Role of serum urate in neurocognitive function and dementia: new evidence contradicts old thinking

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In ARD, Latouret et al used the data from a community-based prospective French cohort study of healthy 4931 elderly people 65 years or older, examined at six clinical visits (including cognitive examinations) over 12 years, and analysed 1598 participants with a baseline serum urate level (serum uric acid (sUA)), no diagnosis of dementia, a Mini-Mental State Examination (MMSE) score of >24 and at least one follow-up visit.1 Dementia was diagnosed in a 3-step process, screening using the MMSE and the Isaccs Set Test by trained psychologists, additional neuropsychological testing by a physician, and adjudication based on criteria by an independent committee of neurologists. Dementia developed in 110 subjects during the 13,357 person years of follow-up. Multivariable-adjusted HR with the highest (≥5.8 mg/dL in men, ≥4.9 mg/dL in women) versus the lowest sUA quartile (≥4.37 and ≤3.51 mg/dL, respectively) was 1.79 for incident dementia (95% CI 1.17 to 2.73; p = 0.007). A strong association was seen with vascular or mixed dementia (HR=3.66 (95% CI 1.29 to 10.41), p=0.015), and no significant association was noted with Alzheimer’s disease (HR=1.55 (95% CI 0.91 to 2.61), p=0.10). Several important aspects of this study need to be carefully considered while interpreting findings: (1) patients on urate-lowering therapies (ULTs) were excluded; (2) there was no significant association between sUA levels and MRI markers of cerebrovascular disease or hippocampal volume; and (3) the association between sUA and vascular or mixed dementia was no longer significant, when adjusted for interim strokes. The authors carefully noted that these findings were not generalisable to hyperuricaemia or gout cohorts or to those younger than 65 years.

When one examines other studies in this area, the evidence is contradictory. Some studies showed that hyperuricaemia was associated with a lower risk of dementia,2–4 while other studies showed an opposite effect.5–10 A major limitation is that most of these studies providing the evidence were cross-sectional.

Two recently published systematic reviews carefully examined these data and provide a more comprehensive synthesis of the evidence. The first systematic review assessed whether sUA was associated with cognitive impairment and dementia.11 Across 31 studies, using mostly case–control data, sUA was lower in cases of dementia compared with non-dementia controls with a standardised mean difference (SMD) of −0.33 (95% CI −1.23 to 0.22), not statistically significant.12 Therefore, based on these systematic reviews, there is no convincing evidence to date that higher sUA levels are associated with a lower risk of dementia, except possibly in Parkinson’s disease-related dementia.

The current cohort study draws our attention to the association of the sUA level (hyperuricaemia) with the risk of dementia in the elderly using a population-based sample of the French elderly.1 The current study reported an association opposite to what has been a past concern, by showing a significant association of the highest baseline sUA quartile with a 1.8-times higher risk of dementia with up to 12-year follow-up. The study showed that the association of higher sUA level was stronger with vascular or mixed dementia compared with Alzheimer’s disease, hinting at different pathogenic mechanisms for these types of dementia as it relates to sUA levels.1 The lack of association of sUA levels and MRI markers of cerebrovascular disease is an equally interesting negative finding. This negative finding might be related to a small number of incident cases despite a large cohort sample size and/or low sensitivity of this MRI marker for early/incident dementia.

In general, a key challenge to any study of dementia or associated risk factors is its long asymptomatic period and a gradual onset in most cases. These challenges can be addressed by the development of more accurate biomarkers of early dementia, an active area of research that holds promise for the future.13–15 Thus, this study adds significantly to the current knowledge base that contains few prospective cohort studies.

This study1 like any well-done study, raises several important questions that future studies should attempt to address: (1) What impact would a change in sUA over time have on the risk of dementia in the elderly? (2) Would the effect be similar in somewhat younger patient populations, that is, those younger than 65 years? (3) Do these risks vary by the presence of cardiovascular or cerebrovascular disease? These are a few questions that this study raises, which can guide the planning of well-designed studies investigating these relationships in the future.
from patients with gout. The 2016 European League Against Rheumatism (EULAR) gout treatment guideline stated that for patients with gout being treated with ULT ‘sUA level <3 mg/dL is not recommended in the long term’, as part of one of the recommendations. They cited few, but not all the studies included in the systematic reviews, and the evidence cited was not from patients with gout. Therefore, the observational evidence used was low-quality evidence due to serious indirectness. The discrepancy in studies included by the EULAR task force and these systematic reviews may be due to the differences in the inclusion/exclusion criteria. A cautionary EULAR recommendation in the absence of high-quality evidence may appear clinically justified to many but may be viewed by others as needing more evidence before implementation. Lowering of sUA <3 mg/dL in patients with gout is not a common occurrence in the clinical practice, and therefore, clinical and research evidence related to it is very limited to none. This recommendation helps to draw more attention to this interesting clinical area, by generating a healthy debate. The data available to date, including the two systematic reviews discussed above, provide reassurance that the serum urate lowering that is typically achieved with oral ULT is unlikely to contribute to development of dementia (and could have potential benefits).

In the field of gout, there are several unanswered questions about sUA and the risk of dementia and the effect of ULTs on the risk of dementia. Currently, we do not know: (1) if a threshold for sUA exists in gout that is associated with a higher or lower risk of dementia; and (2) whether such lowering is safe for short term, but not long term, that is, is there a time threshold for sUA lowering? Longitudinal observational studies of an adequate sample of patients with gout as well as observational and randomised studies of ULT in patients with gout can shed some light on these issues. Such studies are now needed to clarify the role of sUA in dementia risk in patients with gout.

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