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Figual Vacuu	EV 2012	Trada I a di II di I	EV 04/01/2012
Fiscal Year:	FY 2013	Task Last Updated:	F1 U4/U1/2U13
PI Name:	Ullrich, Robert Ph.D.	and an Datieria Control	
Project Title:	NSCOR: NASA Specialized Center of Resea	arch on Radiation Carcinogenesis	
Division Name:	Human Research		
Program/Discipline:	HUMAN RESEARCH		
Program/Discipline Element/Subdiscipline:	HUMAN RESEARCHRadiation health		
Joint Agency Name:		TechPort:	No
<b>Human Research Program Elements:</b>	(1) SR:Space Radiation		
Human Research Program Risks:	(1) Cancer: Risk of Radiation Carcinogenesis	s	
Space Biology Element:	None		
Space Biology Cross-Element Discipline:	None		
Space Biology Special Category:	None		
PI Email:	<u>bullrich@utmb.edu</u>	Fax:	FY
PI Organization Type:	UNIVERSITY	Phone:	409-747-1935
Organization Name:	University of Texas Medical Branch		
PI Address 1:	301 University Blvd		
PI Address 2:	Comprehensive Cancer Center, MS 1048		
PI Web Page:			
City:	Galveston	State:	TX
Zip Code:	77555-5302	<b>Congressional District:</b>	14
Comments:	NOTE: PI moved to UTMB from Colorado S	State University in late 2008 (6/2009	)
Project Type:	GROUND	Solicitation / Funding Source:	2008 NSCOR Space Radiation NNJ08ZSA003N
Start Date:	06/01/2009	End Date:	09/30/2015
No. of Post Docs:	2	No. of PhD Degrees:	0
No. of PhD Candidates:	0	No. of Master' Degrees:	0
No. of Master's Candidates:	0	No. of Bachelor's Degrees:	0
No. of Bachelor's Candidates:	0	<b>Monitoring Center:</b>	NASA JSC
Contact Monitor:	Simonsen, Lisa	Contact Phone:	
Contact Email:	lisa.c.simonsen@nasa.gov		
Flight Program:			
Flight Assignment:	NOTE: End date changed to 9/30/2015 (from 5/31/2015) per NSSC information (Ed., 6/1/15) NOTE: End date changed to 5/31/2015 per NSSC information (Ed., 3/3/14)		
Key Personnel Changes/Previous PI:	none		
COI Name (Institution):	Le Beau, Michelle (University of Chicago) Bacher, Jeff (Promega Corporation) Yu, Yongjia (University of Texas Medical Branch) Story, Michael (University of Texas Southwestern Medical Center at Dallas) Bedford, Joel (Colorado State University) Weil, Michael (Colorado State University) Ray, F (Colorado State University) Ding, Lianghao (University of Texas Southwestern Medical Center at Dallas) Xie, Yang (University of Texas Southwestern Medical Center)		
Grant/Contract No.:	NNX09AM08G		
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The goal of this NSCOR is to provide the information required to develop a rational scientific basis for estimation of risks for carcinogenesis in humans from exposure to radiation during space flight. Previous results from this Program found an unexpectedly low RBE value for acute myeloid leukemia (AML) induction by 1 GeV 56Fe ions. Systematic cytogenetic analyses suggested both microdosimetric factors related to the track structure of 1 GeV 56Fe ions and biological factors could account for this observation. In addition, these studies found an unexpected increase in hepatocellular carcinoma (HCC) at doses as low as 0.1 Gy of 1 GeV 56Fe ions but very little, if any, increase following gamma-ray exposure. These data suggest that processes associated with expansion and progression of initiated cells may play a more prominent role in HCC. If this is the case, it is possible that there are qualitative differences as well as quantitative in the effects of HZE irradiations. To expand on these results and to address the overall goal of this NSCOR a series of coordinated activities will conducted in 5 Projects and 3 Cores aimed at: (1) providing quantitative animal tumorigenesis data on the relative effectiveness of specific HZE particles and SPE protons compared with gamma-rays in mouse models of AML and HCC; (2) providing a better understanding of the impact of radiation exposure on the processes involved in the initiation and in the progression of initiated cells toward the neoplastic phenotype; 3) delineating potential differences between low LET radiation and high LET radiation such as those encountered in space travel on these processes; 4) developing links between animal data and radiation-induced effects for AML in humans; and (5) developing biologically-based modeling approaches which are critical to link these biological effects to risks in humans.

Program Overview: The Radiation Carcinogenesis NSCOR was initiated in June 2009 and builds upon results obtained in its predecessor, the Leukemogenesis NSCOR. The Radiation Carcinogenesis NSCOR consists of four projects supported by three cores. The projects and cores are briefly described below.

Project 1. Dose response relationships for induction of AML and HCC as a function of radiation quality (project leader, Dr. Robert. L. Ullrich). This project is designed to compare the effects of irradiation with gamma-rays, select HZE particles, and protons on the induction of AML and hepatocellular carcinoma (HCC) using the C3H murine model.

Project 2. Mechanisms of radiation leukemogenesis (project leader, Dr. Michael M. Weil). The goal of Project 2 is to better understand how radiation leads to AML in a murine model and to generate data for the development of a biologically based model that can be used to predict AML risks from various HZE or high energy proton exposures.

Project 3. Pathogenesis of radiation-induced hepatocellular carcinoma (project leader, Dr. Robert L. Ullrich). The overall hypothesis of this project is that the dose response is likely to reflect both quantitative as well as qualitative differences in high LET effects. This overall hypothesis will be tested in 3 specific aims:

- 1. Quantify the frequency and progression of preneoplastic foci (including both hyperplastic and dysplastic foci) in livers of C3H/HeNCrl mice irradiated with either 137Cs gamma-rays or HZE ions.
- 2. Examine irradiated liver for evidence of increased oxidative damage and alterations in the regulation of inflammatory processes.
- 3. Determine tumorigenic effects following HZE and gamma-ray irradiation in murine models of hepatocellular carcinoma in which secondary "promoting" events play a significant role.

Project 4. Molecular and cytogenetic targets in murine and human AML. (Project leader, Dr. Michelle Le Beau). This project is designed to develop a cytogenetic and molecular profile of human radiation-induced AML, leading to an understanding of the key events and genetic pathways involved in the pathogenesis of this disease.

Core A (Core Director, Dr. F. Andrew Ray). The Biology Core facilitates the distribution of irradiated and control animals, tissues, cells, and other biological samples to investigators. This core is also responsible for conducting the irradiations required at the various sites for all projects.

Core B (Core Director, Dr. Michael Story). The Genomics and Biostatistics core provides appropriate genomic analyses, innovative statistical modeling, simulations, and data analyses for the projects.

Core C (Core Director, Dr. Robert Ullrich). The Administrative Core provides administrative, fiscal and management support for the Radiation Carcinogenesis NSCOR. This core also oversees the overall scientific conduct of the NSCOR and facilitates interactions between projects, core leaders and project investigators as well as interactions with the internal and external advisors.

**Task Description:** 

# Rationale for HRP Directed Research:

## **Research Impact/Earth Benefits:**

This work will provide basic information on mechanisms of carcinogenesis as well as mechanisms specific to radiation-induced cancer.

## Task Progress:

Progress on this NSCOR has been made with respect to defining the dose response relationships for the induction of acute myeloid leukemia AML and hepatocellular carcinoma HCC using 600 MeV/n 56Fe and 350 MeV/n 28Si as well as gamma rays and SPE1972 protons. We have confirmed a low RBE for the induction of AML and a very high RBE for the induction of HCC. We have also been able to define in more depth mechanisms for the induction of radiation-induced AML in both mouse models and in cancer patients developing AML as a second cancer following treatment for a primary. Progress has also been made on defining important tissue changes in the liver that suggest that the high RBE for the induction of HCC is a result of promoting effects due to upregulation of inflammatory and cytokine pathways.

# Bibliography Type:

Description: (Last Updated: 07/25/2021)

## Articles in Peer-reviewed Journals

Sarkar S, Kantara C, Ortiz I, Swiercz R, Kuo J, Davey R, Escobar K, Ullrich R, Singh P. "Progastrin overexpression imparts tumorigenic/metastatic potential to embryonic epithelial cells: phenotypic differences between transformed and nontransformed stem cells. "Int J Cancer. 2012 Oct 1;131(7):E1088-99. Epub 2012 May 17. http://dx.doi.org/10.1002/ijc.27615; PubMed PMID: 22532325, Oct-2012

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Articles in Peer-reviewed Journals	Bielefeldt-Ohmann H, Genik PC, Fallgren CM, Ullrich RL, Weil MM. "Animal studies of charged particle-induced carcinogenesis." Health Physics. 2012 Nov;103(5):568-76. <a href="http://dx.doi.org/10.1097/HP.0b013e318265a257">http://dx.doi.org/10.1097/HP.0b013e318265a257</a> ; PubMed <a href="http://dx.doi.org/10.1097/HP.0b013e318265a257">PMID: 23032886</a> , Nov-2012
Articles in Peer-reviewed Journals	McNerney ME, Brown CD, Wang X, Bartom ET, Karmakar S, Bandlamudi C, Yu S, Ko J, Sandall BP, Stricker T, Anastasi J, Grossman RL, Cunningham JM, Le Beau MM, White KP. "CUX1 is a haploinsufficient myeloid tumor suppressor on chromosome 7 frequently inactivated in acute myeloid leukemia." Blood. 2013 Feb 7;121(6):975-83. Epub 2012 Dec 3. <a href="http://dx.doi.org/10.1182/blood-2012-04-426965">http://dx.doi.org/10.1182/blood-2012-04-426965</a> ; PubMed <a href="http://dx.doi.org/10.1182/blood-2012-04-426965">PMID: 23212519</a> , Feb-2013
Articles in Peer-reviewed Journals	Peng Y, Nagasawa H, Warner C, Bedford JS. "Genetic susceptibility: radiation effects relevant to space travel." Health Phys. 2012 Nov;103(5):607-20. <a href="http://dx.doi.org/10.1097/HP.0b013e31826945b9">http://dx.doi.org/10.1097/HP.0b013e31826945b9</a> ; PubMed <a href="PMID: 23032891">PMID: 23032891</a> , Nov-2012