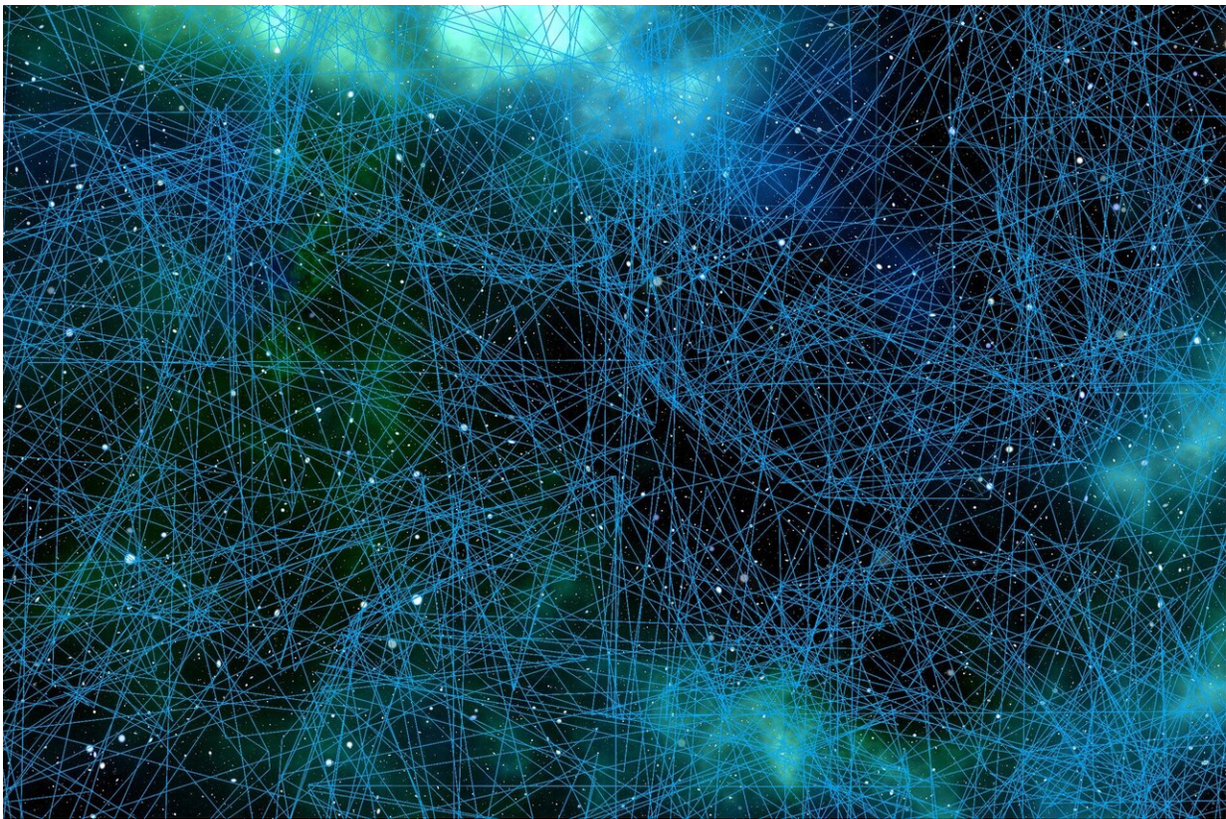


# Scientists succeed in reducing the levels of the molecules that kill neurons in elderly mice with Alzheimer's

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Alzheimer's disease is the most common dementia worldwide and it is characterized by learning impairment, memory loss, disorientation and

anxiety, among other behavioral and psychological symptoms. Since Alois Alzheimer described the first case at the beginning of the 20th century, important advances have been achieved in terms of understanding physiopathology, but, unfortunately, an effective treatment has not been developed yet.

What triggers Alzheimer's disease it is not well established, but several researches claim that the aggregation of the A $\beta$  peptide and of the tau protein, both present in normal brains, plays a pivotal role in Alzheimer's progression. Moreover, it is worth noting that there is a crosstalk between both, A $\beta$  peptide and tau [protein](#), which means that a reduction or an increase in one of them could affect the other. Thus, therapies directed to reduce A $\beta$ , and recently to reduce tau, have been explored. However, the use of full-length antibodies targeting A $\beta$  has not been effective in ameliorating cognitive decline and produced severe side effects in terms of neuroinflammation.

In previous studies, the Protein Design and Immunotherapy Group at the Department of Biochemistry and Molecular Biology of the UAB, led by Prof. Sandra Villegas, developed an [antibody fragment](#), scFv-h3D6, able to reduce the levels of A $\beta$  and to ameliorate [memory loss](#) in young, model of Alzheimer's disease, mice. However, what happens in advanced stages of the disease has never been studied before.

In a recent study published in the *International Journal of Molecular Sciences*, the group has demonstrated that scFv-h3D6 is also effective in reducing A $\beta$  levels in old mice, where the pathology is advanced the most. Interestingly, A $\beta$  reduction because of scFv administration also produced a reduction in total tau levels and did not elicit any detectable side effect. Furthermore, a slight reduction in the neuroinflammatory extent featuring the AD brain was observed. Therefore, the treatment is efficient and safe.

"The main point of this study is that we have shown that therapy with antibody fragments works in the late phase of the disease, and not only decreases the main molecule in Alzheimer's, A $\beta$ , but also reduces the levels of a second one, the [tau protein](#), which ultimately kills the neuron and causes dementia," explains Sandra Villegas.

Future studies targeting both A $\beta$  and tau pathologies in concert could show a putative pharmacological approach to treat this devastating [disease](#).

**More information:** Alejandro R. Roda et al. Both Amyloid- $\beta$  Peptide and Tau Protein Are Affected by an Anti-Amyloid- $\beta$  Antibody Fragment in Elderly 3xTg-AD Mice, *International Journal of Molecular Sciences* (2020). [DOI: 10.3390/ijms21186630](https://doi.org/10.3390/ijms21186630)

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