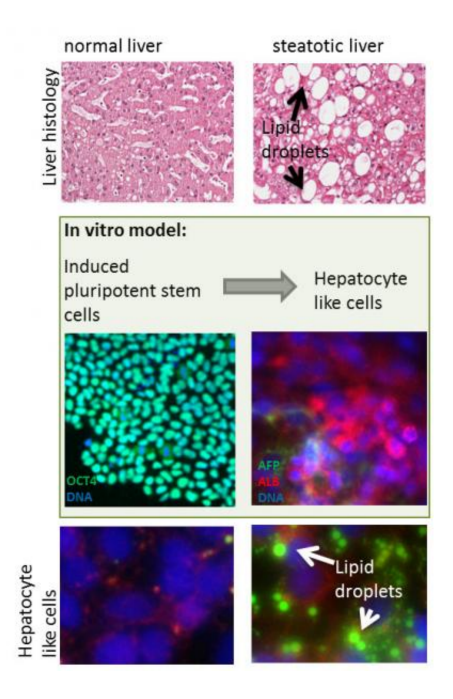


Modeling NAFLD with human pluripotent stem cell derived immature hepatocyte like cells

June 29 2016





Histological section of a liver biopsy of a healthy individual (upper left) and an NAFLD patient (upper right) (1). Lipid droplets are visible as vacuoles. Induced pluripotent stem cells, which express the pluripotency marker OCT4 (middle left, green), are in vitro differentiated into hepatocyte-like cells, characterized by the expression of ALBUMIN (ALB, middle right, red) and alpha-fetoprotein (AFP, middle green right). Unstimulated in vitro derived hepatocyte-like cells do not incorporate lipid droplets (lower left) while they develop massive lipid droplets after stimulation with oleic acid (lower right, green). Credit: Graffmann



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Researchers from the Institute for Stem Cell Research and Regenerative Medicine at the University Clinic of Düsseldorf have established an in vitro model system for investigating nonalcoholic fatty liver disease (NAFLD). The study led by Prof. James Adjaye has now been published in the journal *Stem Cells and Development*.

Nonalcoholic fatty liver <u>disease</u> (NAFLD), also called steatosis, is a dramatically under-estimated liver disease, with increasing incidences throughout the world. It is frequently associated with obesity and type-2 diabetes. Approximately one third of the general population in Western countries are affected, often without even having symptoms. It is a result of a high caloric diet in combination with a lack of exercise, where the liver starts incorporating fat as lipid droplets. Initially, this is a benign state, which can, however, develop into NASH /steatohepatitis, an inflammatory disease of the liver. Then many patients develop fibrosis, cirrhosis or even liver cancer. However, in many cases patients die of heart failure before they develop severe liver damage.

A major obstacle for NAFLD research was, up to now, that biopsies of patients and healthy individuals were required. The Düsseldorf researchers elegantly solved this problem by reprogramming skin cells into so called induced pluripotent stem cells which they differentiated into hepatocyte like cells. "Although our hepatocyte-like cells are not fully mature, they are already an excellent model system for the analysis of such a complex disease", explains Dr. Nina Graffmann, first author of the study. The researchers recapitulated important steps of the disease in vitro. For example they demonstrated up-regulation of PLIN2, a protein that covers lipid droplets. Mice without PLIN2 do not become obese, even when overfed with a high fat diet. Also the key role of PPAR α , a



transcription factor involved in controlling glucose and lipid metabolism, was reproduced in the in vitro system. "In our system, we can efficiently induce lipid storage in hepatocyte-like cells and manipulate associated proteins or microRNAs by adding various factors into the culture. Thus, our in vitro model offers the opportunity to analyse drugs which might reduce the stored fat in hepatocytes" says Dr. Graffmann.

The team now expands the model using induced pluripotent stem <u>cells</u> derived from NAFLD patients, hoping to discover differences which might explain the course of the disease. "Using as reference the data and biomarkers obtained from our initial analyses on patient <u>liver</u> biopsies and matching serum samples, we hope to better understand the etiology of NAFLD and the development of NASH at the level of the individual, with the ultimate aim of developing targeted therapy options," states Professor James Adjaye, senior author of the current study.

More information: Graffmann N, Ring S, Kawala MA, Wruck W, Ncube A, Trompeter HI, et al. Modelling NAFLD with human pluripotent stem cell derived immature hepatocyte like cells reveals activation of PLIN2 and confirms regulatory functions of PPARalpha. *Stem Cells and Development*. 2016.

Provided by Heinrich-Heine University Duesseldorf

Citation: Modeling NAFLD with human pluripotent stem cell derived immature hepatocyte like cells (2016, June 29) retrieved 12 May 2024 from https://medicalxpress.com/news/2016-06-nafld-human-pluripotent-stem-cell.html

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