

# Gene and cell therapy on the acquisition and relapse-like binge drinking in a model of alcoholism: translational options

Yedy Israel,

Quintanilla, María Elena

Ezquer, Fernando

Morales, Paola

Rivera-Meza, Mario

Karahanian, Eduardo

Ezquer, Marcelo

Herrera-Marschitz, Mario

Studies reviewed show that lentiviral gene therapy directed either at inhibiting the synthesis of brain acetaldehyde generated from ethanol or at degrading brain acetaldehyde fully prevent ethanol intake by rats bred for their high alcohol preference. However, after animals have chronically consumed alcohol, the above gene therapy did not inhibit alcohol intake, indicating that in the chronic ethanol intake condition brain acetaldehyde is no longer the compound that generates the continued alcohol reinforcement. Oxidative stress and neuroinflammation generated by chronic ethanol intake are strongly associated with the perpetuation of alcohol consumption and alcohol relapse ?binge drinking?. Mesenchymal stem cells, referred to as guardians of inflammation, release anti-inflammatory cytokines and antioxidant products. The intravenous delivery of human mesenchymal stem cells or the intranasal administration of mesenchymal stem cell-generated exosomes rever