Fiscal Year:	FY 2017	Task Last Updated:	FY 07/18/2017
PI Name:	Bigley, Austin Ph.D.		
Project Title:	The Role of Microgravity and Stress	s-related Humoral Factors in Dysregulat	ed NK-cell Function during Spaceflight
Division Name:	Human Research		
Program/Discipline:	NSBRI		
Program/Discipline Element/Subdiscipline:	NSBRIRadiation Effects Team		
Joint Agency Name:		TechPort:	No
Human Research Program Elements:	None		
Human Research Program Risks:	None		
Space Biology Element:	None		
Space Biology Cross-Element Discipline:	None		
Space Biology Special Category:	None		
PI Email:	abbigley@uh.edu	Fax:	FY
PI Organization Type:	UNIVERSITY	Phone:	7138286634
Organization Name:	University of Houston		
PI Address 1:	3875 Holman St.		
PI Address 2:	Rm 104		
PI Web Page:			
City:	Houston	State:	TX
Zip Code:	77204-6015	Congressional District:	18
Comments:			
Project Type:	Ground	Solicitation / Funding Source:	2015 NSBRI-RFA-15-01 First Award Fellowships
Start Date:	10/01/2015	End Date:	03/31/2017
No. of Post Docs:	1	No. of PhD Degrees:	0
No. of PhD Candidates:	1	No. of Master' Degrees:	0
No. of Master's Candidates:	0	No. of Bachelor's Degrees:	0
No. of Bachelor's Candidates:	0	Monitoring Center:	NSBRI
Contact Monitor:		Contact Phone:	
Contact Email:			
Flight Program:			
Flight Assignment:)17 (previously 5/31/2017) per NSBRI ()17 (previously 9/30/2016) per NSBRI (
Key Personnel Changes/Previous PI:			
COI Name (Institution):	Simpson, Richard (MENTOR/Un	iversity of Houston)	
Grant/Contract No.:	NCC 9-58-PF04307		
Performance Goal No.:			
Performance Goal Text:			

Task Description:	POSTDOCTORAL FELLOWSHIP Before we can ethically send Astronauts to distant locations in space, such as Mars or asteroids, it is critical that we understand how spaceflight affects the human immune system. For years, post-flight data has suggested that spaceflight has a negative impact on the immune system. Unfortunately, the majority of this data was obtained following short-duration missions after the Astronauts had returned to Earth, meaning that no in-flight data was collected. The recent 'Integrated Immune' study filled many of these knowledge gaps by analyzing blood obtained from Astronauts on the Space Shuttle (or International Space Station (ISS)) while they were still in space. Of particular interest were the findings that anti-viral immune responses were compromised during spaceflight. Consequently, latent viral reactivation can result in many negative consequences including Shingles, reduced vaccine efficacy, and increased susceptibility to infection. While the 'Integrated Immune' study focused primarily on the adaptive immune system, our 'Salivary Markers' flight experiment (Principal Investigator Dr. Richard Simpson) has focused on the innate immune system. The current proposal focuses on Natural Killer (NK)-cells, which are able to kill virally-infected and malignant cells without prior exposure. Our in-flight data shows that NK-cell anti-tumor activity is greatly reduced during spaceflight, while cytomegalovirus (CMV)-driven responses are amplified. The current proposal will explore two plausible mechanisms for these observationsmicrogravity and stress. The effects of microgravity can be simulated with a rotating wall vessel that keeps the cells in a constant state of freefall, while stress can be simulated microgravity and spaceflight-drived serum will mimic the deleterious effects of spaceflight on NK-cells. Once we have determined the mechanisms underpinning the adverse effects of spaceflight on NK-cells, we can begin to develop countermeasures that will protect future space explorers from be
Rationale for HRP Directed Researc	h:
Research Impact/Earth Benefits:	Our research has extensive Earth benefits. Using NASA funding, we have gained insight into many of the mechanisms governing NK-cell activity during spaceflight including CMV reactivation and stress. Many of the mechanisms of immunosuppression at play during spaceflight are also found on Earth particularly in the setting of solid organ or hematopoietic stem cell transplantation. CMV plays a major role in spaceflight-associated alterations to immune function and it also plays a major role in transplant outcome and cancer immunology. The findings that we have made through NASA and National Space Biomedical Research Institute (NSBRI) funding have led to two published first-author manuscripts during my NSBRI First Award Fellowship. These manuscripts focus on the beneficial effects of CMV on NK-cell-mediated cytotoxicity against hematologic malignancies and the mechanisms underpinning this anti-tumor effect. This research has resulted in an NIH R21 grant that will enable us to determine the effects of CMV seropositivity on multiple myeloma remission rates and also enable us to develop new ex vivo expansion protocols that selectively expand tumor-reactive NK-cells (like we see with CMV) for adoptive transfer without the negative side effects and decreased mortality associated with CMV reactivation in vivo.
Task Progress:	We have provided major insights into the mechanisms underpinning the impaired NK-cell-mediated immunity observed during spaceflight. Specifically, we have shown that simulated microgravity impairs NK-cell function similarly to spaceflight and identified the specific cellular adaptations responsible, thus enabling the development of countermeasures to help maintain immunity during exploration class missions. Our findings have contributed to closing gaps Immune (IM) IM1, IM2, and IM8. For IM1, we have shown that spaceflight inhibits NK-cell function and we have shown that this inhibition can be recapitulated using simulated microgravity as an analog. We have also identified the specific cellular adaptations responsible for the microgravity-induced inhibition of NK-cells. For IM2, we have shown that an improved immunological standard is necessary for spaceflight. Specifically, NK-cell function will be monitored as impaired NK-cell function has been linked to increased cancer risk. Maintaining NK-cell function will be particularly important for long-duration space exploration when astronauts are being exposed to high levels of cosmic radiation for long periods of time. For IM8, our serum stimulation experiment will allow us to mechanistically link stress (through hormones and cytokines present in serum) during spaceflight to impaired NK-cell function.
Bibliography Type:	Description: (Last Updated: 03/12/2021)
Articles in Peer-reviewed Journals	Bigley AB, Rezvani K, Shah N, Sekine T, Balneger N, Pistillo M, Agha N, Kunz H, O'Connor DP, Bollard CM, Simpson RJ. "Latent cytomegalovirus infection enhances anti-tumour cytotoxicity through accumulation of NKG2C+ NK cells in healthy humans." Clinical Experimental Immunology. 2016 Aug;185(2):239-51. https://doi.org/10.1111/cei.12785; PubMed PMID: 26940026; PubMed Central PMCID: PMC4955006, Aug-2016
Articles in Peer-reviewed Journals	Bigley AB, Spielmann G, Agha N, O'Connor DP, Simpson RJ. "Dichotomous effects of latent CMV infection on the phenotype and functional properties of CD8+ T-cells and NK-cells." Cell Immunol. 2016 Feb;300:26-32. Epub 2015 Nov 24. <u>https://doi.org/10.1016/j.cellimm.2015.11.005</u> ; PubMed <u>PMID: 26651951</u> , Feb-2016
Articles in Peer-reviewed Journals	Simpson RJ, Bigley AB, Agha N, Hanley PJ, Bollard CM. "Mobilizing immune cells with exercise for cancer immunotherapy." Exerc Sport Sci Rev. 2017 Jul;45(3):163-72. <u>https://doi.org/10.1249/JES.000000000000114</u> ; PubMed <u>PMID: 28418996</u> , Jul-2017
Articles in Peer-reviewed Journals	Gupta P, Bigley AB, Markofski M, Laughlin M, LaVoy EC. "Autologous serum collected 1h post-exercise enhances natural killer cell cytotoxicity." Brain Behav Immun. 2018 Jul;71:81-92. <u>https://doi.org/10.1016/j.bbi.2018.04.007</u> ; <u>PMID: 29656052</u> , Jul-2018

Articles in Peer-reviewed Journals	Bigley AB, Agha NH, Baker FL, Spielmann G, Kunz HE, Mylabathula PL, Rooney B, Laughlin MS, Pierson DL, Mehta SK, Crucian BE, Simpson RJ. "NK-cell function is impaired during long-duration spaceflight." J Appl Physiol (1985). 2019 Apr 1;126(4):842-53. Epub 2018 Nov 1. PubMed <u>PMID: 30382809</u> ; https://doi.org/10.1152/japplphysiol.00761.2018, Apr-2019	
Articles in Peer-reviewed Journals	Mylabathula PL, Li L, Bigley AB, Markofski MM, Crucian BE, Mehta SK, Pierson DL, Laughlin MS, Rezvani K, Simpson RJ. "Simulated microgravity disarms human NK-cells and inhibits anti-tumor cytotoxicity in vitro." Acta Astronaut. 2020 Sep;174:32-40. <u>https://doi.org/10.1016/j.actaastro.2020.03.023</u> , Sep-2020	