Detection of Cognitive Impairment and Dementia Using the Animal Fluency Test: The DECIDE Study

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ABSTRACT: Objectives: To evaluate the performance of a one-minute screening test measured against a validated 10-minute screening test for mild cognitive impairment (MCI) in detecting CI in patients aged ≥ 65 years with two or more vascular risk factors (VRF). Methods: Patients (n=1523) aged 65 years or older without documented CI symptoms or dementia with two or more VRF participated in this study set in Canadian primary care practice. Baseline data was collected, followed by the 1-minute animal fluency (AF) test and the 10-minute Montreal Cognitive Assessment (MoCA). Physicians (n=122) completed case reports during patient interviews and reported their diagnostic impression. AF test sensitivity, specificity, and accuracy in predicting a positive MoCA was assessed. Results: Study sample mean age was 79.7 years, 55% were female, 97.6% were Caucasian and 75% had ≤ 12 years of education. The AF test and MoCA detected CI in 52 and 56 percent of the study population, respectively. The AF test demonstrated sensitivity, specificity, and accuracy in predicting a positive MoCA of 67 percent each. Physicians diagnostic impression of MCI was reported for 37% of patients, and of dementia for 6%. Conclusion: In an elderly population with at least two VRF, using AF can be useful in detecting previously unknown symptoms of CI or dementia. Screening for CI in this high risk population is warranted to assist physician recognition of early CI. The short AF administration time favours its incorporation into clinical practice.

RÉSUMÉ: Détection du déficit cognitif et de la démence au moyen de l’Animal Fluency Test : l’étude DECIDE. Objectifs : Le but de l’étude était de comparer le résultat d’un test de dépistage d’une minute au résultat d’un test de dépistage validé de dix minutes pour le déficit cognitif léger (DCL) pour identifier les patients atteints d’un déficit cognitif (DC), chez des individus âgés de 65 ans ou plus qui présentent deux facteurs de risque vasculaires (FRV) ou plus. Méthodes : Des patients (n = 1 523) âgés de 65 ans ou plus, sans symptôme de DC établi ou de démence mais qui présentaient deux FRV ou plus, ont participé à cette étude effectuée dans le contexte de soins de première ligne au Canada. Les données initiales étaient recueillies et l’Animal Fluency Test (AF) d’une minute et le Montreal Cognitive Assessment test (MoCA) de dix minutes étaient ensuite administrés. Les médecins (n = 122) complétaient le cahier d’observation pendant l’entrevue avec le patient et rapportaient leur impression diagnostique. La sensibilité du test AF, sa spécificité et son exactitude pour prédire un MoCA positif ont été évaluées. Résultats : L’âge moyen des sujets était de 79,7 ans, 55% étaient des femmes, 97,6% étaient des caucasiens et 75% avaient 12 ans ou moins de scolarité. Le test AF et le test MoCA ont détecté une DC chez 52% et 56% des sujets étudiés respectivement. La sensibilité du test AF, la spécificité et l’exactitude à prédire un test MoCA positif étaient toutes deux de 67%. Selon l’impression diagnostique du médecin il s’agissait d’une DCL chez 37% des patients et d’une démence chez 6%. Conclusion : Dans une population âgée présentant au moins deux FRV, l’AF peut être utile pour détecter les symptômes d’un DC ou d’une démence jusque là non reconnus. Le dépistage d’un DC chez cette population à haut risque est justifié pour aider les médecins à identifier le début d’un DC. Le peu de temps requis pour compléter l’AF facilite son intégration en pratique clinique.


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Dementia rates increase with age; an analysis of 22 studies reported a doubling of dementia prevalence rates every 5.1 years until the age of 95. The yearly incidence of dementia in Canada, among individuals aged 65 years and older, is estimated at 19.1/1000 for men and 21.8/1000 for women. The prevalence of dementia is expected to increase over the next decades, as a result of the aging population.

Vascular events have been implicated in the pathogenesis of cognitive impairment (CI). Risk factors for CI leading to dementia include vascular disease, hypertension, diabetes,
circuits. Animal fluency has been shown to distinguish patients with dementia from normal controls. Semantic fluency has been demonstrated to be a sensitive and specific measure of early cognitive decline. The animal fluency (AF) test provides an assessment of semantic fluency by asking patients to name as many animals as they can in one minute. Deficits in these cognitive domains are thought to be associated with damage to the temporal and parietal lobes as well as frontal-subcortical circuits. Animal fluency has been shown to distinguish patients with dementia from normal controls. Semantic fluency has been shown to be reduced in patients with MCI compared to cognitively intact controls.

Canadian consensus guidelines recommend that physicians maintain a high index of suspicion for dementia and follow-up on their concerns when dealing with elderly patients that demonstrate memory loss and functional decline. It is recommended that physicians perform a comprehensive assessment for elderly patients with symptoms of CI, cognitive complaints or concerns. In real world clinical practice, proceeding from suspicion to diagnosis of dementia is often a lengthy process that does not occur at early stages when it would be the most helpful. As such, a care gap exists with the prevalence of undiagnosed dementia in its early stages remaining high; in a recent study, as few as 19% of patients with confirmed dementia had documentation of the illness on their medical record.

The objective of this study was to evaluate— in the setting of routine primary care practice— the performance of a one-minute screening test measured against a validated ten-minute screening test for MCI in detecting CI in patients aged ≥ 65 years who had two or more vascular risk factors (VRF).

**Methods**

**Study Design**

DECIDE was an evaluation of a screening tool for MCI conducted at primary care settings across Canadian provinces (from September, 2005 to March, 2006). The study adhered to Good Clinical Practice and International Conference on Harmonization Guidelines, and the protocol was approved by appropriate independent research ethics committees (Institutional Review Board Services, Aurora, Ontario, and the University of Calgary Office of Medical Bioethics).

**Subjects**

To participate in the study, the patients (n=1523) had to be at least 65 years-of-age, provide written consent, and have two or more of the following risk factors: type 2 diabetes; hypertension; hypercholesterolemia; cigarette smoking (current or past); obesity (BMI ≥ 30); coronary artery disease (myocardial infarction or angina or coronary artery bypass graft); cerebrovascular disease (stroke or transient ischemic attack where deficits did not preclude administration of the AF test and MoCA); atrial fibrillation; congestive heart failure; or peripheral vascular disease. Patients were enrolled as they visited their primary care physicians for a routine visit.

Study exclusion criteria included: a diagnosis of dementia or documented CI prior to the study visit; evidence of current clinically significant depression or current alcohol abuse as defined by DSM-IV-TR criteria; enrollment in the present study on a previous occasion; and concurrent participation in a study with an experimental drug or a cholinesterase inhibitor. Patients
who had any condition which, in the physician’s judgment, might decrease the chance of obtaining satisfactory data to achieve the objectives of the study, were unable or unlikely to understand the nature, scope and possible consequences of the study, showed evidence of an uncooperative attitude, or were not sufficiently proficient in English or French to be assessed by the study instruments were also excluded from the study.

A stratified approach to patient enrollment was undertaken to optimize the balance between obtaining patient enrollment across all age groups and obtaining sufficient data on detected cases of CI. Each physician was asked to enroll three patients aged 65-74 years, seven patients aged 75-84 years, and five patients aged 85 years or older. Physicians continued their normal prescribing practices throughout the study.

Data Collection

During a single study visit for each patient, baseline data was collected (two-page clinical data form with: inclusion/exclusion criteria; socio-demographics; vascular risk factors and their levels of control; current and past medication use; family history; information regarding memory changes; current clinical impression and intended management approach). Primary care physicians completed case report forms during the patient interviews. They participated in a training session detailing the administration of the AF test and MoCA. The tests were administered by physicians and completed for each patient. Scores of 25 or less on the 30 point MoCA were considered indicative of CI, whereas the threshold for CI on the AF test was a score below 15.

Data Analyses

We used descriptive statistics to summarize the demographic and clinical characteristics of the sample expressed as mean for continuous variables and count (percent) for categorical variables. The purpose of the analysis was to compare the effectiveness of AF in detecting CI with the MoCA. As such, positive detection results with the AF test were considered true positive when they concurred with corresponding positive MoCA results and positive AF test results were considered false positive when the corresponding MoCA result was negative.

In the primary analysis, the AF positive predictive value was calculated as: (number of true positive AF test results)/(all positive AF test results), and the AF negative predictive value was determined by: (number of true negative AF test results)/(all negative AF test results). Compared to MoCA, AF test sensitivity was calculated as: (number of true positive AF test results)/(number of positive MoCA results), specificity was calculated using: (number of true negative AF test results)/(number of negative MoCA results), and accuracy was determined by the formula: (number of true positive AF test results + number of true negative AF test results)/(all positive AF test results + all negative AF test results). The discriminating power of the AF test relative to the threshold for CI (i.e., fewer than 15 animals named) was evaluated using Receiver Operator Characteristic (ROC) analysis, which is a technique that illustrates the trade-offs between the true positive rate (sensitivity) and the false positive rate (1-specificity) for every possible cutoff. The global index of screening accuracy is measured by the area under the ROC curve (AUC), where an AUC of 1 represents a perfect screening test, and 0.5 corresponds to random assignment. All analyses were performed using SAS 9.2 (Cary, NC).

RESULTS

Patient characteristics are listed in Table 1. The mean age of the study population was 79.7 years [stratified into three groups: 65-74 years (n = 363), 75–84 years (n = 704), 85 years and older (n = 443)]. Fifty-five percent were female, Caucasians represented 97.6 percent of subjects, and 75 percent had less than 12 years of education. The most common cardiovascular disease (CV) risk factors were: high blood pressure (81%); elevated cholesterol (58%); coronary artery disease (37%); and diabetes (33%). The presence of multiple CV risk factors was common; 63 percent of subjects had three or more CV risk factors. The prevalence of concomitant drug treatments for CV risk factors was high among the study population with 87 percent of patients taking antihypertensive medications, 65 percent taking aspirin, 55 percent taking statins, and 27 percent taking anti-diabetic medications. Twenty-five percent of patients were taking sedatives or hypnotics.

Table 1: Patient characteristics at baseline

<table>
<thead>
<tr>
<th>Patient characteristic</th>
<th>1512</th>
</tr>
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<tbody>
<tr>
<td>65-74 yrs/75-84 yrs/85+ years (%)</td>
<td>24/47/29</td>
</tr>
<tr>
<td>Sex, M:F (%)</td>
<td>45:55</td>
</tr>
<tr>
<td>Caucasian (%)</td>
<td>97.6</td>
</tr>
<tr>
<td>≤ 12 years education (%)</td>
<td>75</td>
</tr>
<tr>
<td>Taken university courses (%)</td>
<td>17</td>
</tr>
<tr>
<td>Prevalence of individual vascular risk factors (%)</td>
<td></td>
</tr>
<tr>
<td>High blood pressure</td>
<td>81</td>
</tr>
<tr>
<td>Elevated cholesterol</td>
<td>58</td>
</tr>
<tr>
<td>Coronary artery disease</td>
<td>37</td>
</tr>
<tr>
<td>Diabetes</td>
<td>33</td>
</tr>
<tr>
<td>Past smoker</td>
<td>26</td>
</tr>
<tr>
<td>Current smoker</td>
<td>7</td>
</tr>
<tr>
<td>Elevated BMI</td>
<td>19</td>
</tr>
<tr>
<td>Cerebrovascular disease</td>
<td>14</td>
</tr>
<tr>
<td>Atrial fibrillation</td>
<td>13</td>
</tr>
<tr>
<td>Peripheral vascular disease</td>
<td>11</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td>9</td>
</tr>
<tr>
<td>2/3/4/5+ vascular risk factors (%)</td>
<td>37/34/19/10</td>
</tr>
<tr>
<td>Family history of dementia or depression (%)</td>
<td></td>
</tr>
<tr>
<td>Alzheimer’s disease</td>
<td>10</td>
</tr>
<tr>
<td>Dementia</td>
<td>5</td>
</tr>
<tr>
<td>Depression</td>
<td>7</td>
</tr>
</tbody>
</table>
Although none of the study subjects had documented CI prior to the study visit, the percentage of patients identified as having CI (Table 2) was 52 percent as assessed by the AF test, and 56 percent according to the MoCA test. The primary analysis of this study compared the overall effectiveness of the AF test relative to the MoCA in detecting CI (Table 2). Compared to MoCA, AF demonstrated sensitivity of 67 percent, specificity of 67 percent, and accuracy of 67 percent. The AF test provided a positive predictive value of 72 percent and a negative predictive value of 62 percent relative to the MoCA. The global index of screening accuracy (AUC under the ROC curve) was 0.74, which corresponds to an acceptable level of discrimination.30 The comparison of the sensitivity and specificity as a function of different cutoff points indicated that the number of false positives increased rapidly beyond a threshold of 15 or greater animals named, with relatively limited increase of the true positives (<14: specificity 58%, sensitivity 76%; <16: specificity 76%, sensitivity 57%).

Plotting the data by patient score on both the AF test and the MoCA, Figure A shows the results in the quadrant labelled ‘true positive’ for those patients scoring positive for CI on both tests, and in the quadrant labeled ‘true negative’ for those scoring negative for CI on both tests. The scatter plot of results indicates that as the MoCA score increases, so does the number of correct answers in the AF test. The association between AF and MoCA scores demonstrates a statistically significant relationship between the two measures, with most false negative and false positive test results localized near the AF test and MoCA thresholds for CI.

**Table 2: Overall effectiveness of the animal fluency test relative to the Montreal Cognitive Assessment in detecting cognitive impairment**

<table>
<thead>
<tr>
<th>Outcome measure</th>
<th>AF (n=1507)</th>
<th>MoCA (n=1507)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive for cognitive impairment</td>
<td>787</td>
<td>843</td>
</tr>
<tr>
<td>Negative for cognitive impairment</td>
<td>720</td>
<td>664</td>
</tr>
<tr>
<td>Prevalence (%)</td>
<td>52</td>
<td>56</td>
</tr>
<tr>
<td>Sensitivity n (%)</td>
<td>567/843 (67)</td>
<td></td>
</tr>
<tr>
<td>Specificity n (%)</td>
<td>444/664 (67)</td>
<td></td>
</tr>
<tr>
<td>Accuracy n (%)</td>
<td>1011/1507 (67)</td>
<td></td>
</tr>
<tr>
<td>Positive predictive value n (%)</td>
<td>567/787 (72)</td>
<td></td>
</tr>
<tr>
<td>Negative predictive value n (%)</td>
<td>444/720 (62)</td>
<td></td>
</tr>
</tbody>
</table>

AF: Animal Fluency; MoCA: Montreal Cognitive Assessment

**DISCUSSION**

Administered by physicians in primary care practice, the one-minute AF test is promising in detecting different levels of CI. When administered by physicians to patients without recognized CI or dementia, the AF test identified CI in 52 percent of patients compared to 56 percent by MoCA. Physician impression after administration of the AF and MoCA tests was that 43 percent of patients were cognitively impaired. Compared to MoCA, AF test accuracy in this population was 67 percent. The comparison of the sensitivity and specificity as a function of different cutoff points supported that a threshold of 15 animals named provided a good balance between true positive (sensitivity) and false positive (specificity) of the animal verbal fluency test for the screening of cognitive impairment.

A scatter plot of AF test scores versus MoCA scores demonstrated a trend towards a linear relationship between the number of correct answers in the AF test and the MoCA score. The few outliers with false negative and false positive test results tended to cluster near the AF test and MoCA thresholds for CI.

defined as MCI (37%) or dementia (6%), versus 52 percent by the AF test and 56 percent by the MoCA.

![Figure: A. Scatter plot of animal fluency (AF) versus Montreal Cognitive Assessment (MoCA) results in detecting cognitive impairment. B. Percentage of patients identified as having cognitive impairment using the AF test, the MOCA, or by physician diagnostic impression.](https://www.cambridge.org/core/terms).
The present study assessed the performance of a CI screening instrument in the context of a busy primary care practice. Physicians have identified limited available time as a barrier to dementia diagnosis. A brief screening instrument that serves to identify patients at-risk for CI or dementia and for whom further assessment is warranted would permit physicians to commit resources efficiently. The AF test is quick to administer—an important characteristic for the routine implementation of CI screening in the primary care setting.

The prevalence of dementia is expected to rise substantially in the coming years, as will its impact on patients and society. Dementia is under-diagnosed; more than 50 percent of patients with dementia have not been diagnosed. In the present study, 43 percent of patients were eventually judged by their physicians as having CI and, in some cases, dementia even though a diagnosis of CI or dementia were study exclusion criteria. It is important to identify CI before it progresses to dementia. In cases where dementia has developed, initiating treatment for dementia early in the disease process can be beneficial. In those with early stage disease, cholinesterase inhibitors have been shown to consistently slow the rate of decline in cognitive and global clinical change scores, as well as functional and behavioural measures that can contribute to a reduction of patient and caregiver burden. In the future, when disease modifying agents become available, it will be even more important to have practical useful cognitive screening tests.

Cardiovascular risk factors are positively associated with cognitive decline. Hypertension, heart disease, hypercholesterolemia, and other vascular risk factors are common among the elderly. Significant gains can be made by the timely identification of individuals at-risk and working towards reducing the modifiable risk factors. Among the more than 2800 subjects taking part in the Systolic Hypertension in Europe Study (SYST-EUR), long-term antihypertensive therapy led to a 55 percent reduction in dementia over a period of 3.9 years. Although an infarct in a particularly sensitive brain region can produce a dramatic effect on cognitive function, cognitive decline is often the result of multiple brain lesions accumulated over time. By identifying elderly patients with CV risk factors early, a simple screening test can help determine a course of action to reduce the progression rate to dementia. Delaying the onset of dementia would substantially decrease the prevalence of the disease, and reduce patient, caregiver and societal burdens.

Other tests for CI have been described elsewhere. The MMSE—a test which has high specificity and sensitivity for detecting dementia in outpatients older than 65 years—is the most commonly used test to screen for memory problems and for diagnosing dementia. However, the MMSE has a ceiling effect, demonstrating a limitation of the test in identifying early dementia. The MoCA has been shown to have higher sensitivity in detecting MCI compared to the MMSE. The present study showed that the brief AF test identified only four percent fewer individuals with CI than the MoCA.

Study Limitations

Although the MoCA was used here as a comparator, the true gold standard for the detection of CI is full clinical and psychometric testing. However, given the time requirement to conduct such an evaluation, it was deemed unrealistic to include this for the study or within the context of a busy clinical practice. The AF test is a brief instrument that assesses two domains of cognitive functioning whereas MoCA evaluates several cognitive domains. The simplicity of the AF test allows for brevity, however, some specificity is lost as evidenced by the 67 percent accuracy of the test when compared to another screening instrument—MoCA.

The one minute AF test is a short screening tool that may assist in screening for CI in the primary care setting in patients who are 65 years or older and have two or more vascular risk factors. By identifying patients at risk for progression to dementia, further testing and appropriate measures to counter the progression of the disease can be undertaken.

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References


