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High Glucose Enhances the Proliferation Effects of Stress Hormones in Mesenchymal Stem Cell Cultures

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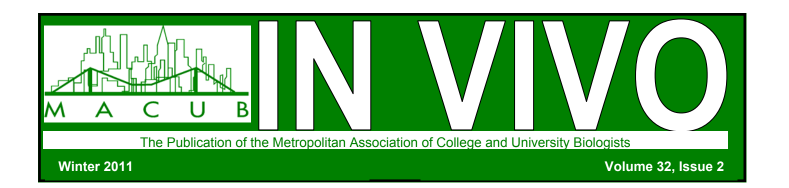
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In This Issue:	
MACUB 2010-2011 Executive Board	inside cover
Instruction for Authors	inside cover
MACUB 2010-2011 Executive Board Election Results	23
MACUB 2010 Conference Poster Presentation Award Winners	24
MACUB 2010 Conference Poster Abstracts	26
MACUB 2010 Conference Member Presentations	53
Highlights of the 43rd Annual MACUB Conference	54
Benjamin Cummings/MACUB Student Research Grants	55
Affiliate Members, 44 th MACUB Conference Announcement	inside back cover

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MACUB 2010 Conference Poster Abstracts

High Glucose Enhances the Proliferation Effects of Stress Hormones in Mesenchymal Stem Cell Cultures. Nancy Abrego^{1,2}, Jodi F. Evans^{1,2} and Louis Ragolia², ¹Molloy College, Rockville Centre, NY and ²Winthrop University Hospital, Mineola, NY.

Mesenchymal stem cells (MSC) are the progenitor cells to connective tissue cells, epithelial cells, and smooth muscle These cells are currently being investigated as a cellular source for tissue regeneration and repair. systemic and local tissue environment may have significant influence on the success of such efforts. We hypothesized that elevated glucose and exposure to stress hormones would influence the proliferation of MSC. MSC from the bone marrow of Wistar-Kyoto (WKY) rats were grown under both low glucose (5 mM) and high glucose (20 nM) conditions in the presence and absence of the synthetic glucocorticoid, dexamethasone, and adrenocorticotropin hormone (ACTH). Relative cell density was used to determine the rate of proliferation and was measured using methylene blue staining. Changes in cell density from initial plating were recorded at various stages of culture. Under low-glucose conditions, stress hormones reduced MSC proliferation and these changes were enhanced when cells were exposed to high glucose conditions. These data indicate that stress and elevated glucose can have a significant effect on the ability of MSC to proliferate and may prove to reduce their efficacy during tissue repair.

Are the Neurotoxic Effects of Manganese Due to Blockage of Post Synaptic Dopamine Receptors. Trevon Adams¹, Danilo Beaubrun², Michael Nelson¹, Margaret A. Carroll¹ and Edward J. Catapane¹, ¹Medgar Evers College and ²Kingsborough Community College.

Manganese, a neurotoxin causing Manganism, a Parkinsons-like disease, disrupts dopaminergic neurotransmission. Lack of effective treatment for Manganism is major obstacle in its management. Recently, paminosalicylic acid (PAS) was reported an effective treatment. Lateral cilia of gill of Crassostrea virginica are controlled by serotonergic-dopaminergic innervations from their ganglia. We showed manganese blocks cilio-inhibitory effects of dopamine and this is prevented by PAS. We sought to determine if manganese exerts its effects by blocking dopamine post-synaptic receptor binding and if PAS prevents manganese from doing this. We observed membrane potentials of lateral ciliated cells with fluorescent dye while measuring cilia beating. Applying dopamine or 20 Hz electrical stimulation after exciting cilia repolarized the cell membrane and decreased beating. Manganese prevented this. PAS prevented the actions of manganese. Adding ATP to gill increased cilia beating without changing membrane potential. Applying MDL-12,330A, an adenylcyclase inhibitor, after manganese decreased cilia beating without affecting membrane potentials. The study shows the correlation between membrane potential of lateral ciliated cells and cilia beating rates. It helps elucidate the neurotoxic mechanism of action of manganese, showing the site of action is after the post-synaptic dopamine receptors. This information is helpful to understand causes and potential therapeutic treatments of Manganism.

The Synergistic Effect of Green Tea Polyphenols with Antiseptics and Antibiotics against the Growth of Potentially Pathogenic Bacteria. Sylvia Chinons Akuwudike, Bobby Haghjoo and Lee H. Lee, Montclair State University, Montclair, New Jersey, USA

Green tea leaves contain many polyphenolic compounds as (-)-epicatechin, (-)-epicatechin-3-gallate, epigallocatechin, and (-)-epigallocatechin-3-gallate(EGCG). Green tea polyphenols (GTPs) have been implicated to have distinct properties that combat the harmful effects of cell proliferation. These compounds contain certain anti-viral and anti-microbial mechanisms that inhibit growth and perhaps reverse the process in which replication occurs. In this study, 2% GTP was used alone or with antiseptics and antibiotics to study its effect on different gram + and gram - bacteria. For the antiseptic study, disk diffusion test was carried out and for the antibiotics; Kirby-Bauer method was used. The zones of inhibition were measured in millimeter and bacterial resistant, intermediate, or susceptible was determined. The results suggested that GTP works best on the gram positive bacteria and had very little effect on the gram negative bacteria. The most powerful GTP effect can have zone of inhibitions reaching to more than 8mm.

Effect of Gap Junction Inhibitors on Breast Cancer Cell Migration. Vanessa Almonte¹ Maria L. Cotrina² and Regina Sullivan¹, ¹Queensborough Community College, Bayside, NY and ²Columbia University, NY.

Connexins, a family of transmembrane proteins, form intercellular gap junctions in vertebrate cells. Gap junctions allow for cell-cell communication and the passage of small molecules between cells. Defective gap junctions have been identified in cancer cells however their role in cancer progression and the maintenance of a metastatic phenotype remains elusive. This study focused on the role of gap junction hemichannels in the migration of MDA 231, a highly metastatic breast cancer cell line. Experiments were done to assess the levels of functional gap functions in MD231 cells and compared to mouse astrocytes, a cell line that shows abundant gap junctions and to a human embryonic kidney cells which show low levels of gap junctions. The cells were grown to a confluent monolayer and treated with carboxydichloroflurescein. The monolayers were injured with a razor and the cells were imaged using an inverse phase fluorescent microscope fitted with a green filter. The wound healing assay was used to determine the effect of two gap junction inhibitors, carbenoxolone and meclofenamic acid, on the migration of 231 cells. The results of the dye transfer assay revealed that 231 cells have a low level of gap junctions. Interestingly, however ,carbenoxolone significantly inhibited cell migration while meclofenamic acid caused cellular morphological changes. Further studies will investigate the specificity of these effects.