

<b>Fiscal Year:</b>	FY 2022	<b>Task Last Updated:</b>	FY 09/23/2021
<b>PI Name:</b>	Ott, C. Mark Ph.D.		
<b>Project Title:</b>	Spaceflight-Induced Changes in Microbial Virulence and Impact to the Host Immune Response		
<b>Division Name:</b>	Human Research		
<b>Program/Discipline:</b>			
<b>Program/Discipline-- Element/Subdiscipline:</b>			
<b>Joint Agency Name:</b>		<b>TechPort:</b>	No
<b>Human Research Program Elements:</b>	(1) <b>HHC:</b> Human Health Countermeasures		
<b>Human Research Program Risks:</b>	(1) <b>Microhost:</b> Risk of Adverse Health Effects Due to Host-Microorganism Interactions		
<b>Space Biology Element:</b>	None		
<b>Space Biology Cross-Element Discipline:</b>	None		
<b>Space Biology Special Category:</b>	None		
<b>PI Email:</b>	<a href="mailto:c.m.ott@nasa.gov">c.m.ott@nasa.gov</a>	<b>Fax:</b>	FY
<b>PI Organization Type:</b>	NASA CENTER	<b>Phone:</b>	281-483-7155
<b>Organization Name:</b>	NASA Johnson Space Center		
<b>PI Address 1:</b>	2101 NASA Parkway, SF24		
<b>PI Address 2:</b>			
<b>PI Web Page:</b>			
<b>City:</b>	Houston	<b>State:</b>	TX
<b>Zip Code:</b>	77058	<b>Congressional District:</b>	36
<b>Comments:</b>			
<b>Project Type:</b>	Ground	<b>Solicitation / Funding Source:</b>	Directed Research
<b>Start Date:</b>	10/01/2019	<b>End Date:</b>	09/30/2025
<b>No. of Post Docs:</b>		<b>No. of PhD Degrees:</b>	
<b>No. of PhD Candidates:</b>	1	<b>No. of Master' Degrees:</b>	
<b>No. of Master's Candidates:</b>		<b>No. of Bachelor's Degrees:</b>	
<b>No. of Bachelor's Candidates:</b>		<b>Monitoring Center:</b>	NASA JSC
<b>Contact Monitor:</b>	Brocato, Becky	<b>Contact Phone:</b>	
<b>Contact Email:</b>	<a href="mailto:becky.brocato@nasa.gov">becky.brocato@nasa.gov</a>		
<b>Flight Program:</b>			
<b>Flight Assignment:</b>			
<b>Key Personnel Changes/Previous PI:</b>			
<b>COI Name (Institution):</b>	Nickerson, Cheryl Ph.D. ( CoPI-- Arizona State University grant 80NSSC20K0016 ) Barrila, Jennifer Ph.D. ( Arizona State University ) Oubre, Cherie Ph.D. ( NASA Johnson Space Center ) Crucian, Brian Ph.D. ( NASA Johnson Space Center )		
<b>Grant/Contract No.:</b>	Directed Research		
<b>Performance Goal No.:</b>			
<b>Performance Goal Text:</b>			

<p><b>Task Description:</b></p>	<p>One of the critical factors to ensure crew health, safety, and performance is anticipating the risk for infectious disease during human deep space exploration and habitation. In 2006 and 2007, our spaceflight experiments aboard the Space Shuttle and International Space Station (ISS) demonstrated that the foodborne pathogen, <i>Salmonella enterica</i> serovar Typhimurium (S. Typhimurium), increased its virulence in response to culture in the spaceflight environment. These findings were in agreement with our initial studies using spaceflight analogue conditions for this same organism. Since those experiments, unexpected alterations in other microbial pathogen characteristics that may or may not be related to disease have also been documented in response to both spaceflight and spaceflight analogue culture. However, the range of bacteria whose true virulence in animals is altered in response to spaceflight (and spaceflight analogue) culture and the degree to which their virulence may be altered remains poorly understood.</p> <p>In parallel to the observations regarding microgravity exposure and bacterial virulence, multiple studies onboard ISS have documented and characterized the dysregulation of the human immune system associated with spaceflight. The phenomenon is generally characterized by altered leukocyte reductions in T and Natural Killer (NK) cell function, altered plasma cytokine profiles, and the reactivation of latent herpesviruses. Most of these studies were generally performed by returning astronaut biosamples for evaluation. Akin to microbial studies, spaceflight analogue cell culture is also well established as a terrestrial analogue that mimics key aspects of microgravity on immune cell activation. For example, both T and NK cells exhibit inhibited activation during spaceflight analogue cell culture.</p> <p>Unknown, and almost completely uninvestigated, is the possible synergistic effect between increased microbial virulence and reductions in immune cell function during microgravity/spaceflight conditions. Should virulence increase for multiple pathogens during crewed deep space missions, synergy with diminished immunity could increase crew health risk for infectious diseases during pending missions of exploration. The NASA Human Research Program created a specific 'Knowledge Gap' in their Integrated Research Plan regarding this issue, but to date, no study has provided an integrated systematic approach to address this Gap.</p> <p>Using spaceflight analogue technology, the proposed studies will incorporate microbial and mammalian cell culture, animal studies, and ISS crew immune cell studies in an integrated systematic approach to better understand how these systems shape the dynamics of the host-pathogen interaction that lead to infectious disease in microgravity conditions. Our hypothesis is that the incidence of higher virulence observed in both spaceflight and spaceflight analogue culture for the foodborne pathogen S. Typhimurium is not limited to this organism, and that multiple bacteria will exhibit similar increases in virulence when cultured under spaceflight analogue conditions. We further hypothesize that spaceflight-induced alterations in crew immune cell function will lead to compromised defenses against pathogen infection, which when combined with alterations in microbial virulence, will lead to a synergistic response that will reflect greater risk to crew health.</p> <p>Accordingly, the goal of this study is to gain an understanding of medically important microorganisms relevant to crew health that exhibit altered virulence and pathogenesis-related properties and the impact of those changes on the crew immune cell response using spaceflight analogue culture conditions. This study will incorporate relevant ISS bacterial pathogens that have either been identified from operational microbial monitoring activities or that have a clear route of infection for the crew, including potential foodborne pathogens applicable to future development of bioregenerative food systems.</p>
<p><b>Rationale for HRP Directed Research:</b></p>	<p>The MicroHost research plan aims to determine which microorganisms develop altered virulence when exposed to spaceflight conditions and understand the synergistic effect of altered microbial virulence and dysregulated immunity on crew health risks for deep space missions.</p> <p>Insufficient time for solicitation: Continued delays in initiating the proposed study will continue to impact the schedule and decrease our likelihood of gaining the knowledge needed to close the risk. Note that the delay in this work may impact the Path to Risk Reduction (PRR) color change from yellow to green and put the studies outside of the window for use of the International Space Station (ISS). Two prior solicitations have been released (in 2009 and 2014) for ground-based proposals to understand microbial responses to simulated microgravity. Even though the prior solicitations were written clearly, the selected studies did not focus on identifying the microbial alterations that would gain the understanding needed to inform the risk, and they did not produce the needed ground-based investigations on mechanisms. The 2009 selection addresses collective changes of organisms within the human microbiome, and the 2014 selection addresses viral reactivation. The selected studies will provide information applicable to the gaps Micro-101 to better understand the potential impact of microgravity on microbial virulence and Micro-201 to better understand the contribution of these changes on adverse health events. Completion of the proposed work will provide clear evidence as to the operational applicability of these original microbial virulence data to a variety of microorganisms and will include measurements of host immune responses to microbial challenge.</p> <p>Access to Previous Crew Data: This proposed study will leverage previous microbiology operational and research data as well as previously published immunology research data to provide a better understanding of impacts of microbial changes to the host and to determine the need for countermeasure evaluation as outlined in our PRR.</p>
<p><b>Research Impact/Earth Benefits:</b></p>	<p>This research will enrich life on Earth through the use of space technology and the application of biomedical knowledge. Specifically, this study will utilize the microgravity spaceflight platform to 1) to broaden our knowledge of the host-pathogen interaction that leads to infectious disease, and 2) for the development of new therapeutic strategies to combat infectious disease for the general public.</p>
	<p>Work at Arizona State University (ASU) on the 5 test microorganisms to support this project are listed below and have included extensive interactions via routine Zoom video telecons and emails with the Principal Investigator and Co Principal Investigator and external Consultants:</p> <p><i>Salmonella Enteritidis</i>:</p> <ul style="list-style-type: none"> <li>• Characterization of spaceflight analogue culture on growth kinetics, stress responses, and in vitro infection of 3-D biomimetic intestinal cell culture models, and fixation of samples for RNA has been completed.</li> </ul> <p><i>Pseudomonas aeruginosa</i>:</p> <ul style="list-style-type: none"> <li>• Growth kinetics completed for this organism multiple times for statistical relevance; • Initiating RNA isolation</li> </ul>

<b>Task Progress:</b>	<p>Burkholderia cepacia:</p> <ul style="list-style-type: none"> <li>• Growth kinetics completed multiple times for statistical relevance ;</li> <li>• Several in vitro stress assays performed multiple times for statistical relevance.</li> </ul> <p>Streptococcus pneumoniae:</p> <ul style="list-style-type: none"> <li>• Optimization of growth conditions and media requirements for Streptococcus pneumoniae.</li> </ul> <p>Enterohemorrhagic E. coli:</p> <ul style="list-style-type: none"> <li>• Growth kinetics completed multiple times for statistical relevance ;</li> <li>• Several in vitro stress assays performed multiple times for statistical relevance.</li> </ul> <p>ASU Institutional Animal Care and Use Committee (IACUC) approval has been obtained for all animal infection studies using respiratory pathogens.</p> <p>Participation of NASA Johnson Space Center (JSC) in animal infection studies using intestinal pathogens has been discussed with JSC IACUC.</p> <p>JSC IACUC panel review of animal studies occurred in summer 2021.</p> <p>ASU supplied four test organisms to NASA JSC in support of the Immunological component of this study.</p> <p>ASU provided technical support to JSC for growth conditions in support of the Immunological component of this study.</p>
<b>Bibliography Type:</b>	<p>Description: (Last Updated: 10/14/2024)</p>
<b>Abstracts for Journals and Proceedings</b>	<p>Krieger S, Madedonas G, Mehta S, Rooney B, Nelman M, Castro C, Colorado A, Ott CM, Barrila J, Stafford P, Nickerson CA, Oubre C, Crucian B. "Microgravity influence on bacterial pathogen virulence and immune cell function –relevance for spaceflight infectious disease risk." 2020 NASA Human Research Program Investigators' Workshop, Galveston, TX, January 27-30, 2020.</p> <p>Abstracts. 2020 NASA Human Research Program Investigators' Workshop, Galveston, TX, January 27-30, 2020. , Jan-2020</p>
<b>Abstracts for Journals and Proceedings</b>	<p>Nickerson C. "Infectious Diseases and Global Health Training Program." Invited speaker, Infectious Diseases and Global Health Training Program, University of Manitoba, Nov 26, 2020.</p> <p>Infectious Diseases and Global Health Training Program, University of Manitoba, Nov 26, 2020. Presentation. , Nov-2020</p>
<b>Articles in Other Journals or Periodicals</b>	<p>Nickerson CA, Colorado A, Barrila J, Poste G, Ott CM. "A Vision for the Next Generation of Spaceflight Microbiology: Human Health and Habitat Sustainability. Invited Review. " Nature Microbiology. In revision as of September 2021. , Sep-2021</p>