## STRONG CHILDREN'S RESEARCH CENTER

# Summer 2016 Research Scholar

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#### **ABSTRACT**

Title:

A Comparison of Lung Derived Mesenchymal Stromal Cells and Fibroblasts Using qRT-PCR

## Background:

In recent years, stem cells have shown great promise for the repair and regeneration of tissues and organs through stem cell therapy. Access to developing human lung tissue is a major barrier to advancing medical treatments for neonatal and pediatric lung disease. Many recent studies suggest mesenchymal stromal cells could be the start to an answer for treating these chronic developmental barriers faced by a premature infant. The LungMAP was established by NHLBI to provide an atlas of normal human lung development to the scientific community for improved understanding of lung disease. The data presented is a continuation of a study that showed the diversity of samples isolated from 6 donor pilots. We have continued to characterize these 6 donor pilots but now in comparison to 7 fibroblast donors. MSCs are fibroblast-like shaped cells residing in mesodermal tissues of bone marrow, adipose tissue, lung, placenta, and the umbilical cord stroma<sup>1</sup>. MSCs have identifying features including their ability to adhere to plastic surfaces in cell culture laboratory practices, their expression of CD105, CD73, and CD90, and their

#### **Conclusion:**

MSCs are quintipotential cells at the top of their differentiation hierarchy and have the ability to become osteocytes, adipocytes, chondrocytes. Fibroblasts are terminally differentiated cells at the bottom of the MSC differentiation hierarchy. RNA from MSCs and fibroblasts from 6 and 7 respective donors was collected at P4/5 and P10/11 for analysis by qRT-PCR. Primers were selected from published literature to test the expression of various markers within the cells to characterize at the early and late passages. After analysis, it was found that the MSCs and fibroblasts from P4/5 and P10/11 were not as different as predicted. Future directions for the project include continuing to expand the cells to P15 and test the same primers with RT-PCR to see if there is a significant difference after more time. DO42 adult lung MSCs and fibroblasts as well as 5 cord donor cells are currently being grown; once at P15, another project could be to test these cells alongside the MSCs at P5, P10, and P15. Lastly, it would be beneficial to review new published literature to look for genes that may show larger differences, including an exploration of the difference in contact inhibition between the MSCs and fibroblasts.