

## Natural plant compound prevents Alzheimer's disease in mice

January 28 2014



Pamela Maher and Antonio Currais. Credit: Salk Institute for Biological Studies

(Medical Xpress)—A chemical that's found in fruits and vegetables from strawberries to cucumbers appears to stop memory loss that accompanies Alzheimer's disease in mice, scientists at the Salk Institute for Biological Studies have discovered. In experiments on mice that normally develop Alzheimer's symptoms less than a year after birth, a daily dose of the compound—a flavonol called fisetin—prevented the progressive memory and learning impairments. The drug, however, did not alter the formation of amyloid plaques in the brain, accumulations of proteins



which are commonly blamed for Alzheimer's disease. The new finding suggests a way to treat Alzheimer's symptoms independently of targeting amyloid plaques.

"We had already shown that in normal animals, fisetin can improve memory," says Pamela Maher, a senior staff scientist in Salk's Cellular Neurobiology Laboratory who led the new study. "What we showed here is that it also can have an effect on animals prone to Alzheimer's."

More than a decade ago, Maher discovered that fisetin helps protect neurons in the brain from the effects of aging. She and her colleagues have since—in both isolated cell cultures and mouse studies—probed how the compound has both antioxidant and anti-inflammatory effects on cells in the brain. Most recently, they found that fisetin turns on a cellular pathway known to be involved in memory.

"What we realized is that fisetin has a number of properties that we thought might be beneficial when it comes to Alzheimer's," says Maher.

So Maher—who works with Dave Schubert, the head of the Cellular Neurobiology Lab—turned to a strain of mice that have mutations in two genes linked to Alzheimer's disease. The researchers took a subset of these mice and, when they were only three months old, began adding fisetin to their food. As the mice aged, the researchers tested their memory and learning skills with water mazes. By nine months of age, mice that hadn't received fisetin began performing more poorly in the mazes. Mice that had gotten a daily dose of the compound, however, performed as well as normal mice, at both nine months and a year old.

"Even as the disease would have been progressing, the fisetin was able to continue preventing symptoms," Maher says.

In collaboration with scientists at the University of California, San



Diego, Maher's team next tested the levels of different molecules in the brains of mice that had received doses of fisetin and those that hadn't. In mice with Alzheimer's symptoms, they found, pathways involved in cellular inflammation were turned on. In the animals that had taken fisetin, those pathways were dampened and anti-inflammatory molecules were present instead. One protein in particular—known as p35—was blocked from being cleaved into a shorter version when fisetin was taken. The shortened version of p35 is known to turn on and off many other molecular pathways. The results were published December 17, 2013, in the journal *Aging Cell*.

Studies on isolated tissue had hinted that fisetin might also decrease the number of <u>amyloid plaques</u> in Alzheimer's affected brains. However, that observation didn't hold up in the mice studies. "Fisetin didn't affect the plaques," says Maher. "It seems to act on other pathways that haven't been seriously investigated in the past as therapeutic targets."

Next, Maher's team hopes to understand more of the molecular details on how fisetin affects memory, including whether there are targets other than p35.

"It may be that compounds like this that have more than one target are most effective at treating Alzheimer's disease," says Maher, "because it's a complex disease where there are a lot of things going wrong."

They also aim to develop new studies to look at how the timing of fisetin doses affect its influence on Alzheimer's.

"The model that we used here was a preventive model," explains Maher. "We started the mice on the drugs before they had any memory loss. But obviously human patients don't go to the doctor until they are already having memory problems." So the next step in moving the discovery toward the clinic, she says, is to test whether fisetin can reverse declines



in memory once they have already appeared.

**More information:** Currais, A., Prior, M., Dargusch, R., Armando, A., Ehren, J., Schubert, D., Quehenberger, O. and Maher, P. (2013), Modulation of p25 and inflammatory pathways by fisetin maintains cognitive function in Alzheimer's disease transgenic mice. *Aging Cell*. DOI: 10.1111/acel.12185

## Provided by Salk Institute

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