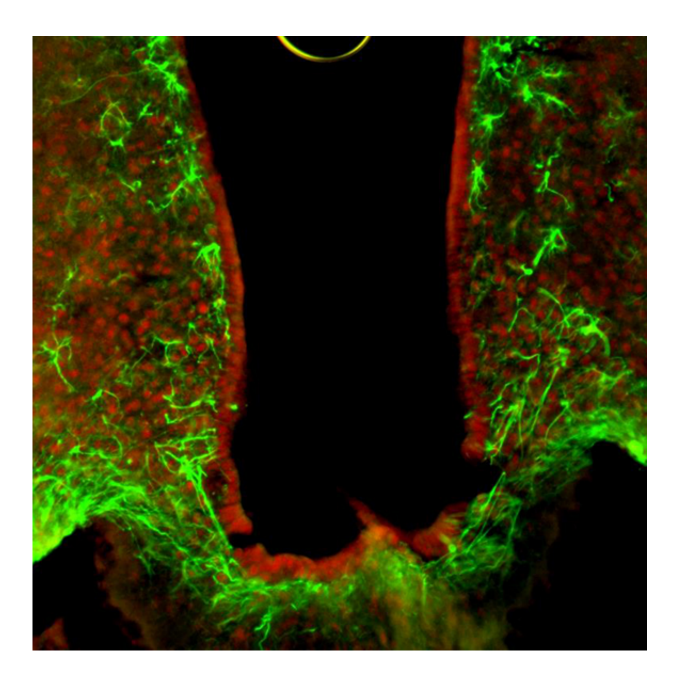


Researchers identify genetic switch regulating satiety and body weight

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HDAC5 (red) is a key factor in neurons for the control of food intake, astrocytes are stained in green. Credit: Helmholtz Zentrum München

A team of researchers at Helmholtz Zentrum München, Technische Universität München and the German Center for Diabetes Research (DZD) has identified a new mechanism that regulates the effect of the satiety hormone leptin. The study published in the journal *Nature Communications* identified the enzyme HDAC5 as key factor in our control of body weight and food intake and potential target against the Yoyo dieting effect.

Why do we get <u>fat</u> and why is it so difficult for so many people to keep off excess weight? Researchers in the Reseach Unit Neurobiology of Diabetes led by Dr. Paul Pfluger and at the Institute for Diabetes and Obesity led by Prof. Dr. Matthias Tschöp have now identified a new component in the complex fine-tuning of <u>body weight</u> and food intake. They found that the enzyme histone deacetylase 5 (HDAC5) has a significant influence on the effect of the hormone leptin. This hormone plays a crucial role in triggering satiety and thus on how the body adapts to a changing food environment.

Pfluger describes the results as follows: "HDAC5 is an important link in the communication between our fat tissue and hunger centers in the brain." Initial experiments have shown that the production and activity of HDAC5 in our control center for energy balance, the hypothalamus, is increased by a high-fat diet, by enlarged fat deposits and by the satiety hormone leptin. Dhiraj Kabra, first author of the study, added: "HDAC5 is a molecular switch in the brain that helps the body to recognize how full the fat depots are. Without HDAC5, we cannot adapt our food intake or curb our fat deposition."



Disruption of the signaling pathway leads to obesity

According to the scientists, mice unable to produce HDAC5 respond significantly worse to leptin – a condition referred to as leptin resistance. They show a continuously increased food intake and – in comparison to mice with intact HDAC5 – show increased obesity, i.e. they get fat. Through targeted activation of HDAC5 the team was able to reverse this effect, and thereby enabling obese animals to loose fat mass and body weight.

"The restoration of leptin sensitivity is an important step on the path towards sustainable weight loss and towards combating potential sequelae of obesity such as type 2 diabetes," said study leader Pfluger. "In addition to the essential changes in diet and exercise behavior, in the future the individual components of the leptin effect could be potential drug targets to support the weight loss process." One of these essential components – HDAC5 – has already been identified. "However," Pfluger concluded, "it remains to be seen in the coming years whether this enzyme will be a suitable target for combating obesity in humans."

Leptin is secreted by <u>fat tissue</u> in direct correlation to body fat mass, thereby signaling to neurons in the brain that the fat stores are full. This leads to a negative energy balance whereupon the body reduces food intake, which ultimately leads to a reduction of body weight and fat storage. As a consequence, in the reduced fat stores less leptin is released, food intake is increased, and the fat stores and body weight are increased.

Specifically, the researchers showed that HDAC5 deacetylates the transcription factor STAT3, whereupon STAT3 can migrate to the <u>leptin</u> receptor. If STAT3 is phosphorylated (activated) there, it migrates back to the cell nucleus and binds to the DNA to activate gene and behavior programs that ultimately cause the reduction of <u>food intake</u>. If HDAC5



is absent (such as here after genetic deletion), STAT3 accumulates as hyper-acetylated and thus inactive protein in the cell nucleus. The leptin signaling pathway is thus switched off, subsequently suppressing the feeling of satiety. This study describes for the first time a direct interaction of the two molecules.

<u>Hungry or Full? – New Switch of Feeling of Satiety Found</u> from <u>Helmholtz Zentrum München</u> on <u>Vimeo</u>.

More information: Dhiraj G. Kabra et al. Hypothalamic leptin action is mediated by histone deacetylase 5, *Nature Communications* (2016). DOI: 10.1038/NCOMMS10782

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