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PI Name:	Brenner, David Ph.D.	
Project Title:	Physical and Biological Modulators of Space Radiation Carcinogenesis: Mechanistically- Bas Space Radiation Risk Assessment	ed Model Development for
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Program/Discipline:		
Program/Discipline Element/Subdiscipline:	HUMAN RESEARCHRadiation health	
Joint Agency Name:	TechPort:	Yes
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Human Research Program Risks:	(1) Cancer: Risk of Radiation Carcinogenesis	
Space Biology Element:	None	
Space Biology Cross-Element Discipline:	None	
Space Biology Special Category:	None	
PI Email:	djb3@cumc.columbia.edu Fax:	FY
PI Organization Type:	UNIVERSITY Phone:	(212) 305-5660
Organization Name:	Columbia University	
PI Address 1:	Center for Radiological Research	
PI Address 2:	630 W. 168th St.	
PI Web Page:		
City:	New York State:	NY
Zip Code:	10032 Congressional District:	13
Comments:		
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	This project is designed to use state-of-the-art mechanistic modeling of the experimental data from NASA Specialized Center of Research (NSCOR) programs and other available data as a basis to generate HZE (high energy particle) related cancer risk and uncertainty estimates in humans. There are four components: First, development of practical mechanistically motivated models, emphasizing the significance of individual radiation sensitivity. Second, based on model-based analysis of our and other NSCOR experimental data, estimate site-specific and consensus quality functions for HZE ions. Third, generate realistic uncertainty estimates for these estimates. Finally, our results and uncertainties will be critically compared with the current NASA projections and uncertainties. In order to answer the critical question of how to reliably estimate heavy ion-induced cancer risks in astronauts embarking on long-distance space exploration missions such as a flight to Mars, we are developing a	
Task Description:	mechanistically-motivated mathematical model that can predict radiogenic carcinogenesis as a	

	rate using both targeted effect (TE) and non-targeted effect (NTE) contributions. Importantly, such models are needed to predict low dose rate risks based on data at higher dose rates because the very low heavy ion dose rates relevant for space missions are difficult to achieve in terrestrial experiments. Our goal is to calibrate the carcinogenesis model using available human and animal data and to generate scaling factors such as the recently proposed radiation effects ratio (RER), which compares carcinogenic effectiveness of heavy ions and gamma rays at the dose of interest. The scaling factors would then be used to estimate human heavy ion-induced cancer risks, based on human gamma-ray-induced risks. An important focus of our work is generation of realistic uncertainties for model parameters and predictions, which ultimately translate into realistic uncertainties on astronaut risk estimates.
Rationale for HRP Directed Research:	This research is directed because it contains highly constrained research, which requires focused and constrained data gathering and analysis that is more appropriately obtained through a non-competitive proposal. The timing of this work supports current efforts by the Risk Assessment project to quantify uncertainties due to radiation quality factors and use of the dose and dose-rate effectiveness factor (DDREF). Work is highly synergistic with on-going work in the Fornace NSCOR as well as in assessing tissue-specific quality factors and DDREF specific to GI (gastronintestinal) cancers. The study will integrate data from multiple NSCORs (NASA Specialized Centers of Research).
Research Impact/Earth Benefits:	Cancer is the second leading cause of death in the United States, exceeded only by heart disease (<u>https://</u>). It accounts for one of every four deaths in the United States. More than 1.7 million new cancer cases and over 600,000 cancer-related deaths are predicted to occur in the US in 2018 (<u>https://</u>). Considering this high frequency and lethality of cancer, even a small increase by space radiation would have a major impact on planning and design of future interplanetary manned space missions. Accurate estimation of space radiation-related cancer risks is, therefore, very important for NASA mission planning.
Task Progress:	Heavy ion bombardment can be much more carcinogenic per unit dose than exposure to sparsely-ionizing radiation such as gamma-rays, and therefore heavy-ion induced carcinogenesis is an important challenge for long-distance human space exploration such as manned missions to Mars. Mechanistically-motivated mathematical models are needed to predict low dose rate risks relevant for space missions based on data at higher dose rates, which are more easily achievable in terrestrial experiments. We developed such a model, which quantifies targeted and non-targeted radiation effects. An important goal for using the model was to estimate dose rate effects (DRE) for heavy ion exposures at doses and exposure times relevant for a Mars mission, relative to much shorter (effectively acute) exposure times at the same dose. For tractable model development, we assumed that traversal of a cell by densely ionizing radiation such as the core of a heavy ion track can cause the release of NTE signals. Under continuous irradiation, such as during a space mission, the signal concentration is expected to quickly (in much less time than the duration of exposure) reach a steady-state equilibrium probability for cells susceptible to NTE signals to enter into and remain in an "activated" state, e.g., a state of perturbed signaling, altered gene expression, and/or oxidative stress. We assume, for simplicity, that two components contribute to the excess cancer yield due to NTE: (1) the effects of cell activation during irradiation, which docer are elevant for space missions) of heavy ions are also dose rate independent. We fitted the model to lung carcinogenesis data in radon-exposed miners and rats. These data sets are valuable because they provide information nature and animal lung carcinogenesis induced by protracted exposure to densely ionizing radiation and several types of space-relevant for space missions. The super the dows at dose rate explendent. We fitted the model to lung carcinogenesis data in radon-exposed miners and rats. These
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