further by genomic PCR amplification, subcloning and Sanger sequencing. Whether CD34⁺ cells are generated from iPSCs or from CRISPR/Cas9 editing, these cells will be examined for ROS levels (DCF), proliferation, differentiation (FACS, methylcellulose) and expression levels (RNA-Seq).

2017 Accomplishments: Specific Objectives

Our objectives for Year 2 were to try and complete preparatory and pilot experiments related to Aim1 and most of Aim 3. All preparation has been completed in regards to Aim 1 leaving only analyses related to ROS and proliferation to be finished in addition to sequencing related to

CRISPR library experiments. In regards to Aim 3, we have verified means to induce DC mutations into normal cells as a means to circumvent issues related to DC iPSCs. Although a bit further behind than we anticipated on Aim 3 at this time, we project that Aim 3 will be finished on schedule prior to the end of this grant. Finally, Aim 2 will be the primary focus of Year 3.

2017 Accomplishments: Significant results or key outcomes

We have performed a number of RNA-Seq analyses and have found many novel candidate genes that are differentially expressed in DC cells compared to their control and/or rescued counterparts. Given our pipeline to test genes related to the NRF2 and antioxidant pathway we can easily test these genes as well. Many of these new genes from these analyses were expected based on previous literature investigating senescence that serve as internal controls (e.g. cell cycle related genes, miRNAs, MMPs, cytokines). Furthermore, the use of CRISPR is of utmost importance for use in CD34⁺ cells as an alternative means to evaluate this difficult-to-study cell type from DC patients. We have found that we are able to successfully knockout DC-related genes at a high rate and believe that we can improve upon this percentage. Cells generated from these experiments will serve as an important source to evaluate the effects of telomere-related ROS in hematopoietic progenitors.

(Progress report June 2016)

2016 Accomplishments: Major activities

The first nine months of our research focused primarily on the initial stages of Aim 1 and Aim 3. **Aim 1, Major Task 1:** all cDNAs have been cloned into a lentiviral backbone for the purpose of overexpression within DC and control fibroblasts. This will permit the assessment of whether restoration of antioxidant gene expression, found suppressed in DC cells, can ameliorate deficits found in within these cells. These cDNAs include the NRF2 family members (NRF1, NRF2, NRF3), NRF2 heterodimerizing proteins (MAFF, MAFG, MAFK, ATF4, cJun) and related metabolic genes of interest (Pgc1 α and Pgc1 β). We have also acquired a number of shRNAs from OpenBiosystems that will be used throughout this study that include those that target p53, p21 (CDKN1A), NRF2, the NRF2 ubiquitination complex (KEAP1) and the NRF2 transcriptional competitor (BACH1). Given the extent of this pathway we may need to clone more cDNA or acquire further shRNAs to better characterize and/or verify certain findings. Cell infections and antibiotic selections have been performed on 90% of these cDNAs/shRNAs and cells frozen on each of these conditions. Preliminary oxidative studies have been performed in a subset of these infections (data presented below). **Aim 1, Major Task 2:** RNA-Seq has been performed on a number of cell conditions, including skin fibroblasts and skin fibroblasts overexpressing TERT (Control, Control-TERT, DC, DC-TERT) in addition to control lymphocytes and DC lymphocytes. Bioinformatics have also been performed (DESeq package, DE analysis) on these RNA-Seq experiments to elucidate key differences among these subsets of cells. To bolster the RNA-Seq gene discovery approach we also proposed using chromatin immunoprecipitation (ChIP). At this time, no ChIP experiments have been performed. Major Task 2 also proposed the preparation of a CRISPR knockout (KO) viral library as an additional gene discovery approach. This library relies on CRISPR targeting to each gene within the human genome and the byproduct of CRISPR targeting, non-homologous end joining (NHEJ), to create a frameshift knockout. This library has been acquired, cloned in bacteria to maintain the library diversity, transfected into 293T cells for viral preparation and titered to ensure accurate multiplicity of infection (MOI) when ready for infection. Cells will be infected and selected in the near future.

Specific Aim 2 was designed to test candidate drugs and a library of FDA approved drugs for their ability to decrease ROS within DC cells and improve DC cellular functions (i.e. proliferation, decreased senescence). At this time, only a handful of candidate drugs have been tested for decreases in ROS (DHE staining). Major expansion of mid-passage DC and control cells has been performed to provide a consistent source of cells to be used in these studies. Initial experiments testing of all candidate drugs and the use of the FDA approved drug library will start within Year 3 as initially planned within the grant.

Specific Aim 3 was designed to provide access to hematopoietic progenitor cells (HPCs) from DC cells which are difficult to acquire from patients due to their propensity for bone marrow failure and anemia. At this time induced pluripotent stem cells (iPSCs) have been manufactured from DC patients' cells that harbor mutations in the TERT and DKC1 loci. Greater than 10 clones have been produced from each mutation and have been characterized for pluripotency and frozen for further analyses at a later date. At this time, attempts to generate iPSCs from our

patient's cells with a TINF2 mutation have been unsuccessful after two attempts (highlighted in the above Table by asterisks) however further attempts have remained unsuccessful. We have proposed within Specific Aim 3 to differentiate these iPSCs (and controls) into hematopoietic progenitor cells for further analyses. Initial experiments to derive and characterize DC HPCs will take place within Year 2.

2016 Accomplishments: Specific Objectives

Within the first year our objective was to finish as much of Aim 1 as possible (create all lentiviral vectors and subsequent infections, perform gene discovery experiments [RNA-Seq, CRISPR KO library, ChIP] and initiate experiments within Aim 2 and Aim 3. Since our research funding began in October of 2015, we are on track to accomplish the vast majority of Aim 1 within a full calendar year and are on schedule to pursue the major portions of Aim 2 and 3 with early experiments having been completed.

2016 Accomplishments: Significant results or key outcomes

To date, we have accumulated further evidence supporting the modulation of NRF2 at the protein level (Western) by short telomere signaling that is in agreement with preliminary evidence provided for the application of this grant (RNA expression changes). It appears as telomere shortening takes place, NRF2 RNA and protein levels undergo a concomitant decrease. Upon activation of telomerase (exogenous TERT expression) and subsequent telomere elongation, NRF2 protein (2016 Figure 1) and NRF3 RNA levels (2016 Figure 2 & 3) are restored and oxidative stress is decreased (preliminary data). Consistent with our hypothesis, antioxidant responses are also suppressed during periods of heightened oxidative stress compared to controls (exposure to increasing concentrations of peroxide; data not shown). We hypothesize that p53 may be actively suppressing antioxidant responses as part of the DNA damage response. We have previously found that expression of p53 shRNA decreases ROS and have now found evidence that p53 shRNA also increase NRF2 and NRF3 RNA expression and protein levels (NRF2 data not presented; 2016 Figure 4).

RNA-Seq has provided substantial insight into the changes that take place between controls and DC cells (fibroblasts [2016 Figure 5] and lymphocytes) as well as changes that take place in DC cells 'corrected' by expression of TERT. For example, when looking for gene expression profile differences between DC and control cells that reflect changes found in DC cells compared to 'rescued' DC-TERT expressing cells we found 74 genes that were elevated in both controls and DC-TERT cells and 54 that were decreased in DC cells. Interestingly, NRF3 was found increased in control and DC-TERT expressing cells while the NRF2 heterodimerizing partner FOS was elevated in DC cells suggesting a potential inhibitory element. Furthermore, when performing a gene ontology enrichment analysis, there is an overrepresentation of CXCR chemokine receptor binding genes ($p=0.0007$) suggesting a modified autocrine/paracrine effect within the secretory pathway. Other potentially interesting genes were also uncovered relating to telomere biology and transcriptional regulation that may be of interest for further study. Regarding RNA-Seq performed on DC and control lymphocytes (three controls, three DC patients), the gene most significantly decreased in DC cells was GSTM1 (glutathione S-

transferase mu 1; $p=0.0005$, 600-fold decrease in DC cells). In DC lymphocytes, suppression of two other glutathione-S-transferase (GST) genes, GSTM2 and GSTT1 was also found (4-6-fold, $p<0.0001$). Importantly, GSTM1 and GSTM2 are NRF2 transcriptional targets and GSTM1 and GSTT1 have been previously implicated in the acquisition of aplastic anemia.

Candidate genes within the NRF2 family and pathway have been cloned into lentiviral vectors for the purpose of circumventing the suppression of the endogenous gene by overexpressing the exogenous cDNA in DC cells. If an increase in oxidative stress is causal towards premature entry into senescence then restoration of key antioxidant genes may afford improved parameters to DC cells. One particular example of an unexpected candidate gene that decreased ROS (DHE staining) upon lentiviral overexpression was MAFG (2016 Figure 5). Further research will be required to assess the nature of the relationship between MAFG and the NRF2 family to decrease ROS in this context.

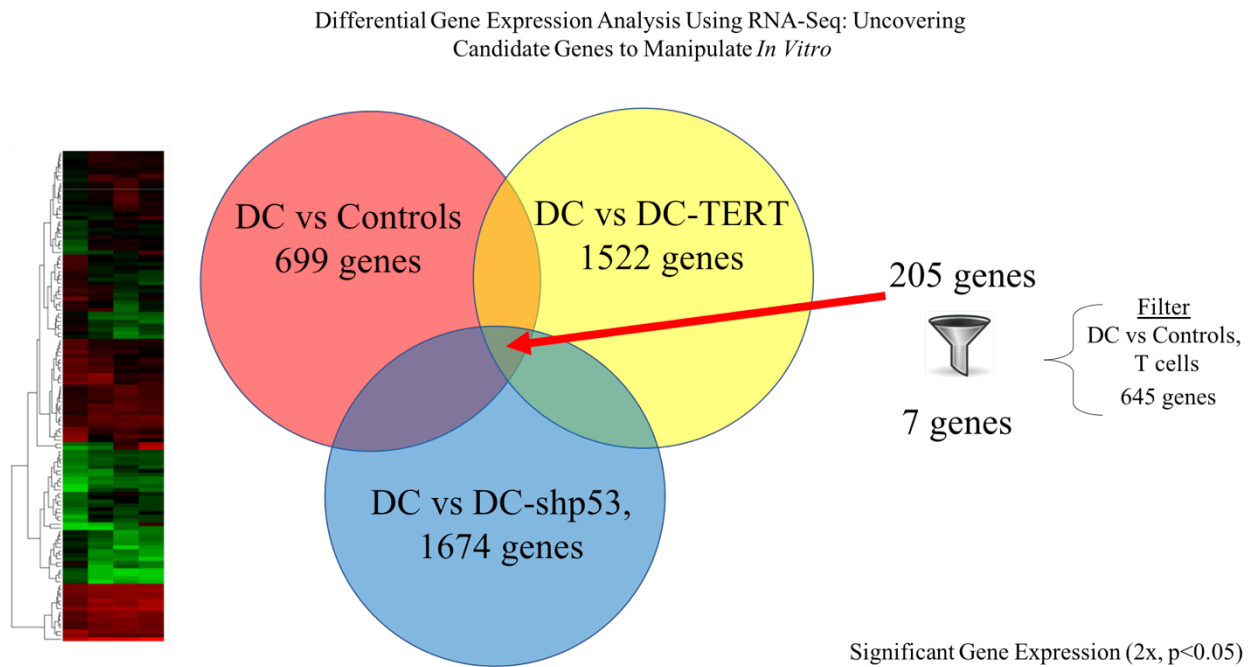
2016 Accomplishments: Results Dissemination

Data from this project has been presented publicly at the American Society of Hematology (ASH) in December of 2015 as a poster presentation by the postdoc, Erik Westin. Further dissemination in the near future regarding data accumulated under this grant will take place at the 2016 ASH Meeting and manuscript preparation will likely be underway prior to that meeting.

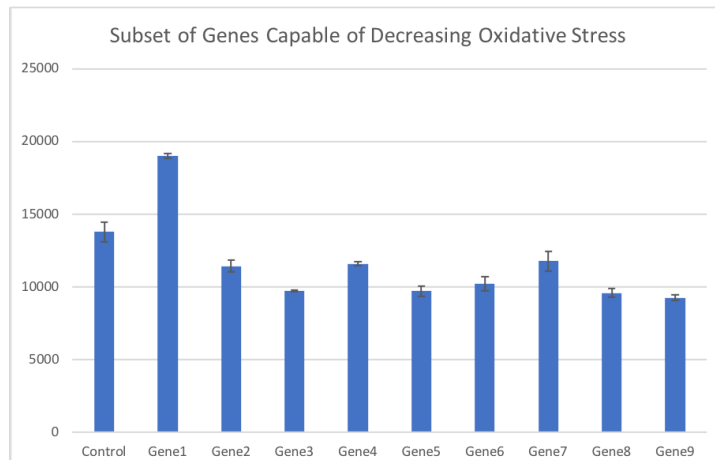
For the next reporting period, we anticipate that we will have accumulated data regarding a number of key experiments. Namely,

1. Oxidative stress and proliferation data regarding cDNA and shRNA expression studies
2. Gene discovery approaches (RNA-Seq, CRISPR-KO, ChIP) initiated
3. Drug Screen: the drug screen will begin in Year 2 and likely completed initial analysis of all compounds
4. DC iPSCs: initial experiments will be underway to evaluate hematopoietic progenitors and their ability to differentiate into multiple lineages (myeloid/erythroid/lymphoid)

2018 Progress Report Figures



2018 Figure 1 Application of Bioinformatic Filter to Isolate Antioxidant Genes

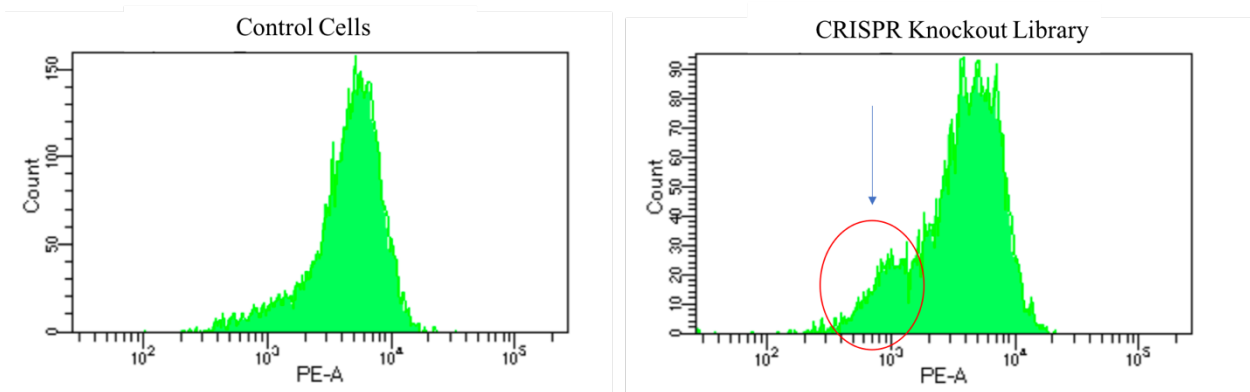


2018 Figure 2 Focus on Genes Capable of Decreasing Oxidative Stress in DC Fibroblasts When Overexpressed (Gene 1 Consistently increased oxidative stress; serves as control)

Gene Name	Fold % of Oxidative Stress Compared to Control
Control	x
Gene1	138
Gene2	82.93
Gene3	70.57
Gene4	84.14
Gene5	70.45
Gene6	74.01
Gene7	85.39
Gene8	69.59
Gene9	67.15

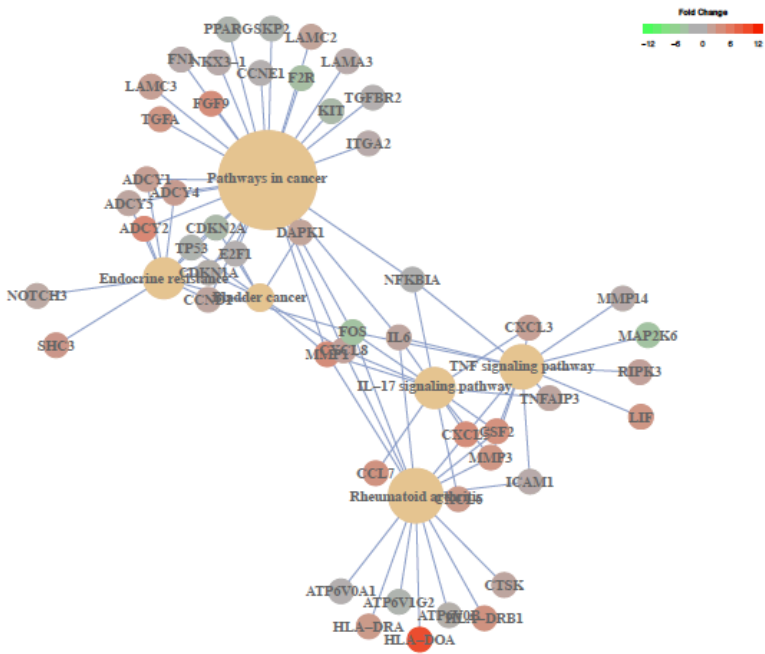
2018 Table 1 Percentage Oxidative Stress Relative to Control (from 2018 Figure 2)

CRISPR Knockout Library, Cells Sorted for Low Oxidative Stress

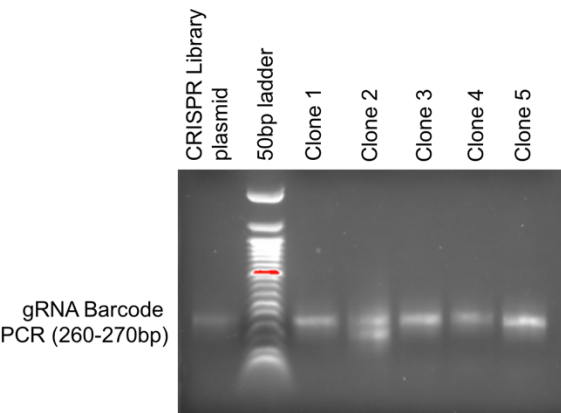


2018 Figure 3 CRISPR-Knockout Library Creates Subset of DC Fibroblasts with Significant Decrease in Oxidative Stress

2017 Progress Report Figures

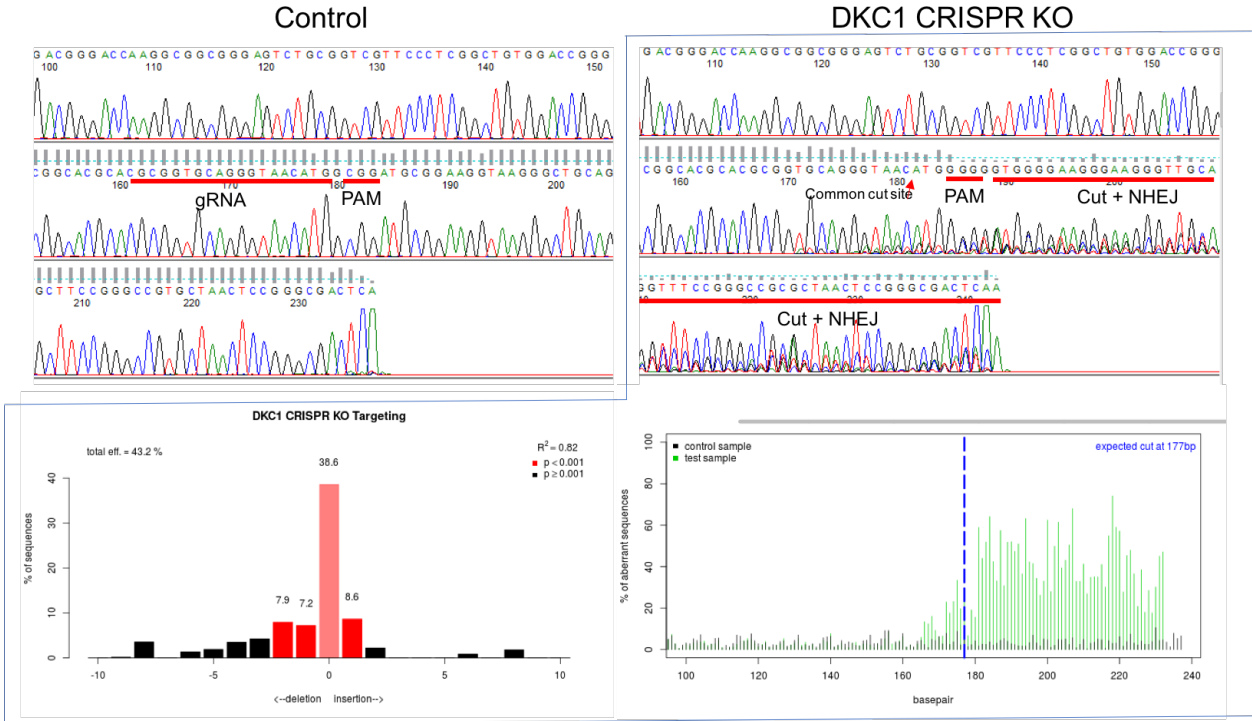


2017 Figure 1 KEGG Analysis of Genes Differentially Expressed in DC Vs Rescued Control Cells

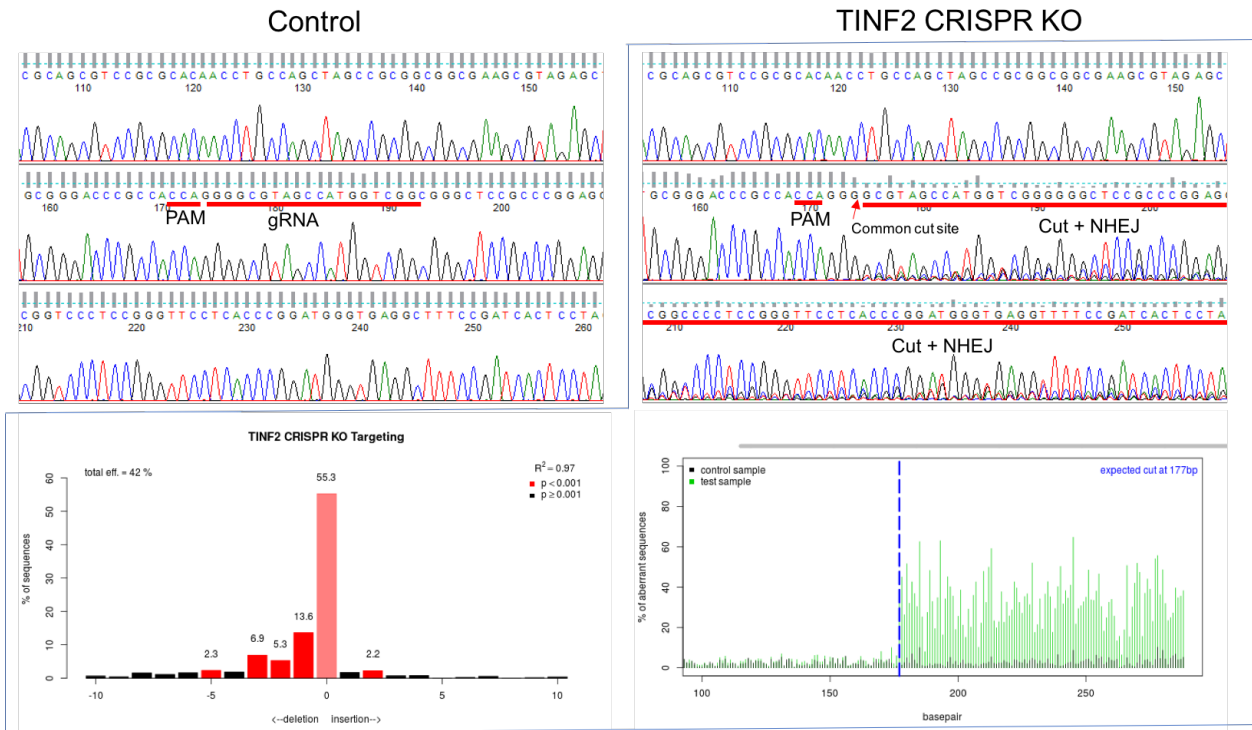


2017 Figure 3 PCR of gRNAs (Barcodes) from Cloned DC Fibroblasts Infected with CRISPR/Cas9 Library

2017 Progress Report Figures, continued

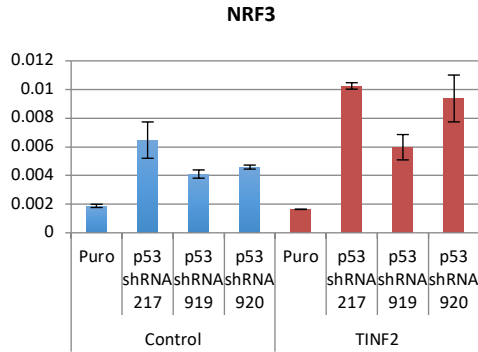


2017 Figure 4 Knockout of DKC1 in CD34+ Cells

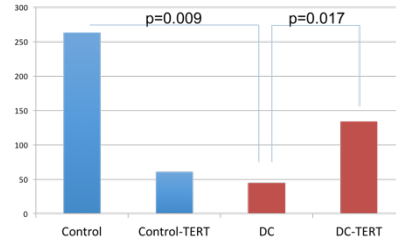


2017 Figure 5 Knockout of DKC1 in CD34+ Cells

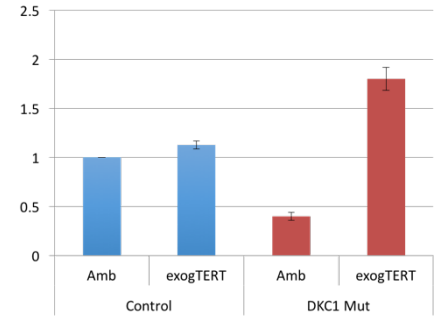
2016 Progress Report Figures



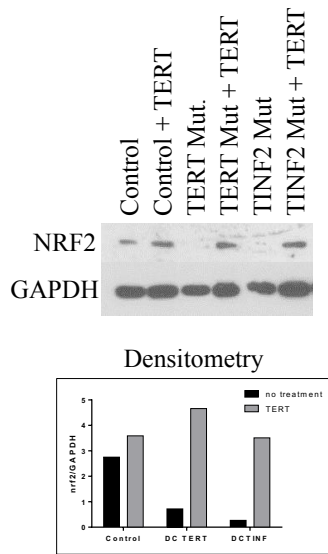
2016 Figure 4 De-repression of NRF3 Expression by p53 shRNA



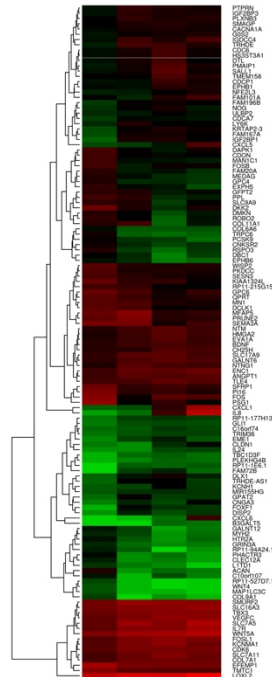
2016 Figure 2 Rescue of NRF3 Expression in TINF2-Mutated Cells Upon Exogenous TERT Expression (RNA-Seq)



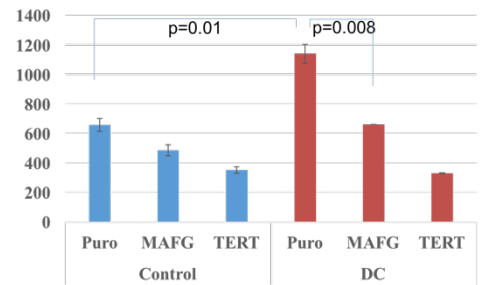
2016 Figure 3 Rescue of NRF3 Expression in DKC1-Mutated Cells Upon Exogenous TERT Expression (QRT-PCR)



2016 Figure 1 Increased NRF2 Protein in DC Cells Upon Exogenous TERT Express



2016 Figure 5 Hierarchical Clustering of Genes Differentially Expressed in Control and DC Cells and TERT-Expressing Cells



2016 Figure 6 Decreased Reactive Oxygen Species in DC Cells Expressing NRF2-Heterodimerizing Partner MAFG

Impact

The impact of this project is multifaceted given the nature of key elements found within this project. We are studying a bone marrow failure disorder that is caused by prematurely shortened telomeres. Telomere attrition and maintenance also have significant biological implications in aging as well as cancer. Furthermore, we have found that telomere shortening within these cells leads to a concomitant increase in reactive oxygen species which have also been implicated in aging and cancer. Through characterization of the signaling pathway by which telomere shortening leads to premature cell death, we can shed light on potential targets within the pathway that can be disrupted/augmented to prevent bone marrow failure. These same findings may also have implications for aging and cancer research.

Impact on other disciplines: Nothing to report

Impact on tech transfer: Nothing to report

Impact beyond science/technology: Nothing to report

Changes/Problems

2018

No major changes have been made from those made in 2017/2016

2017

We have outlined above issues related to the maintenance of an undifferentiated state in DC iPSCs in addition to the derivation of CD34⁺ cells. We have slightly altered the plan-of-action, as stated above, to alleviate these issues by:

1. We are repeating the production of iPSC clones to assess whether different clones may be more permissive for hematopoiesis
2. We are using CRISPR/Cas9 to knockout DC-related genes and alternatively induce DC-centric mutations into normal iPSCs that have been found to be permissive in regards to generating CD34⁺ cells
3. Circumvent iPSCs altogether and use CRISPR/Cas9 to directly knockout or mutate target DC-related genes in normal CD34⁺ cells

We feel that modification of normal cells to carry DC mutations or to knockout DC-related genes will serve as an appropriate alternative approach that will provide data that would be at parity, if not serve better than data generated from iPSCs.

2016

No significant changes have been made to the research. At this time the only significant, unanticipated problem is the inability to manufacture TINF2 iPSCs. We have a number of different methodologies (viral vs episomal; feeder vs feeder-free; addition of inhibitors or additional genes to derivation cocktail) to derive iPSCs and will modify our current strategies to produce these iPSCs.

Products: Nothing to report

Participants and Other Collaborating Organizations:

Name:	<i>Erik Westin</i>
Project Role:	<i>Postdoc</i>
Researcher Identifier (e.g. ORCID ID):	
Nearest person month worked:	12 months
Contribution to Project:	<i>Performed all major experiments</i>
Funding Support:	<i>DoD W81XWH-15-1-0099</i>

Name:	<i>Tim Townes</i>
Project Role:	<i>Collaborator</i>
Researcher Identifier (e.g. ORCID ID):	
Nearest person month worked:	2 months
Contribution to Project:	<i>Provided direction and project oversight</i>
Funding Support:	<i>DoD W81XWH-15-1-0099, Hyundai Hope on Wheels</i>

Name:	<i>Frederick Goldman</i>
Project Role:	<i>PI</i>
Researcher Identifier (e.g. ORCID ID):	
Nearest person month worked:	0.6 month
Contribution to Project:	<i>Provided direction and project oversight</i>
Funding Support:	<i>NIH, DoD W81XWH-15-1-0099</i>

Special Reporting Requirements: Nothing to report

Suppression of Antioxidant Responses in Dyskeratosis Congenita Cells

Erik Westin, PhD^{1*}, Larisa Pereboeva, MD, PhD², Divya Devadasan, MS^{1*}, Tim M. Townes, PhD³ and Frederick D Goldman, MD⁴

¹University of Alabama at Birmingham, Birmingham, AL; ²Univ. of Alabama At Birmingham, Birmingham, AL; ³Department of Biochemistry and Molecular Genetics, UAB Stem Cell Institute, University of Alabama at Birmingham, Birmingham, AL; ⁴Department of Pediatrics, Division of Hematology Oncology, Children's Hospital of Alabama, University of Alabama at Birmingham, Birmingham, AL

Dyskeratosis Congenita (DC) is a bone marrow failure disorder characterized by a triad of leukoplakia, skin dyspigmentation and nail dystrophy. Pathologies found in these patients arise due to mutations found within a number of genes (*DKC1*, *TERT*, *TERC*, *TINF2*, *TCAB1*, *CTCI*, *NOP10*, *C16orf57*, *NHP2* and *PARN*) that limit telomere maintenance/elongation, resulting in severely shortened telomeres. Previous studies in our lab have demonstrated impaired proliferation, limited lifespan and aberrant DNA damage response pathways in DC cells. These studies have also uncovered a significant reactive oxygen species (ROS) increase within every cell type investigated thus far. This ROS increase correlates with telomere dysfunction and the subsequent activation of the p53 DNA damage response pathway, which can be rescued by exogenous *TERT* or p53-shRNA expression. We have acquired skin punch biopsies from two patients with DC carrying either a *TERT* or *DKC1* mutation. Here, we have investigated a potential candidate pathway largely characterized as a key antioxidant regulator in hematopoietic cells, NRF2 (NFE2L2). NRF2 is a redox-sensitive basic leucine zipper transcription factor that, together with its heterologous partners (small MAF proteins, cJun, ATF, etc), binds to antioxidant response elements (AREs) within gene promoters in a pro-oxidant environment. We compared the RNA expression via QRT-PCR of *NRF2* in control and DC skin fibroblasts and found a significant reduction in DC cells (*TERT* mutation: 1.5 fold; *DKC1* mutation: 2.6 fold). Protein levels of NRF2 were also decreased in DC fibroblasts compared to controls. *TXN* is a gene whose expression is increased by NRF2 in a pro-oxidant environment. *TXN* expression was also significantly reduced (*TERT* mutation: 2.1 fold; *DKC1* mutation: 2.2 fold). To test whether *NRF2* suppression in DC cells is due to telomere dysfunction, we exogenously expressed *TERT* via retrovirus in DC and control fibroblasts. *TERT* expression led to dramatic increases in *NRF2* (*TERT* mutation: 3.4 fold, *DKC1* mutation: 3.7 fold) and *TXN* (*TERT* mutation: 3.7 fold, *DKC1* mutation: 1.6 fold). In contrast, *TERT* expression in control cells increased *NRF2* only 1.3 fold while *TXN* decreased 1.4 fold. Finally, we wanted to compare the expression of NRF2/*TXN* in low and elevated oxidative environments (4% vs 21% O₂). Control cells increased the *TXN* expression in 21% O₂ (*NRF2*: no change, *TXN*: 2.8 fold) while DC cells suppressed *NRF2* (*TERT* mutation: no change, *DKC1* mutation: 3 fold decrease) and *TXN* expression (*TERT* mutation: 1.4 fold decrease, *DKC1* mutation: 2.3 fold decrease). Functional studies have found DC cells grown in low oxygen increase their proliferative capacity perhaps due to, in part, the NRF2 pathway. Together, these data support a hypothesis whereby shortened/dysfunctional telomeres suppress NRF2 activity and an antioxidant response to a pro-oxidant environment. Based upon previous research, this pathway is likely dependent on the activation of p53 as an intermediary between dysfunctional telomere signaling and the subsequent suppression of NRF2 activity. An abrogated antioxidant response in shortened telomere cells may promote entry into senescence

and pathologies related to aging. Systemic pharmacological intervention that reduces ROS could reverse this process and form the basis to alleviate DC and related symptomology associated with this multi-organ disorder.

List of Personnel Receiving Pay for Research Effort

1. Erik Westin
2. Tim Townes
3. Frederick Goldman