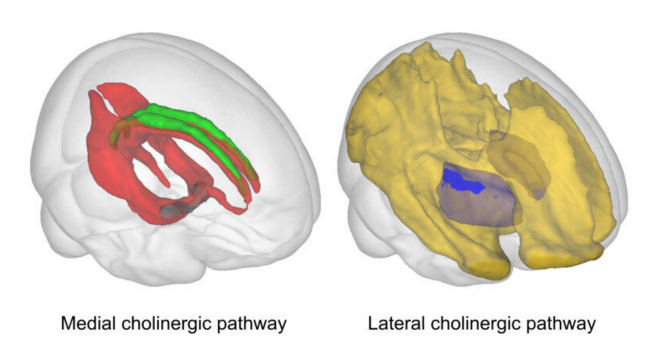


Novel imaging marker reveals very early brain changes in Alzheimer's disease

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Credit: Milan Nemy

A new collaborative study from Karolinska Institutet, Deutsches Zentrum für Neurodegenerative Erkrankungen (DZNE), and Czech Technical University suggests a novel imaging marker of brain connectivity might be a very early indicator of pathological changes in Alzheimer's disease. This imaging marker could be capable of detecting alterations when formal neuropsychological tests and even standard clinical image-based markers fail.



Alzheimer's disease can be regarded as a continuum. In that continuum, the very first stage includes individuals with positive biomarkers of Alzheimer's disease, among which individuals with self-reported memory complaints are at increased risk of progressing to later stages in the continuum. The later stages include individuals with <u>mild cognitive</u> <u>impairment</u> and, eventually, full-blown Alzheimer's disease dementia.

The neural cells in a strategic brain region—the cholinergic basal forebrain—and their long cholinergic axons are selectively vulnerable to Alzheimer's disease pathology. It is known that the hippocampus and basal forebrain are one of the first structures which show volumetric changes in the progressing disease.

We also know that neurons with long axons tend to follow a dying-back pattern of degeneration when defects in these long nerve fibers and in the transfer of information forego the death of the cell body. For this reason, it is desirable to investigate the neuronal pathways to uncover the very early phases of the disease.

For Alzheimer's disease, a very attractive but also perhaps slightly neglected target is the long and scattered pathways originating from the minuscule cholinergic nuclei of the basal forebrain. The importance of these cholinergic neural cells is highlighted by the only currently available treatment for Alzheimer's disease in Sweden, which targets these cholinergic cells.

Brain cholinergic pathways in vivo

This research includes three studies led by Dr. Daniel Ferreira from Karolinska Institutet in close collaboration with Milan Nemy, MSc. from Czech Technical University and Professor Stefan Teipel from University of Rostock—DZNE. In their studies, the researchers chose to investigate the brain fibers using a technique based on <u>magnetic resonance imaging</u>



and its diffusion-weighted sequence. This is a sequence sensitive to microscopic water movement and able to capture the direction of brain fibers.

The quality of tracking the desired pathways is dependent on a precise determination of the location of axonal bodies and other important landmarks facilitating the tracking process. The partner from DZNE came up with a high-resolution segmentation mask of the cholinergic basal forebrain nuclei, which was crucial to serve as starting points of the imaging method.

The researchers began the first study, published in *NeuroImage*, by locating these cholinergic pathways in cognitively unimpaired individuals from a community-based study from Spain led by Dr. Daniel Ferreira (the GENIC cohort). The researchers could observe that indeed the pathways found by the diffusion-weighted tracking followed the pattern known so far only from post-mortem studies, hence visualizing these pathways for the first time in a healthy brain.

These results were successfully replicated and extended in a second study, published in *Neurology*, that included an independent cohort—the Gothenburg H70 Birth Cohort Studies led by Professor Ingmar Skoog from Gothenburg University.

Having identified the pathways and their location, the researchers investigated their microstructural properties in several diagnostic groups in the third study, which was published in *Brain*. This time, the researchers had quite an extensive cohort at their hands with 172 patients with subjective cognitive decline, 66 with mild cognitive impairment, and 52 with Alzheimer's disease dementia from DZNE in Germany (the DELCODE cohort).

The researchers' initial hope was to see some significant differences



between cognitively healthy subjects and patients with Alzheimer's disease dementia. This would add additional validity to their approach having the a-priori knowledge about cholinergic depletion in Alzheimer's disease.

First, the researchers tested for the overall integrity of the cholinergic pathways. Not only did they observe significant differences between these two extreme groups of subjects in the Alzheimer's disease continuum, but also, to their surprise, they noticed that there were already pronounced alterations in the individuals with subjective cognitive decline—a clinical stage with the absence of objective cognitive impairment.

What's more, the <u>subjective cognitive decline</u> individuals did not differ in cerebrospinal fluid-based biomarkers with healthy controls, a set of biomarkers that are currently used in Sweden to support the clinical diagnosis of Alzheimer's disease in several specialized clinics. The researchers demonstrated these results in the whole clinical cohort but also in the amyloid-stratified subsample, suggesting the integrity of the cholinergic pathways, as captured by their method, might be an indicator of very early changes in the Alzheimer's disease continuum.

In this cross-sectional study, the researchers were able to uncover the spatial distribution of the differences in the white matter integrity of the cholinergic pathways across the stages of the disease. Already in the preclinical stage of Alzheimer's disease, the posterior cholinergic white matter showed early regional vulnerability.

With the more advanced stages of the disease, these differences were even more pronounced with additional anterior areas and in the white matter underneath the cholinergic neural bodies only affected in the most advanced stage of dementia. This observation could be a manifestation of the dying-back pattern of neuronal degeneration—now



observable in living people.

More information: Milan Nemy et al, Cholinergic white matter pathways make a stronger contribution to attention and memory in normal aging than cerebrovascular health and nucleus basalis of Meynert, *NeuroImage* (2020). DOI: 10.1016/j.neuroimage.2020.116607

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Provided by Karolinska Institutet

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