The effect of microgravity on mammalian system is an important and interesting topic for scientific investigation, since NASA’s objective is to send manned flights to planets like Mars and eventual human colonization. The Astronauts will be exposed to microgravity environment for a long duration of time during these flights. Our objective of research is to conduct in vitro studies for the effect of microgravity on mammalian immune system. We did our preliminary investigations by exposing mammalian lymphocytes to a microgravity simulator cell bioreactor designed by NASA and manufactured at Synthecon Inc (USA). Our initial results showed no significant change in cytokine expression in these cells for a time period of forty eight hours exposure. Our future experiments will involve exposure for a longer period of time.

INTRODUCTION

During space flight the function of the immune system changes significantly. Several papers reported that postflight the number and the proportion of circulating lymphocytes in astronauts are modified (Uchakin et al 2002), the in vitro mitogen induced T cell activation is depressed (Cogoli et al. 2002, Konstantinova et al. 1993) and there are detectable differences in cytokine production. Lymphocytes as well (Chapes et al. 1992). One of the possible modifying forces is the microgravity condition itself. Our aim was to analyse mechanisms responsible for changing lymphocyte functions in low gravity environment. For terrestrial simulation of microgravity we used a Rotary Cell Culture System (RCCS) developed by NASA.

In this experiment we exposed mouse B Lymphocyte cells to microgravity conditions and then analyzed the cells for cytokine expression. We exposed the cells to different time periods, however, our initial results failed to show any significant changes in cytokine expression under microgravity conditions.

MATERIALS AND METHOD

Mouse B Lymphocyte cells were purchased from ATCC, VA, USA and cultured in L-15 medium at 37°C in a cell culture incubator. Cells were exposed to microgravity conditions in a Rotary Cell Culture System (RCCS) developed by NASA for 24 hours, 48 hours and 72 hours with proper control. After exposure cells were collected, lysed by antigen lysis buffer and cytokine expression (TGF-beta1 and IL-6) was determined by standard ELISA technique according to manufacturers instructions.

RESULTS

No significant changes in any cytokine expression tested was found during that particular exposure period under microgravity simulations.
DISCUSSION

Several attempts have been made to investigate the effects of microgravity on the growth and function of animal cells. Cellular activation of immune T lymphocytes is greatly affected by microgravity. On the other hand, little is known about the effects of microgravity on B lymphocytes. Thus, we attempted to study the effect of microgravity simulation on mouse B lymphocytes.

Our current experiment did not show any significant change in cytokine expression in microgravity exposed immune cells. We look forward to do experiments with longer exposure time, since astronauts are exposed to microgravity conditions for months and years. Several researchers have reported alteration of the immune system due to microgravity conditions (Uchakin et. al 2002). However, we did not notice any significant change in cytokine expression in any of our experiments done for shorter intervals of time i.e., 24 hours, 48 hours and 72 hours.

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BIBLIOGRAPHY

5) Uchakin PN, Tobin BW, Morukov BV, Larina IV, Cubbage ML. Type 1 vs. type 2 cytokine secretion in vitro and its regulation by hydrocortisone in humans subjected to 120-day anti-orthostatic bed-rest regime. J Gravit Physiol. 2002 Dec;9(2):71-82.