How food intake modifies the gut

Researchers at UNIGE identified that the amount of food regulate the gut size and its capacity to absorb calories, thus shedding light on a fundamental mechanism at the very origin of obesity.

Throughout the world, hundreds of millions of adults and children are clinically obese; a condition closely associated with leading causes of death such as heart diseases or stroke. Obesity mainly results from an imbalance between energy expenditure and caloric uptake, which takes place in the gut where food, previously broken down, is absorbed and passes into the bloodstream for distribution throughout the body. To absorb enough calories, the intestinal wall is layered with millions of convolutions called villi and microvilli that together could cover the surface of a football field.

A striking and rapid effect

“A few years ago, we discovered that the gut could get longer or shorter according to environmental triggers and physiological needs”, recalls Mirko Trajkovski, professor in the Department of Cellular Physiology and Metabolism and the Diabetes Centre at UNIGE Faculty of Medicine, and corresponding author of this study. “We therefore wanted to understand what propels this remarkable intestinal plasticity.” Using different mouse models combined with human intestinal biopsies — 3D artificial structures — the research team observed that the amount of food consumed was the main regulator of gut length. “We saw a relatively fast and physiologically striking response to elevating the amount of ingested food: gut length increased for over 30%, coupled
with a major growth of the villi and microvilli, contributing to an enhanced caloric uptake capacity of the gut.” adds Mirko Trajkovski. Importantly, these changes were reversible: when the amount of food was reduced, the gut length and morphology were reverted close to normal.

**A plasticity under the control of the PPARα protein**

Gut expansion requires a lot of energy. The UNIGE scientists found that increasing the intestinal absorptive surface mobilises different metabolic pathways in the gut, i.e. steps by which cells convert food into energy. While they found several potential pathways that may contribute to gut expansion, one, the PPARα pathway, was found to be indispensable. Indeed, PPARα is a protein that appears of critical importance for increasing the length of villi, as well as for increasing caloric uptake capacity from the food by raising the level of another protein, PLIN2, which, in turn, promotes formation of lipid droplets in the intestinal cells, thereby favouring fat absorption. By inactivating PPARα in the gut of mice, the researchers were able to confirm this mechanism. “Intestinal PPARα deletion, or its pharmacological inhibition, showed marked effects in reducing the absorptive function of the gut. The gut-specific PPARα inhibition was sufficient to revert the fat accumulation and obesity, as well as the glucose intolerance caused by high-caloric feeding”, explains Mirko Trajkovski.

The impressive gut plasticity that permits gut and villi to be shrunk or enlarged is an asset to be further explored as a reversible alternative to gastric bypass surgery, or other interventions that aim at reducing body weight gain and obesity-related complications. “PPARα is a key protein in many metabolic functions, and it is expressed in various tissues throughout the body. It is therefore necessary to develop ways to inhibit it selectively in the gut without reaching the other organs before these discoveries can be applied to patients”, conclude the authors.

---

Sections of mouse intestine. Up, a normal gut circumference (in black) and villi (pink convolutions). Bottom, expanded gut after overeating-induced obesity with a bigger circumference and longer villi.

High resolution pictures