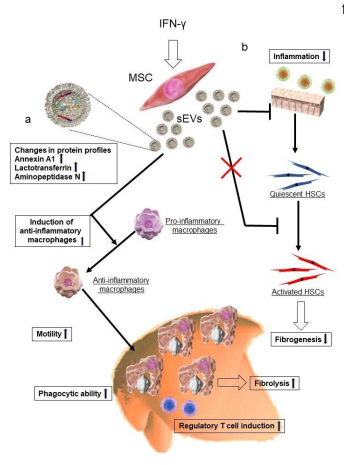


A glimmer of hope: New weapon in the fight against liver diseases

3 May 2021



Increased levels of annexin A1, lactotransferrin, and aminopeptidase N. Increase the counts of antiinflammatory macrophages with high motility and phagocytic ability, which increase repair of damaged tissue. Induces regulatory T cells and fibrolysis. b) The ?sEVs also have the ability to reduce inflammation as well as downregulate fibrogenesis. Credit: Niigata University

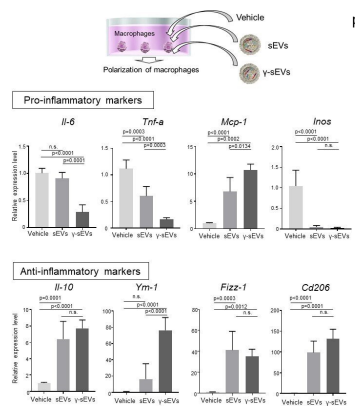
Researchers from Niigata University, the University of Tokyo, Osaka University and Tokyo Medical University, Japan, have developed a new approach that could revolutionize the treatment, prevention, and possibly reversal of the damage caused by liver diseases. This novel strategy exploits small extracellular vesicles (sEVs) derived

from interferon-? (IFN-?) pre-conditioned MSCs (?sEVs).

Cirrhosis and other chronic <u>liver diseases</u> remain a global health concern, with close to 2 million deaths reported annually; these account for approximately 3.5% of annual worldwide deaths. All these statistics are largely driven by the fact that 75 million of the 2 billion people who consume alcohol worldwide are diagnosed with alcohol-use disorders and are at risk of developing alcohol- induced liver disease. In addition, the high prevalence of viral hepatitis-induced liver damage continues to be on the rise.

These sobering numbers inspired a team of scientists led by Prof. Shuji Terai of the Division of Gastroenterology and Hepatology, Graduate School of Medical and Dental Sciences, Niigata University, to build upon previous knowledge regarding loss of the ability to control fibrosis, the development of fibrous connective <u>tissue</u> as a reparative response to injury or damage, in livers under advanced cirrhosis. In an interview Prof. Terai said, "Our results showed that modified extracellular vesicles can become a new therapeutic strategy for liver cirrhosis."





The AD-MSC-?-sEVs effectively downregulate proinflammatory (IL-6, Tnf-a, Mcp-1, and Inos) and induce anti-inflammatory (iL-10, Ym-1, Fizz-1, Cd206) macrophage responses and in vitro. Credit: Niigata University

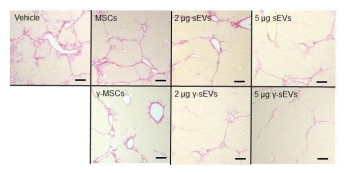
Since clinically advanced cirrhosis is an end-stage disease that can effectively be treated only by liver transplantation at present, there is a race in the field with many scientists developing targeted therapies for modulating fibrosis and aiding tissue regeneration.

One of the most popular approaches is cell therapy, where mesenchymal stromal cells (MSCs) and macrophages have shown potential for inducing liver fibrosis regression. The popularity of this approach is centered on its cost-effectiveness; because MSCs are not only obtainable from the bone marrow, but also from medical waste that include umbilical cord tissue, adipose tissue, and dental pulp. Apart from the ease of availability, MSCs can also be grown relatively easily in the lab. sEVs improved liver fibrosis more than the AD-MSC-Furthermore, rather than acting directly by replacing the damaged tissues, MSCs have

previously been shown to be medical signaling cells that indirectly produce cytokines, chemokines, growth factors and exosomes that are crucial for repairing and regenerating damaged tissue. Over the years, considerable progress has been made toward capacity building for research and clinical trials for novel treatment strategies against liver diseases. This is exemplified by previous demonstrations that MSCs have anti-inflammatory, anti-fibrotic and anti-oxidative effects through these humoral factors.

Tissue rejection is a barrier to cell/tissue transplantation interventions; MSCs are thus attractive for possessing low immunogenicity, and this can facilitate their use for both autologous (same individual) and allogeneic (different individuals of the same species) transplantation, as evidenced by applications in nearly 1000 clinical trials involving other fields, including the treatment of liver diseases.

In a series of experimental mice studies, this team of researchers, from Niigata University, the University of Tokyo, Osaka University and Tokyo Medical University, Japan may have discovered that IFN-? pre-conditioned human AD-MSC-derived sEVs (AD-MSC-?-sEVs) can induce antiinflammatory macrophage counts, which are the key players in tissue repair, including the regression of fibrosis and promotion of tissue regeneration in vitro.



There is dose dependent improvement of liver fibrosis improvement a) PBS (vehicle) treated-most damaged b) AD-MSC, e) AD-MSC-?, f-g) 2 ?g and 5 ?g of AD-MSC-?sEVs (c-d). Credit: Niigata University



MSCs (AD-MSC-sEVs) and AD-MSC-?-sEVs can promote macrophage motility and phagocytic activity. In addition, they also show that AD-MSC-?sEVs contain anti-inflammatory macrophage inducible proteins and can effectively control inflammation and fibrosis in a mouse model of cirrhosis. Following single-cell RNA-seq application, (2021). DOI: 10.1038/s41536-021-00132-4 they confirmed AD-MSC-?-sEVs therapy can induce multidimensional transcriptional changes. Taken together, these results suggest that AD-MSCderived sEVs can affect the shape and function of macrophages and effectively recruit them into damaged areas, thereby promoting damaged liver tissue repair.

Dr. Atsunori Tsuchiya of the Division of Gastroenterology and Hepatology, Graduate School of Medical and Dental Sciences, Niigata University, who was part of the research team said, "Both mesenchymal stromal cells and macrophages are reported to have therapeutic effects for liver cirrhosis, however relationship of both cells and mechanisms of action was not clear. We challenged this problem. We found the important fact that extracellular vesicles from interferon-? can induce the tissue repair macrophages, which can regress fibrosis and promote liver regeneration effectively."

Dr. Suguru Takeuchi of the Division of Gastroenterology and Hepatology, Graduate School of Medical and Dental Sciences, Niigata University, who was also part of the research team, said, "In our previous study, we reported that intravenous administration of mesenchymal stromal cells migrated to the lung, can work as 'conducting cells' and affect to macrophages 'working cells' in the liver. In this study we first elucidated that extracellular vesicles from mesenchymal stromal cells are key molecules to affect the macrophages."

This proof-of-concept pilot study holds potential as a strategy for treating liver diseases using IFN-? preconditioned sEVs. However, further development and determination of the mechanisms underlying Treg cell count induction by IFN-? pre-conditioned MSCs and sEVs still form part of their future research plans before these findings can be

They report that both human adipose tissue-derived translated to humans in phased and controlled clinical trials.

> More information: Suguru Takeuchi et al, Small extracellular vesicles derived from interferon-? preconditioned mesenchymal stromal cells effectively treat liver fibrosis, npj Regenerative Medicine

Provided by Niigata University



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