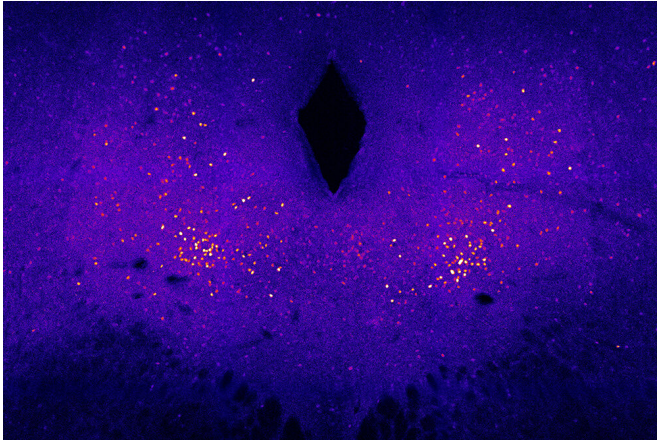


New research raises prospect of better anti-obesity drugs

27 June 2019



Neurons in the brainstem become activate in response to heat. By inhibiting these cells in mice, researchers were able to make the animals eat less and burn more calories. Credit: Laboratory of Molecular Genetics at The Rockefeller University

Effective weight-loss strategies call for eating less food, burning more calories—or ideally, both. But for the more than 90 million Americans who suffer from obesity, a disease that contributes to conditions ranging from cancer to heart disease, behavioral change is hard to accomplish or not effective enough—which is why scientists have long sought drugs that would help people shed pounds. Yet effective, long-lasting treatments have thus far eluded them.

In a new report published in *Cell*, researchers in the laboratory of Jeffrey M. Friedman propose a new avenue in the search for anti-obesity drugs. Collaborating with a Princeton University team, they have found that a group of brain cells previously shown to regulate hunger also controls energy expenditure. And since our body weight depends both on the calories we consume and the energy we burn, these findings could lead to a new type of weight-loss medication that acts on both

sides of the energy equation.

Energy in, energy out

Thus far, most obesity research has focused on the biological mechanisms that govern how much we eat. Yet manipulating the [neural circuits](#) that control our calorie intake hasn't led to broadly successful anti-obesity drugs. So, Marc Schneeberger Pané, a postdoctoral Kavli NSI fellow in Friedman's lab, launched a project to investigate the processes by which we burn energy instead.

Mammals like mice and humans expend energy in many ways, and producing heat is among most important. When the ambient temperature drops, we burn more fuel to maintain a steady body temperature; when it rises, we burn less. We even possess a special form of fat tissue, known as brown fat, that is burned to produce heat directly.

Scientists knew that some populations of temperature-sensitive [neurons](#) in the hypothalamus region of the brain play a role in regulating heat production and, therefore, energy expenditure. But they did not know exactly how those neurons exerted their influence, or if other cells outside the hypothalamus might play a similar role.

Double duty

Schneeberger Pané and his colleagues, including graduate student Luca Parolari, began by mapping the brain regions activated by a rise in ambient temperature. They used an advanced 3-D-imaging technique called iDISCO, [developed at Rockefeller](#), to scan the brains of mice exposed to hot temperatures, looking for signs of neuronal activity.

As expected, the team saw activity in the hypothalamus. But they also saw activity among a particular group of neurons in a part of the brainstem known as the dorsal raphe nucleus—and, to the researchers' surprise, these happened to be

the very same neurons that, just two years earlier, the lab had found to be crucial for controlling hunger.

"Our new findings demonstrate that these neurons regulate energy balance by modulating both food intake and energy expenditure through partially overlapping circuit mechanisms," says Alexander R. Nectow, who led the Princeton University team.

The possibility that these cells, which the lab had previously dubbed "hunger neurons," might regulate both hunger and energy expenditure raised the prospect that they could serve as powerful levers for managing [body weight](#).

"We were very excited," recalls Schneeberger Pané, who adds that he and his colleagues view these multitasking neurons as "a new horizon in obesity research."

The researchers used sophisticated biochemical techniques to alternately turn these temperature-sensitive brainstem neurons on and off. They discovered that activating the neurons reduced the temperature of the animals' brown fat, which is burned to generate heat, core body temperature as well. Suppressing the neurons, meanwhile, amped up heat production—and, as the scientists had shown previously, it also made the animals less hungry.

Eat less, burn more

But generating heat by burning brown fat isn't the only way to expend energy. Physical activity burns calories too, as do all the basic tasks that keep the body alive: breathing, digesting food, and so on. So the researchers put the mice in special cages tricked out with sensors to track their movements and gauge how much carbon dioxide they produce and how much food, water, and oxygen they consume. The goal was to see if the temperature-sensitive hunger neurons dorsal raphe nucleus could control energy expenditure not only by regulating temperature, but by other means as well.

Again, the results were clear: just as activating the neurons caused heat production to plummet, so too did it cause movement, metabolic activity, and

overall energy expenditure to nosedive. Suppressing the neurons, on the other hand, caused all of them to rise.

The team has already begun searching for unique receptors in these dual-purpose neurons that govern both hunger and [energy](#) expenditure. The idea is to identify targets that can be used to create novel anti-obesity drugs capable of delivering a one-two punch.

"When you inhibit these neurons, they suppress food intake and increase [energy expenditure](#) at the same time," says Schneeberger Pané. Figuring out how to quiet them in people could enable a doubly effective assault against a massive public health problem.

More information: Marc Schneeberger et al, Regulation of Energy Expenditure by Brainstem GABA Neurons, *Cell* (2019). [DOI: 10.1016/j.cell.2019.05.048](#)

Provided by Rockefeller University

APA citation: New research raises prospect of better anti-obesity drugs (2019, June 27) retrieved 29 November 2021 from <https://medicalxpress.com/news/2019-06-prospect-anti-obesity-drugs.html>

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