FY 2021	Task Last Updated:	FY 12/01/2020
Costes, Sylvain Ph.D.		
Blood-based Multi-scale Model for Cance	r Risk from GCR in Gene	tically Diverse Populations
Human Research		
HUMAN RESEARCHRadiation health		
	TechPort:	No
(1) SR:Space Radiation		
(1) Cancer: Risk of Radiation Carcinogene	esis	
None		
None		
None		
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NOTE: After retiring from NASA, the PI j Laboratory until December 2016.	oined Blue Marble. Dr. C	Costes was previously at Lawrence Berkeley National
Ground	Solicitation / Funding Source:	2014-15 HERO NNJ14ZSA001N-RADIATION. Appendix D: Ground-Based Studies in Space Radiobiology
02/04/2016	End Date:	10/01/2021
1	No. of PhD Degrees:	3
0	No. of Master' Degrees:	
1	No. of Bachelor's Degrees:	
1	Monitoring Center:	NASA ARC
Zawaski, Janice	Contact Phone:	
janice.zawaski@nasa.gov		
NOTE: End date changed to 10/1/2021 per		
NOTE: End date changed to 9/30/2021 per	r L.Lewis/ARC HRP (Ed.	, 12/9/20)
NOTE: End date changed to 9/30/2021 per NOTE: Extended to 5/31/2021 per L. Lew	`	· · · ·
0 1	is/ARC HRP (Ed., 9/24/2	· · · ·
	Costes, Sylvain Ph.D. Blood-based Multi-scale Model for Cancer Human Research HUMAN RESEARCHRadiation health (1) SR:Space Radiation (1) Cancer:Risk of Radiation Carcinogene None None None None Sylvain.costes@bmsis.org GOVERNMENT Blue Marble Space Institute of Science 600 1st Avenue, 1st Floor Blue Marble Space Institute of Science 600 1st Avenue, 1st Floor Seattle 98104 NOTE: After retiring from NASA, the PI J Laboratory until December 2016. Ground 02/04/2016 1 0 1 1 2 1 2 2 awaski, Janice janice.zawaski@nasa.gov	Costes, Sylvain Ph.D. Blood-based Multi-scale Model for Cancer Risk from GCR in General Human Research HUMAN RESEARCHRadiation health IUMAN RESEARCHRadiation health II Cancer:Risk of Radiation Carcinogenesis None None None None None Sylvain.costes@bmsis.org Fax: GOVERNMENT Phone: Blue Marble Space Institute of Science G00 1st Avenue, 1st Floor Seattle Statte:

Key Personnel Changes/Previous PI:	NOTE (January 2018): The lab moved from Lawrence Berkeley National Lab (LBNL) to NASA Ames Research Center in 2017, where it was established as the Radiation Biophysics Lab in Space Biosciences Division. Dr. Costes will continue collaborating with LBNL and some funding will be left at LBNL to cover more plate processing in collaboration with Dr. Weil, Colorado State University (CSU) and for support from Dr. Snijders for the writing of the animal data. April 2017 report: - Elodie Guiet was a full time technician with a Bachelor in microbiology and biotechnology, working on this project from March 2016 until February 2017 she did not stay on the project when the lab moved to NASA Ames ; - Louise Viger was a Postdoc working partly on this project from June 2016 to January 2017 she was only here for a quick postdoc, focused primarily on modeling ; - Charlotte Degorre was a Postdoc who helped executing BNL run 16C visiting scientist for 1 month ; - Sebastien Penninckx was a PhD student who has been helping on data analysis visiting scientist for 3 months ; - Shayoni Ray is a new recruit at NASA Ames, postdoctoral fellow working on doing genomic analysis between animal DNA repair phenotypic data and their individual genes new postdoc full time at NASA Ames, started on April 10 2017.
COI Name (Institution):	
Grant/Contract No.:	Internal Project-ARC ; NNJ16HP24I
Performance Goal No.:	
Performance Goal Text:	
Task Description:	NOTE (January 2018): The lab moved from Lavrence Berkeley National Lab (LBNL) to NASA Ames Research Center in 2017, where it was established as the Radiation Biophysics Lab in Space Biosciences Division. Dr. Costes will continue collaborating with LBNL and some funding will be left at LBNL to cover more plate processing in collaboration with Dr. Weil, Colorado State University (CSU), and for support from Dr. Snijders for the writing of the animal data. Crew on future exploration missions to Mars and other destinations in our solar system will be exposed to acute low doses (<100 mSV) and chronic low doses (<0.1 mSV/min) of high-LET (linear energy transfer) ionizing radiation from solar particle events (SPE) and galactic cosmic radiation (GCR). Predicting cancer risk associated with these radiation types is a mission-critical challenge for NASA radiation health scientists and mission planners. Epidemiological methods lack sensitivity and power to provide detailed risk estimates for cancer, mainly because the number of exposed individuals to date is relatively small, limited to several hundred individual sexposed to trapper adiation in low Earth orbit and fewer than two dozen Apollo astronauts exposed to GCR for several days at a time. Moreover, population-based studies do not take individual radiation sensitivity into account, are sensitive to the presence of other confounding environmental insults, and require long follow-up times. In collaboration with the radiation Biodosimetry Laboratory and the modeling group at NASA Johnson Space Center and with the International Computer Science Institute (ICSI) at University of California (UC) Berkeley, our team will archived by NASA under the various Human Research Program (HRP) funded projects. The main goal of this proposal is to identify factors influencing radiation-induced carcinogenesis and integrate them into a multi-scale model already fary types of HZE (high energy particels). Because expremimental data are dispered facross may different cancer models, radiation
Rationale for HRP Directed Research	:
Research Impact/Earth Benefits:	A current radiobiology challenge is the ability to predict cancer risk associated with exposure to acute (<100 mSv) and chronic (<0.1 mSv/min) low doses of high-LET ionizing radiation. Epidemiological methods lack the sensitivity and power to provide detailed risk estimates for cancer, mainly because the astronaut cohort exposed to galactic cosmic rays (GCR) is relatively small. Moreover, population-based studies do not take individual radiation sensitivity into account, are affected by the presence of other confounding environmental insults, and require long follow-up times. We have hypothesized that characterizing the dose and time dependence of 53BP1 radiation induced foci (RIF) after exposure to a systematic array of X-ray doses and time points is sufficient to describe someone's ability to respond to any other LET. The main concept is that the non-physiological response to high doses of radiation is modulated by different pools of genes. Such work provides a new approach combining novel biomarkers with sophisticated mathematical analysis to better characterize individual sensitivity to space radiation. Once validated across mice and eventually a large cohort of humans, this approach could be generalized to establish individualized health risk management for astronauts and for the population at large being exposed to ionizing radiation.

We are continuing sample processing and analysis to uncover the genomic associations with human ex vivo immune cell responses to simulated space radiation. All mouse sample analysis has been completed. All human DNA samples have been successfully sequenced using low-throughput whole-genome sequencing, and the sequencing results are in the process of being analyzed. Human immune cell analysis has been severely delayed by shelter-in-place, which contributed to sample degeneration, so this experiment will have to be partially repeated in the next Brookhaven National Laboratory (BNL) run to finish all sample collection and analysis. We have published 6 peer-reviewed articles based on mouse and human data analysis, with more articles in preparation. We have presented our work in talks and posters in the NASA Human Research Program (HRP) Investigators' Workshop (presentations: Sylvain Costes, Eloise Pariset) and the Radiation Research Society Annual Meeting (presentation: Sylvain Costes, posters: Eloise Pariset, Egle Cekanaviciute) and have been accepted to present it again at COSPAR (Committee on Space Research) 2021 (invited presentation: Sylvain Costes).
Furthermore, we have been awarded a NASA HRP Tissue Sharing grant ("Mapping peripheral immune signatures of mouse and human responses to space radiation for biomarker identification," PI: Costes, Co-I/Science PI: Cekanaviciute, Co-I: Pariset) to follow up on this work. We will utilize the samples from this project together with samples collected by Dr. Susanna Rosi's group on her HRP-funded project on mouse neuroimmune responses to combined exposures to ionizing radiation and simulated microgravity, in order to identify a shared signature of immune responses to spaceflight that could be utilized for biomarker and countermeasure development.
Description: (Last Updated: 05/01/2025)
Penninckx S, Pariset E, Acuna AU, Lucas S, Costes SV. "Considering cell proliferation to optimize detection of radiation-induced 53BP1-positive foci in 15 mouse strains ex vivo." Radiat Res. 2021 Jan 1;195(1):47-59. https://doi.org/10.1667/RADE-20-00165.1 ; PMID: 33181852 , Jan-2021
Pariset E, Penninckx S, Kerbaul CD, Guiet E, Macha AL, Cekanaviciute E, Snijders AM, Mao JH, Paris F, Costes SV. "53BP1 repair kinetics for prediction of in vivo radiation susceptibility in 15 mouse strains." Radiat Res. 2020 Nov 10;194(5):485-99. <u>https://doi.org/10.1667/RADE-20-00122.1</u> ; <u>PMID: 32991727</u> , Nov-2020
Pariset E, Malkani S, Cekanaviciute E, Costes SV. "Ionizing radiation-induced risks to the central nervous system and countermeasures in cellular and rodent models." Int J Radiat Biol. Published online: 20 Oct 2020. https://doi.org/10.1080/09553002.2020.1820598 ; PMID: 32946305 , Oct-2020
Pariset E, Bertucci A, Petay M, Malkani S, Lopez Macha A, Paulino Lima IG, Gomez Gonzalez V, Tin AS, Tang J, Plante I, Cekanaviciute E, Vazquez M, Costes SV. "DNA damage baseline predicts resilience to space radiation and radiotherapy." Cell Rep. 2020 Dec 8;33(10:108434. <u>https://doi.org/10.1016/j.celrep.2020.108434</u> ; <u>PMID: 33242409</u> , Dec-2020
Nikitaki Z, Pariset E, Sudar D, Costes SV, Georgakilas AG. "In situ detection of complex DNA damage using microscopy: A rough road ahead." Cancers (Basel). 2020 Nov 6;12(11):E3288. Review. https://doi.org/10.3390/cancers12113288 ; PMID: 33172046; PMCID: PMC7694657, Nov-2020
Afshinnekoo E, Scott RT, MacKay MJ, Pariset E, Cekanaviciute E, Barker R, Gilroy S, Hassane D, Smith SM, Zwart SR, Nelman-Gonzalez M, Crucian BE, Ponomarev SA, Orlov OI, Shiba D, Muratani M, Yamamoto M, Richards SE, Vaishampayan PA, Meydan C, Foox J, Myrrhe J, Istasse E, Singh N, Venkateswaran K, Keune JA, Ray HE, Basner M, Miller J, Vitaterna MH, Taylor DM, Wallace D, Rubins K, Bailey SM, Grabham P, Costes SV, Mason CE, Beheshti A. "Fundamental biological features of spaceflight: Advancing the field to enable deep-space exploration." Cell. 2020 Nov 25;183(5):1162-84. Review. <u>https://doi.org/10.1016/j.cell.2020.10.050</u> ; <u>PMID: 33242416</u> , Nov-2020