**Docket #:** S17-417

# Systemic targeting of inflammatory sites and enhanced immunomodulatory function by introducing the chimeric antigen receptor (CAR) into mesenchymal stem cells for inflammatory and autoimmune diseases

Stanford inventors have developed a mesenchymal stem cell-based gene therapy that can target the inflammatory environment and secrete immunomodulatory cytokines. The model has been demonstrated in bone marrow mesenchymal stem cells in vitro. The inflammation-targeting molecules are expressed in approximately 90% of mesenchymal stem cells at 24 hours post-electroporation. The immunomodulatory cytokines can be detectable for more than 48 hours and convert the pro-inflammatory response into an anti-inflammatory tissue repair/regeneration response.

# **Applications**

 In clinical therapy, to enhance the immunomodulatory and tissue repair capabilities of mesenchymal stem cell transplantation in inflammatoryassociated diseases including diabetes, bone healing, osteoarthritis, myocardial infarction, pulmonary hypertension, spinal cord injury, inflammatory diseases of other organs (hepatitis, nephritis etc.).

# **Advantages**

- Enhanced immunomodulation and targeting specificity of mesenchymal stem cell-based therapy.
- Using transient gene expression instead of genome integration to avoid potential side effects.
- Modulation of the crosstalk between mesenchymal stem cells and immune cells to enhance tissue regeneration.
- Expandable to other cell-based immunomodulation therapies.

### **Patents**

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Published Application: <u>WO2019195142</u>

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