Fiscal Year:	FY 2014		
	11 2014	Task Last Updated:	FY 12/04/2013
PI Name:	Vlkolinsky, Roman Ph.D.		
Project Title:	Functional decline in mice with Alzheimer's-type neurodegeneration is accelerated by charge-particle radiation		
Division Name:	Human Research		
Program/Discipline:	HUMAN RESEARCH		
Program/Discipline Element/Subdiscipline:	HUMAN RESEARCHRadiation health		
Joint Agency Name:	TechPort:		No
Human Research Program Elements:	(1) SR:Space Radiation		
Human Research Program Risks:	(1) BMed:Risk of Adverse Cognitive or Behavioral Conditions and H	Psychiatric Disorders	
Space Biology Element:	None		
Space Biology Cross-Element Discipline	None		
Space Biology Special Category:	None		
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Zip Code:	92350-1700	Congressional District:	41
Comments:			
Project Type:	Ground		2010 Space Radiobiology NNJ10ZSA001N
Start Date:	02/01/2011		01/31/2015
No. of Post Docs:	1	No. of PhD Degrees:	
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:	NASA JSC
Contact Monitor:	Simonsen, Lisa	Contact Phone:	
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Flight Program:			
Flight Assignment:	NOTE: End date is now 1/31/2015 per NSSC information (Ed., 11/5/	13)	
Key Personnel Changes/Previous PI:	Jerome Badaut, PhD terminated participation in our project as of July	v, 2013. Richard E Hartman, PhD ; Gregory Nelson, PhI	D; Attila Szucs, PhD - subcontractor
COI Name (Institution):	Nelson, Gregory (Loma Linda University) Hartman, Richard Ph.D. (Loma Linda University)		
Grant/Contract No.:	NNX11AE41G		
Performance Goal No.:			
Performance Goal Text:			
Task Description:	Exposure of an astronaut's central nervous system (CNS) to solar particle events (SPE) and galactic cosmic rays (GCR) may accelerate neurodegenerative changes and impact neuronal network activity, leading to cognitive deficits. There are similarities between radiation CNS effects and pathological processes found in the Alzheimer's disease (AD). Common functional and structural findings include profound deficits in neuronal communication (synaptic transmission), cognitive impairments, neuro-inflammatory changes and reduced neurogenesis. These similarities lead us to hypothesize that subjects with a genetic propensity to develop AD-pathology may be excessively vulnerable to ionizing radiation. We previously showed in transgenic (TG) APP23 mice, a murine model of AD, that irradiation with 600 MeV in roop particles accelerated the onset of electrophysiological changes in the hippocampus, a brain structure crucially involved in the formation of short-term memory. In this project we use young adult APP/PSEN129 (APP/PSEN129) duble transgenic (TG) mice accesses which will be develop AD-valhology may be excessively not (irradiations performed at LLU) proton treatment facility, 250 MeV/n silicon and incon-particle radiation to compare and quantify their detrimental effects on hippocampal functions and the onset of AD-like pathology. The APP/PSEN1TG mice typically exhibit early-onset of age-related behavioral abnormalities and deficits in synaptic transmission. We hypothesized that exposure to even low radiation does will accelerate the onset of age-related neurodegenerative processes, while in wild-type (WT) animals such damage may stay undetectable. Comparison of proton, silicon and ino radiation on selected neurophysiological and points in APP/PSEN1TG mice will provide valuable information whether exposure to space radiation may exacerbate neurodegenerative processes. The functional and points (e.g. electrophysiological and behavioral changes) will be directly correlated with the expression of immunohistochemical marke		
Rationale for HRP Directed Research:			
Research Impact/Earth Benefits:	may affect basic neuronal processes, such as synaptic transmission, n formation of memory, the ionizing radiation has been shown to impa- onset of neurodegenerative disorders that affect the hippocampus, su protons and high-LET radiation on neurodegenerative processes in m cranial radiotherapies using charged particle radiation. The time-depe severe delayed demyelination and neurodegeneration. Whether low d murine double transgenic model of AD that we exposed to low- and l	euronal excitability, and formation and consolidation of et synaptic excitability and plasticity. In addition, it cam the as Alzheimer's disease (AD). However, this hypothes ammalian CNS is a critical step, not only for the assess neden tchanges in the CNS in patients undergoing crania losses of charged particle radiation may accelerate the on nigh-LET charged-particle radiation to attempt to answe ation of behavioral, electrophysiological, and histologi	hysiological and new behavioral evidence showing that even low doses of ionizing radiation 'spatial memory. Specifically in the hippocampus, a brain structure intimately involved in the not be excluded that ionizing radiation, even at very low doses of 0.1-1 Gy, may promote the sis has not been fully tested with different low- and high-LET particles. Studying the impact of ment of the space radiation risks for astronauts, but also for further development of modern al irradiations have been well documented, and they range from mild memory deficits to set or affect the severity of AD-related pathology is not known. In the current project we used: r this question. We tested whether radiation affects the time course and severity of cal data will help us to identify functional decrements and the neurodegenerative changes in ated and AD-affected CNS tissue.

Task Progress:	We received an approval for one year no-cost extension of our project ending in January 31, 2015. Dr. Badaut, PhD, has been a co-investigator and lead for the immunohistochemical (IHC) aspect of our project. In July, Dr. Badaut had announced his intention to relocate to France and on his request his participation in our project was terminated as of July, 2013. Dr. Shalini Mchrotra, PhD, a first year postdoctoral fellow, was trained by Dr. J. Badaut. She was working with us since June 2012 and she was mostly responsible for performing the IHC analyses. Dr. Mchrotra had announced terminating her employment with LLU in August, 2013. Thus, the HIC part of our study was paused since Dr. Badaut's laboratory equipament (e.g., fluorescent microscope with Mercator software package) became unavailable. Nonetheless, we intend to complete the HIC part in full in collaboration with Dr. Nelson (co-investigator) and students trained by Dr. Badaut. Re-allocation of funds allowed us to hire part time Mr. Gordon Harding as of September 2013, a senior research associate with significant experience in Western blotting and other protein quantification techniques. Mr. Harding performed initial experiments with presynaptic marker synaptophysin in mice irradiated with protons. Technical progress: Summary Results by Aims. Aim 1 & Aim 3 Activities. In accord with our statement of work (SOW), we completed all irradiations, behavioral testing, and in vitro electrophysiological experiments with protons and HZE (ron 600 MeV/n & silicon 250 MeV/n) particles. Proton inradiated APPTESEN1 transgenic (TG) and wild-type (WT) mice were behaviorally tested and 6 months post-irradiation followed by electrophysiological testing at6 -7 months post-irradiation. Pooling two electrophysiological testing to the off and the DW mice with protons at Lona Linda Lunking testing at0 francing sets. This change to the original SOW was opted for a flor observing considerable variability in electrophysiological time point was applied for both HZE species to allow f	
Lisk Frigress:	Electrophysiological data show that proton radiation at doses from 0.1 to 1 Gy may impact synaptic excitability and short term synaptic plasticity mediated by presynaptic glutamate release, but it likely does not affect long-term potentiation (LTP; reported previously), the widely used cellular correlate of memory formation in the hippocampus. We observed that proton radiation-induced changes in synaptic excitability are qualitatively different in APP/PSEN1 TG and WT mice. In accord with our behavioral findings, the WT mice exhibit different sensitivity to radiation and, for example at 0.5 Gy we observed increased postsynaptic excitability in CA1 neurons, whereas the TG mice exhibited opposite responses at the same radiation dose. The stimulation paradigms using two (paired) stimulation pulses were used to evaluate the effect of proton radiation on presynaptic glutamate release (paired-pulse facilitation; PPF). In TG mice at 6 months post-irradiation with protons we observed reduced PPF indicating increased glutamate release and this change became more pronounced at 9 months post-irradiation. Changes in PPF, vere not detected in WT mice. On the other hand, WT mice exhibited sensitivity to proton radiation because at the dose of 0.5 Gy we observed radiation-induced decrements in frequency of sharp wave-ripple complexes, which are implicated in memory consolidation process in the hippocampus. Interestingly, in TG mice, a radiation exposure to protons or HZE particles had no effect on these spontaneous oscillations of β-amyloid deposits in the brain samples (the cortex and the hippocampus) of APP/PSEN1 TG mice irradiated with protons using thioflavin-S staining (fibrillar form of amyloid) and by HIC using 6E10 monoclonal antibody (total amyloid). Both methods confirmed amyloid depositions in the hippocampus) at 0 Sy of protons we observed significant increase of total amyloid by 9 months not-irradiation in detected by 6E10 antibodies. The HHC on brin samples irradiation with the doracle cortex (but not the hippoca	
	HZE particles was temporarily paused due to departure of Drs. Badaut and Methorta. Nonetheless, the IIIC analyses of HZE irradiated samples is planned for the fourth year of the project (the no-cost extension has been approved) by hardware provided by Dr. Nelson (co-investigator) and performed by other team members trained in Dr. Badaut's lab and by student volunteers. Neuroinflammation and neurodegenerative changes in TG (and WT) brains (cortex only) exposed to radiation have been assessed by determination of five cytokines/chemokines (IL-1 beta, IL-6, TNF alpha, MCP-1, and IL-10). These molecules have been previously reported to be elevated in irradiated brains and/or have been shown to affect synaptic plasticity in the hippocampus, thus their elevation may be associated with functional decrements observed in these animals. The Luminex assays have been completed in samples irradiated with protons, the assays with HZE-irradiated brains will be completed by December, 2013. In a cohort of proton-irradiated mice we observed differences in the expression of chemokines IL-10 between TG and VTT mice a 9 months, such dependent on the radiation exposure. The other chemokines were not affected by either genotype or radiation, indicating that at 9 months radiation effects on the CNS are not associated with elevated levels of pro-inflammatory cytokines. This also indicated that the electrophysiological and behavioral decrements reported above are not due to elevated levels of cytokines within the CNS, as previously suggested by us and other investigators. We are currently performing the analyses of synaptic markers in WT and TG mice irradiated with protons by Western blotting. The initial analyses in APP/PSEN1 TG mice irradiated with protons indicates that such exposure to iron radiation, which awaits confirmation in APP/PSEN1 TG mice planned for the next year. Analyses in cortices irradiated with 0.1 and 1 G y of protons and with HZE particles will be ensure.	
Bibliography Type:	Description: (Last Updated: 04/24/2019)	
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Abstracts for Journals and Proceedings	Rudobeck E, Szücs A, Mehrotra S, Vlkolinsky R. "Ionizing radiation impairs hippocampal functions in APP/PSEN1 transgenic mice." Neuroscience 2013, San Diego, CA, November 9-13, 2013. Neuroscience 2013, San Diego, CA, November 9-13, 2013. Program#/Poster#: 802.06/E19. Abstract available at: http://www.skitractonine.com/Plan/ViewAbstract.aspx?sKey=fdeef2fib.ad7c-41efb.b7f1.d21d68sd8565&cKey=2de25863.0125-de1d-baed-6160ce30d33f&mKey=[8D2A5REC_4825-dCD6-9439.R42RR151D1CE] accessed 12/5/13., Nov-2013	
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Articles in Peer-reviewed Journals	Rudobeck E, Szücs A, Vlkolinsky R. "Effects of Proton Radiation on Evoked and Spontaneous Neuronal Activity in the Hippocampus of APP/PSEN1 Transgenic Mice." Journal of Radiation Research. In press, as of December 2013. To be published January 2014. , Dec-2013	