Different brain atrophy patterns may explain variability in Alzheimer's disease symptoms

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Mathematical modeling of brain scans of patients with Alzheimer's disease and others at risk for the devastating neurodegenerative disorder has identified specific patterns of brain atrophy that appear to be related to the loss of particular cognitive abilities. In their report that has been published online in the Proceedings of the National Academy of Sciences, a team of researchers at Massachusetts General Hospital (MGH) and the
National University of Singapore describe how different atrophy patterns may explain the different ways that Alzheimer's disease can be manifested in individual patients.

"The symptom severity and neurodegeneration can vary widely across patients in Alzheimer's disease," says Thomas Yeo, PhD, of the Martinos Center for Biomedical Imaging at MGH. "Our work shows that participants in this study exhibit at least three atrophy patterns - cortical, temporal or subcortical - that are associated with variability in cognitive decline not only in patients diagnosed with Alzheimer's but also in individuals with mild cognitive impairment or those who are cognitively normal but are at risk for Alzheimer's." In addition to his affiliation with the Martinos Center, Yeo is an assistant professor in the Department of Electrical and Computer Engineering, Clinical Imaging Research Centre and Singapore Institute for Neurotechnology at the National University of Singapore.

The study analyzed data collected as part of the Alzheimer's Disease Neuroimaging Initiative (ADNI), a multi-institutional project to develop biomarkers - including blood tests, cerebrospinal fluid tests, and imaging studies - that can be used for diagnosis or in clinical trials. Yeo and his team - including investigators at the MGH and in Singapore - analyzed MR images taken of the brains of 378 ADNI participants when they enrolled in the study. Of these participants, 188 had been diagnosed with Alzheimer's disease; the others - 147 with mild cognitive impairment and 43 who were cognitively normal - were at increased risk based on levels in their brains of the beta-amyloid plaques that are characteristic of the disease.

As a first step, the research team analyzed data from the baseline structural MRIs using a mathematical model that estimated the probability that particular details of each image were associated with atrophy of a specific location within the brain. Based on the location of
atrophy factors, they determined three atrophy factor patterns: cortical -
representing atrophy in most of the cerebral cortex; temporal - indicating
atrophy in the temporal cortex (the cortical lobe behind the ears),
hippocampus and amygdala; and subcortical, indicating atrophy in the
cerebellum, striatum and thalamus, structures at the base of the brain.

Analysis of study participant scans taken two years later indicated that
atrophy factor patterns were persistent in individuals and did not reflect
different stages of disease. Most participants - including those in the
mild cognitive impairment and cognitively normal groups - showed
levels of more than one atrophy factor.

Behavioral and cognitive tests of study participants taken at six-month
intervals indicated associations between particular atrophy factor
patterns and specific cognitive deficits. Individuals in whom temporal
atrophy predominated had greater problems with memory, while cortical
atrophy was associated with difficulties with executive function - the
ability to plan and to accomplish goals. Individual differences in how
atrophy factors are distributed within the brain may allow prediction of
the rate at which cognitive abilities would be expected to decline.

"Most previous studies focused on patients already diagnosed, but we
were able to establish distinct atrophy patterns not only in diagnosed
patients but also in at-risk participants who had mild impairment or were
cognitively normal at the outset of the study," Yeo says. "That is
important because the neurodegenerative cascade that leads to
Alzheimer's starts years, possibly decades, before diagnosis. So
understanding different atrophy patterns among at-risk individuals is
quite valuable.

He adds, "Previous studies assumed that an individual can only express a
single neurodegenerative pattern, which is highly restrictive since in any
aged person there could be multiple pathological factors going on at the
same time - such as vascular impairment along with the amyloid plaques and tau tangles that are directly associated with Alzheimer's. So individuals who are affected by multiple, co-existing pathologies might be expected to exhibit multiple atrophy patterns."

Future research could further determine whether and how these atrophy patterns relate to the distribution of amyloid and tau and the mechanisms by which they affect specific cognitive abilities, Yeo explains. The same analytic approach also could be applied to other types of patient data and extended to other neurodegenerative disorder that produce varying symptom patterns, such as Parkinson's disease and autism.


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