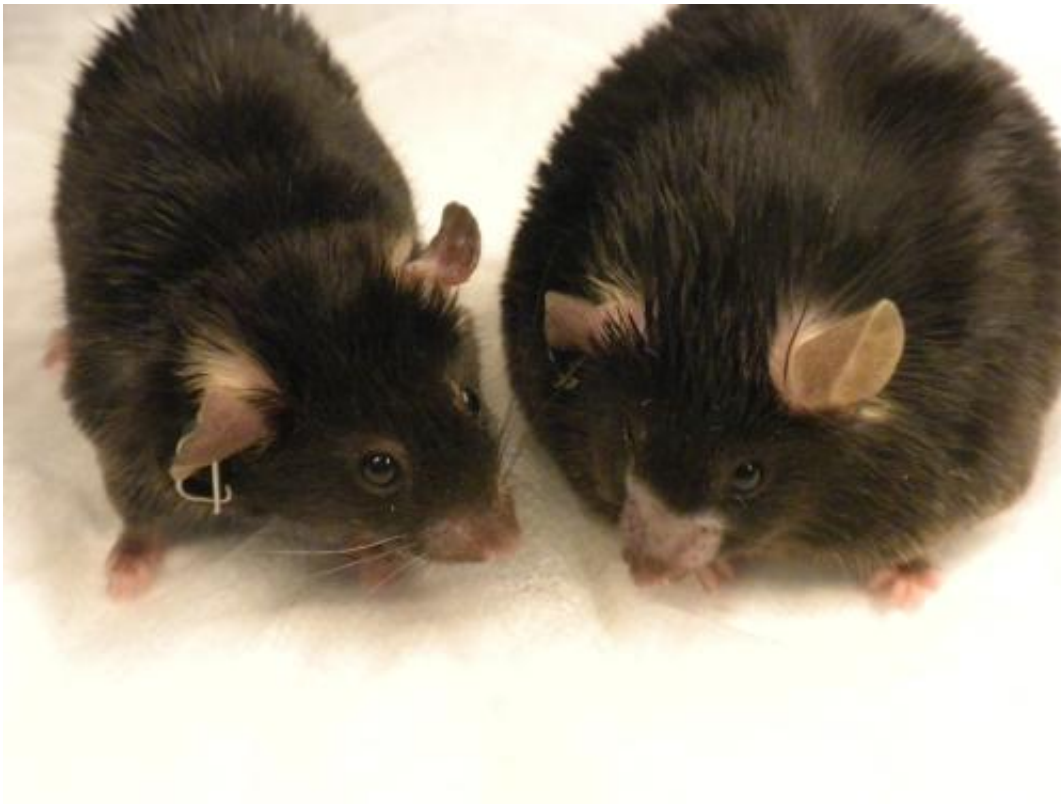


Old drug may point the way to new treatments for diabetes and obesity

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The obese mouse on the right was fed a high-fat diet. The mouse on the left was fed the same diet but is a normal weight after receiving amlexanox. Credit: Shannon Reilly.

Researchers at the University of Michigan's Life Sciences Institute have found that amlexanox, an off-patent drug currently prescribed for the treatment of asthma and other uses, also reverses obesity, diabetes and

fatty liver in mice.

The findings from the lab of Alan Saltiel, the Mary Sue Coleman director of the Life Sciences Institute, are scheduled to be published online Feb. 10 in the journal *Nature Medicine*.

"One of the reasons that diets are so ineffective in producing weight loss for some people is that their bodies adjust to the reduced calories by also reducing their metabolism, so that they are 'defending' their body weight," Saltiel said. "Amlexanox seems to tweak the metabolic response to excessive calorie storage in [mice](#)."

Different formulations of amlexanox are currently prescribed to treat asthma in Japan and canker sores in the United States. Saltiel is teaming up with clinical-trial specialists at U-M to test whether amlexanox will be useful for treating obesity and diabetes in humans. He is also working with medicinal chemists at U-M to develop a new compound based on the drug that optimizes its formula.

The study appears to confirm and extend the notion that the genes IKKE and TBK1 play a crucial role for maintaining metabolic balance, a discovery published by the Saltiel lab in 2009 in the journal *Cell*.

"Amlexanox appears to work in mice by inhibiting two genes—IKKE and TBK1—that we think together act as a sort of brake on metabolism," Saltiel said. "By releasing the brake, amlexanox seems to free the [metabolic system](#) to burn more, and possibly store less, energy."

Using high-throughput [chemical screening](#) at LSI's Center for [Chemical Genomics](#) to search for compounds that inhibit IKKE and TBK1, the researchers hit upon an approved off-patent drug: amlexanox. They then demonstrated that amlexanox had profound [beneficial effects](#) in both genetic and dietary-induced [obese mice](#). The chemical lowered the

weight of obese mice and reversed related metabolic problems such as diabetes and fatty liver.

"These studies tell us that, at least in mice, the IKKE/TBK1 pathway plays an important role in defending body weight by increasing storage and decreasing burning of calories, and that by inhibiting that pathway with a compound, we can increase metabolism and induce weight loss, reverse diabetes and reduce fatty liver," Saltiel said.

The drug has been on the market in Japan for more than 25 years.

However, the researchers don't yet know if humans respond with the same pathway, or if the discovery of amlexanox's effectiveness in mice can lead to a compound that is safe and effective for treating obesity and [diabetes](#) in humans.

"We will be working hard on that," Saltiel said.

More information: [dx.doi.org/10.1038/nm.3082](https://doi.org/10.1038/nm.3082)

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