Fiscal Year:	FY 2010	Task Last Updated:	FY 12/09/2009
PI Name:	Tai, Yu-Chong Ph.D.		
Project Title:	In-flight Blood Analysis Technology for Astronau	ut Health Monitoring	
Disting Name	Hannan Daaaanta		
Division Name:	Human Kesearch		
Program/Discipline:	NSBRI		
Program/Discipline Element/Subdiscipline:	NSBRISmart Medical Systems and Technology	7 Team	
Joint Agency Name:		TechPort:	Yes
Human Research Program Elements:	(1) ExMC :Exploration Medical Capabilities		
Human Research Program Risks:	(1) Medical Conditions : Risk of Adverse Health that occur in Mission, as well as Long Term Heal	Outcomes and Decrements in Po th Outcomes Due to Mission Ex	erformance Due to Medical Conditions
Space Biology Element:	None		
Space Biology Cross-Element Discipline:	None		
Space Biology Special Category:	None		
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Zip Code:	91125	Congressional District:	29
Comments:			
Project Type:	Ground	Solicitation / Funding Source:	2007 NSBRI-RFA-07-01 Human Health in Space
Start Date:	10/01/2007	End Date:	09/30/2011
No. of Post Docs:	1	No. of PhD Degrees:	2
No. of PhD Candidates:	3	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:			
Key Personnel Changes/Previous PI:			
COI Name (Institution):	Ho, Chih-Ming (University of California, Los A Kasdan, Harvey (IRIS International) Adams, Thomas (IRIS International)	Angeles)	
Grant/Contract No.:	NCC 9-58-TD01301		
Performance Goal No.:			
Performance Goal Text:			

Rationale for HRP Directed Research: Research Impact/Earth Benefits: The devices under development can be used for earth-based applications. The proposed device use cartridge and hand held system. The cartridge will be cheap and disposable. The results will be available almost immediately to the patients without going through central lab facilities. The device an be used in emergency room, on ambulance as well as at home. As the senior population continuous to grow, this kind of device will find more and more appealing in point-of-care applications. For the first funding year, we proposed to optimize the blood staining and testing procedure for 5-part WBC differential, optimize hydrodynamic separator for WBC subtype separation, and plasma preparation for plasma biomarker detection. We are on the schedule. 1. Blood staining and testing procedure optimization: We demonstrated 4-part WBC differential count including Lymphocyte, Monocyte, Neutrophil and Eosinophil, with a combination staining of FTIC and PI. We also demonstrated 3-part WBC differential count by an optimized Acridine Orange staining. 2. Fluorophore conjugated antibody staining of whole blood: As an alternative to chemical staining, we demonstrated WBC differential and WBC subtype counting with fluorophore conjugated antibody. By using a laser module for excitation, the sensitivity of the prototype is largely improved to detect signals from fluorophore conjugated antibody. Monocyte count and WBC subtype counts such as CD4+ have been demonstrated on the proposed antibody. Whonocyte count and WBC subtype count such as CD4+ have been demonstrated on the prototype on a zero-G parabolic flight test, which is important to validate using the prototype in the rate of more restored antibody test as been achieved. Experiences from the flight test are being used for improvement of t	Task Description:	The specific aims of the project include (a) 5-part MBC differential, (b) analysis of WBC subtypes (e.g., CD4+ T helper and natural killer cells), and (c) serum/plasma protein biomarker analysis (e.g., for infection, radiation and bone loss monitoring). This project is a continuation of a related project entitled "Handheld Body-Fluid Analysis System for Astronaut Health Monitoring." in which we explored electrical impedance sensing, fluorescence optical sensing, and flow separation of blood cells in microfluidic devices and portable platforms. We successfully demonstrated fluorescent assessing and counting for WBC count and 2-part differential with a portable prototype microf lowcytometer. For the current project, a major effort is proposed to extend the 2-part WBC differential to a 5-part WBC differential, add cell surface marker detection and analysis capability to the platform repertoire, and add plasma protein detection and analysis eqability to the platform repertoire. Our approach to achieve the objectives is to extend the capability of the microf lowcytometer to enable a more comprehensive WBC differential, and allow detection of fluorescent labels attached to ligands used for cell surface marker and plasma protein detection. The second component necessary for extending the platform capability is the offline data analysis software. This software is being developed in Matlab to facilitate both quantitativa assessment of fluorescence detection and cell and analyte recognition and quantitation. In the last funded year, we successfully tested the proposed microf lowcytometer in a zero-G parabolic flight test in collaboration with research scientists from Wyle cooperation. The test demonstrated the facility of doing WBC differential count in zero/micro gravity environment with the proposed prototype. Results similar to on-ground test are obtained. The prototype was shown to be convenient for operation. One flight crew learned to operate the prototype and carried out the test after a brief training.	
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	reinforcing the mechanical structure and increasing the pumping efficiency under low environmental pressure.
Task Progress:	4. Hardware improvement: The excitation source of the prototype has been upgraded to a blue laser from a LED. The laser excitation provides stronger fluorescence intensity for detection. Now we are working on improving the detection component. Analysis of fluorescence spectrum would be explored with a mini-spectrometer detector.
	5. Optimization of hydrodynamic separator for WBC subtype separation: The geometry of the sorting region has been further optimized to improve sorting efficiency and enhance continuous operation. Computational simulations were utilized to improve the design of various components including the hydrodynamic focusing region, the sorting region and the collection bins. Experiments have been performed to successfully separate particles and embryoid bodies into size-dependant groups.
	6. WBC subtype analysis with EIS: Characterization of single-cell dielectric properties using EIS requires immobilization of the target cells onto detection electrodes with accurate position control at the single-cell level. We have refined our previously developed cell patterning technique through optimizing the fabrication process to achieve high selectivity protein patterns enabling for the precise formation of single-cell arrays. We have also improved the detection sensitivity by increasing the effective electrode surface area through a polypyrrole (PPy)-electrode coating and by using a low conductivity cell suspension buffer.
Bibliography Type:	Description: (Last Updated: 08/30/2018)
Articles in Peer-reviewed Journals	Li N, Ho CM. "Patterning functional proteins with high selectivity for biosensor applications." Journal of Laboratory Automation. 2008 Aug;13(4):237-42. <u>http://dx.doi.org/10.1016/j.jala.2008.04.001</u> , Aug-2008
Articles in Peer-reviewed Journals	Li N, Ho CM. "Photolithographic patterning of organosilane monolayer for generating large area two-dimensional B lymphocyte arrays." Lab Chip. 2008 Dec;8(12):2105-12. <u>http://dx.doi.org/10.1039/b810329a</u> ; <u>PMID: 19023473</u> , Dec-2008
Dissertations and Theses	Li N. "Lab-on-a-chip systems for blood cell separation, counting, and characterization." Dissertation, University of California, Los Angeles, June 2009. , Jun-2009
Papers from Meeting Proceedings	 Shi W, Zheng S, Kasdan HL, Fridge A, Tai YC. "Leukocyte count and two-part differential in whole blood based on a portable microflow cytometer." 15th International Conference on Solid-State Sensors, Actuators and Microsystems (Transducers 2009), Denver, CO, June 21-25, 2009. 15th International Conference on Solid-State Sensors, Actuators and Microsystems (Transducers 2009), Denver, CO, June 21-25, 2009. 15th International Conference on Solid-State Sensors, Actuators and Microsystems (Transducers 2009), Denver, CO, June 21-25, 2009.
Papers from Meeting Proceedings	Zheng S, Kasdan HL, Fridge A, Tai YC. "Blood cell analysis using portable flow cytometer with microfluidic chips as cartridge." 12th International Conference on Miniaturized Systems for Chemistry and Life Sciences, San Diego, CA, October 12-16, 2008. Proceedings of the 12th International Conference on Miniaturized Systems for Chemistry and Life Sciences, San Diego, CA, Oct. 12-16, 2008. , Oct-2008