Evaluation of the Montreal Cognitive Assessment as a screening tool for cognitive dysfunction in SLE

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ABSTRACT

Objectives Cognitive dysfunction in SLE is common and associated with significant morbidity but is currently underdetected. Early detection requires the use of screening tests, as formal diagnostic cognitive testing is time-consuming. This study aims to evaluate the Montreal Cognitive Assessment (MoCA) as a screening tool for cognitive dysfunction in SLE.

Methods Patients with SLE (n=95) and demographically matched healthy control participants (n=48) underwent cognitive testing using the 1-hour neuropsychiatric test battery recommended by the American College of Rheumatology for use in SLE and the MoCA. We used regression analyses to determine associations between MoCA and cognitive test scores. We assessed several MoCA cut-offs for predicting cognitive impairment in terms of sensitivity, specificity, positive predictive value and negative predictive value. Receiver operating curve analyses were used to determine the diagnostic accuracy of the MoCA cut-off thresholds.

Results We found a significant correlation between MoCA score and 9 of the 10 cognitive endpoints studied (all p<0.001). Receiver operating curve analysis suggested that a MoCA cut-off of <27 had highest diagnostic accuracy across the cognitive impairment definitions (area under the curve 0.76–0.78). Using a screening cut-off of <28, the MoCA had sensitivity of 83%–94% and specificity of 46%–59%, depending on the impairment definition used.

Conclusions The MoCA correlates strongly with cognitive test results in SLE and has sufficient sensitivity for use as a screening tool with a cut-off of <28 as the optimal threshold. This tool can be incorporated into clinical practice for screening for cognitive dysfunction in SLE.

INTRODUCTION

SLE is a chronic multisystem autoimmune disease associated with significant morbidity and reduced life expectancy. Cognitive dysfunction is common in SLE and can present insidiously. Many patients with SLE report cognitive dysfunction as one of the most distressing symptoms of their condition, adversely impacting function and employment. Although the pathogenesis of cognitive dysfunction in SLE remains poorly understood, early detection and recognition of cognitive changes may help us develop strategies that improve patients' quality of life.

Formal evaluation by a clinical neuropsychologist including cognitive testing remains the gold standard for the diagnosis of cognitive impairment, but is not practical for use on a routine basis in the SLE clinical care setting. Neuropsychological assessments themselves are time-consuming, and are expensive for routine use, because these services are often not covered by healthcare benefits or insurance. This further adds to the potential utility of a screening tool as a first step to identify patients who may benefit from comprehensive cognitive testing.

Key messages

What is already known about this subject?
- Cognitive dysfunction is common in SLE and has significant morbidity, but is currently underdetected as formal diagnostic testing is time-consuming and requires specialised staff.
- The Montreal Cognitive Assessment (MoCA) is a freely available, practical and brief cognitive screening test; the utility of the MoCA as a screening test for cognitive dysfunction in SLE has not been comprehensively studied.

What does this study add?
- Performance on the MoCA correlates strongly with formal cognitive test results and the MoCA has sufficient sensitivity to be used as a screening test for cognitive dysfunction in SLE.
- The optimal screening cut-off threshold for use of the MoCA in SLE is <28, which is higher than the cut-off of <26 used for mild cognitive impairment and dementia.

How might this impact on clinical practice or future developments?
- Screening for cognitive dysfunction with the MoCA should be incorporated into routine assessment of patients with SLE.
The ideal characteristics of a screening test include sufficient sensitivity to detect potential cases and that it should be easily accessible and simple to administer. It is essential that a screening tool is based on objective cognitive tests, as patient-reported symptoms are frequently discordant with objective tests and may be affected by factors such as mood disorders.

The Montreal Cognitive Assessment (MoCA) and the Mini Mental State Examination (MMSE) are both cognitive screening tools that meet these characteristics and have undergone preliminary assessment for use in SLE. The MoCA is a freely available brief screening tool that was initially designed in 1996 to screen for early cognitive impairment in the setting of dementia. Three studies have assessed the MoCA for use in SLE, and have suggested that it is a more sensitive test than the MMSE. Of the studies assessing the use of the MoCA in SLE, only one used a broad conventional cognitive test battery to define cognitive dysfunction. However, this study did not specify the method, definition or threshold used to define cognitive impairment on the cognitive test battery comparator and did not evaluate a range of MoCA cut-off thresholds. In addition, this study excluded patients with SLE with cerebrovascular disease or mood disorders.

The MoCA was initially designed in 1996 to screen for early cognitive impairment in the setting of dementia. Three studies in patients with SLE is known to affect a wide range of cognitive domains and hence it is also important that screening covers a broad range of domains.

The Montreal Cognitive Assessment (MoCA) and the Mini Mental State Examination (MMSE) are both cognitive screening tools that meet these characteristics and have undergone preliminary assessment for use in SLE. The MoCA is a freely available brief screening tool that was initially designed in 1996 to screen for early cognitive impairment in the setting of dementia. Three studies have assessed the MoCA for use in SLE, and have suggested that it is a more sensitive test than the MMSE. Of the studies assessing the use of the MoCA in SLE, only one used a broad conventional cognitive test battery to define cognitive dysfunction. However, this study did not specify the method, definition or threshold used to define cognitive impairment on the cognitive test battery comparator and did not evaluate a range of MoCA cut-off thresholds. In addition, this study excluded patients with SLE with cerebrovascular disease or mood disorders despite these being common comorbidities in SLE which may contribute to cognitive dysfunction, making these results less applicable to a typical SLE cohort in clinical practice. The Automated Neuropsychological Assessment Metrics test has also been evaluated in SLE but practical considerations limit its use as a screening tool, including accessibility, cost and testing time.

We aimed to address methodological deficiencies in these previous studies by analysing across different definitions of cognitive dysfunction using the American College of Rheumatology (ACR) recommended conventional cognitive test battery and testing multiple MoCA cut-offs to determine the optimum threshold for clinical application. The first objective was to evaluate the construct validity of the MoCA for screening in SLE by determining if there was an association between MoCA scores and performance on conventional cognitive testing. The second objective was to evaluate the performance of the MoCA as a screening tool in SLE by determining the sensitivity, specificity, positive predictive value, negative predictive value and diagnostic accuracy of various MoCA cut-off thresholds.

METHODS

Participants

Study participants for the SLE group (N=95) were recruited consecutively from the Monash Lupus Clinic site of the Australian Lupus Registry and Biobank (ALRB) between October 2018 and February 2020. The ALRB is a national registry of patients with SLE prospectively collecting longitudinal clinical data, blood and tissue samples since 2007. All enrolled patients fulfill either the 1997 ACR or the 2012 Systemic Lupus International Collaborating Clinics (SLICC) classification criteria. For this study, adults over the age of 65 years were excluded to avoid potential comorbid cognitive disorders associated with ageing. Patients with neurological conditions definitively not related to SLE (such as traumatic brain injury) were also excluded. Disease activity and damage were measured with the SLE Disease Activity Index 2000 (SLEDAI-2K) and SLICC-ACR Damage Index (SDI), respectively, as previously described.

A healthy control (HC) group was recruited as a comparator (N=48). The mean and range of age and premorbid IQ of the HC group were matched to the SLE group. HC participants were excluded if they had a history of autoimmune disease (except stable thyroid disease), any organ failure, central nervous system neurological condition or were on immunosuppressive therapy. HCs were recruited from family and friends of the SLE participants and via advertisement in the local community. All participants in both groups were English speaking and had completed at least part of their secondary schooling in English in order to ensure sufficient English language proficiency for the cognitive assessments. Participants provided informed consent and received no monetary compensation. Patients and the public were not involved in developing the study design. The Standards for Reporting of Diagnostic Accuracy Studies (STARD) guidelines for reporting diagnostic accuracy studies were used to ensure completeness and transparency.

The MoCA

The MoCA is a cognitive screening tool that takes approximately 10 min to administer, does not require specialised training and is freely available (it can be accessed here: https://www.moca.test.org/the-moca-test/). It consists of tasks examining visuospatial/executive function, naming, memory (delayed), attention, language, abstraction and orientation. The MoCA is scored out of 30 with 1 point added as an education adjustment for individuals who have completed less than 12 years of education. In screening for mild cognitive impairment or Alzheimer’s disease, the recommended cut-off score is <26. In this study, the MoCA was administered prior to the conventional cognitive test battery for all patients.

Conventional cognitive test battery

A single trained assessor (SR) administered the cognitive assessment using the 1-hour conventional neuropsychological test battery recommended by the ACR for use in SLE. The ACR battery has been validated in SLE against a more comprehensive 4-hour neuropsychological test battery with 90% agreement. The cognitive assessment component of this study was conducted under the guidance of a clinical neuropsychologist (YG-J).

Within the 15 subtest scores obtained from ACR test battery, there is some overlap in the domains tested. For
example, the battery includes several tests of psychomotor speed and tests that tap overlapping aspects of memory. Therefore, for the purpose of defining cognitive impairment, seven subtest scores with significant magnitude of effect in the SLE group were chosen as outcome measures to represent seven domain groups (see online supplemental file for detailed cognitive test descriptions):

- The *Rey-Osterrieth Complex Figure Test* recall task score was used to assess visual memory.
- The *California Verbal Learning Test* immediate recall (sum of trials 1–5) was used to measure verbal learning and memory.
- The *Controlled Oral Word Association Test* F, A and S trials summation score was used to assess verbal fluency.
- The *Letter Number Sequencing Test* from the *Wechsler Adult Intelligence Scale* (WAIS) fourth edition was used to assess verbal working memory and attention.

### Table 1 Demographic characteristics and cognitive test results of study groups

|                                | SLE group, N=95 | HC group, N=48 | Comparison* (p value) |
|--------------------------------|-----------------|----------------|----------------------|
| Age, median (range)            | 45 years (22–64) | 46 years (23–62) | 0.77                 |
| Gender, female                 | 93%             | 92%            | 0.84                 |
| Ethnicity                      |                 |                | 0.27                 |
| Caucasian                      | 62%             | 58%            |                      |
| Asian                          | 34%             | 42%            |                      |
| Other                          | 4%              | 0%             |                      |
| Premorbid IQ†, mean±SD         | 108.5±7.3       | 110.8±8.3      | 0.10                 |
| Education                      |                 |                | 0.14                 |
| Less than secondary            | 12%             | 6%             |                      |
| Secondary                      | 23%             | 13%            |                      |
| Tertiary                       | 54%             | 58%            |                      |
| Postgraduate                   | 12%             | 23%            |                      |
| Paid employment                | 60%             | 92%            | <0.001               |
| History of depression          | 37%             | 6%             | <0.001               |
| History of anxiety             | 27%             | 6%             | 0.003                |
| MoCA score, median (range)     | 26 (19–30)      | 28.5 (21–30)   | <0.0001              |
| Individual cognitive domain test scores‡, mean (range) | | | |
| Visual memory                  | 19 (3–32)       | 25 (12–33)     | <0.0001              |
| Verbal memory                  | 51 (19–70)      | 59 (45–71)     | <0.0001              |
| Verbal fluency                 | 41 (16–80)      | 53 (33–84)     | <0.0001              |
| Working memory                 | 18 (9–26)       | 21 (16–28)     | <0.0001              |
| Processing speed               | 69 (33–114)     | 84 (44–134)    | <0.0001              |
| Complex attention              | 80 (29–267)     | 57 (30–153)    | <0.0001              |
| Psychomotor speed              | 146 (89–193)    | 156 (116–204.5)| 0.008                |
| Cognitive impairment§          |                 |                |                     |
| 2 domains >1.5 SD below HC     | 49%             | 15%            | <0.001               |
| 1 domain >2 SD below HC        | 41%             | 10%            | <0.001               |
| 2 domains > 2 SD below HC      | 19%             | 0%             | 0.001                |
| All 3 definitions pooled       | 52%             | 16%            | <0.001               |

*Sociodemographic variables were compared between the SLE and control groups using Mann-Whitney, χ² and t-tests; cognitive test scores compared using one way analysis of variance.
†Premorbid IQ measured by Test of Premorbid Functioning scaled score.
‡Specific cognitive tests used for each domain are as follows: visual memory (Rey-Osterrieth Complex Figure Test recall score), verbal memory (California Verbal Learning Test trials 1–5), verbal fluency (Controlled Oral Word Association Test FAS sum), working memory (Letter Number Sequencing score), processing speed (Coding Score), complex attention (Trail Making Test B time in seconds (longer indicates worse performance)), psychomotor speed (Finger Tap Test dominant hand score).
§Impairment defined by number of cognitive domains either 1.5 or 2 SD below HC group mean, all three definitions pooled into fourth cognitive impairment category.

HC, healthy control; MoCA, Montreal Cognitive Assessment.
The Coding Test from the WAIS fourth edition (previously known as the Digit Symbol Substitution Test) was used to assess processing speed and attention.26 The Trail Making Test B time was used to assess complex attention and cognitive sequencing.27 The Finger Tap Test using the dominant hand (average number of taps per 25-second trial) was used to measure fine motor speed.28 The Test of Premorbid Functioning (TOPF) was used to estimate premorbid IQ.29 Because the pronunciation of known words is relatively resistant to impairment from brain pathology, the TOPF yields an estimate of premorbid verbal ability that is robust in the context of possible cognitive impairment.

Defining cognitive impairment
The percentage of patients with SLE with cognitive impairment was defined using SD from the HC group. The ACR 2007 response criteria for neurocognitive impairment in SLE clinical trials proposed the use of two SD thresholds to determine cognitive dysfunction:30 (1) two cognitive domains with SD >1.5 below the HC group mean, (2) one cognitive domain with SD >2 below the HC group mean and (3) two cognitive domains with SD >2 below the HC group mean. To capture the spectrum of cognitive impairment in SLE, we applied these three thresholds independently to categorising each participant as cognitively impaired or unimpaired. We added a fourth category of cognitive impairment pooling the first three definitions, categorising patients meeting any definition as being cognitively impaired.

Statistical analysis
To determine associations between MoCA scores and performance on conventional cognitive testing, we used multivariate analysis adjusting for age and premorbid IQ. We used linear regression to examine for relationships between MoCA scores and test performance in the seven individual cognitive domains, which were represented as z-scores in comparison with HC group data. We used logistic regression to examine associations between MoCA scores and cognitive impairment categories derived from the four ways of defining cognitive impairment separately. Pearson’s correlation coefficients were used to assess collinearity between covariates with likelihood ratios used.

Table 2 Clinical characteristics of patients with SLE

|                                | N (95) | %   |
|--------------------------------|--------|-----|
| Disease duration: median (range)| 15.0 years (0.2–38.7) |   |
| ANA positive                    | 94     | 99  |
| dsDNA positive                  | 77     | 81  |
| Anti-Smith positive             | 14     | 15  |
| APLS antibodies (any)           | 55     | 58  |
| APLS antibody triple positive   | 5      | 5   |
| History of cerebrovascular disease | 12    | 13  |
| History of seizures             | 8      | 8   |
| History of cranial neuropathy   | 5      | 5   |
| SLEDAI-2K score: median (range) | 3 (0–12) |   |
| SLICC-SDI score: median (range) | 1 (0–7)  |   |
| Medications (ever exposed)      |        |     |
| Hydroxychloroquine              | 89     | 94  |
| Prednisolone                    | 75     | 79  |
| Mycophenolate                   | 45     | 47  |
| Azathioprine                    | 37     | 39  |
| Methotrexate                    | 23     | 24  |
| Leflunomide                     | 5      | 5   |
| Rituximab                       | 5      | 5   |
| Cyclophosphamide                | 3      | 3   |
| Prednisolone dose (at test): median (range) | 0 mg (0–50) |   |

Serology was recorded as ever positive for each patient, usually ordered at baseline. APLS antibodies (anti-cardiolipin, beta-2 glycoprotein and lupus anticoagulant). APLS, antiphospholipid syndrome; dsDNA, double-stranded DNA; SLEDAI-2K, SLE Disease Activity Index 2000; SLICC-SDI, Systemic Lupus International Collaborating Clinics/American College of Rheumatology Damage Index.

Figure 1 Venn diagram of number of cognitively impaired patients in SLE group using different cognitive impairment definitions (out of n=95 total in SLE group). Impairment defined by number of cognitive domains either 1.5 or 2 SD below healthy control (HC) group mean to form three definitions above as described in coloured text. N impaired by each definition in brackets. N in overlap between definitions indicated by numbers in black text. All three definitions pooled into fourth cognitive impairment category (n=49 impaired).
Table 3  Multivariate analysis of the Montreal Cognitive Assessment (MoCA), age and cognitive test results in SLE

| Cognitive impairment definitions* | Individual cognitive domains† |
|----------------------------------|-----------------------------|
| 2 domains 1.5 SD below | 1 domain 2 SD below | 2 domains 2 SD below | All three definitions pooled |
| 1 domain 2 SD below | Coefficient (CIs) using linear regression |
| OR (Cls) using logistic regression |
| Age | 1.08** (1.02 to 1.13) | 1.07* (1.02 to 1.12) | 1.10* (1.02 to 1.21) | 1.05* (1.02 to 1.08) |
| | -0.04** (-0.06 to -0.02) | -0.02* (-0.03 to -0.01) | 0.004 | -0.02* (-0.03 to -0.01) |
| | -0.03** (-0.05 to -0.02) | -0.03** (-0.05 to -0.02) | -0.02** (-0.03 to -0.01) |
| MoCA score | 0.44** (0.31 to 0.63) | 0.49** (0.36 to 0.67) | 0.49** (0.35 to 0.69) | 0.49** (0.38 to 0.63) |
| | 0.26** (0.17 to 0.36) | 0.16** (0.09 to 0.22) | 0.24** (0.17 to 0.30) | 0.14** (0.08 to 0.20) |
| | 0.12** (0.06 to 0.18) | 0.28** (0.20 to 0.36) | 0.07 |

Multivariate analysis adjusted for age. Premorbid IQ was highly collinear with MoCA score and was therefore not included in the multivariate model. *p<0.05; **p<0.005.

†Specific cognitive tests used for each domain are as follows: visual memory (Rey-Osterrieth Complex Figure Test recall score), verbal memory (California Verbal Learning Test Trials 1–5), verbal fluency (Controlled Oral Word Association Test FAS sum), working memory (Letter Number Sequencing score), processing speed (Coding Score), complex attention (Trail Making Test B time inverse), psychomotor speed (Finger Tap Test dominant hand score). Test scores were expressed as z-scores in comparison with healthy control group data.
Lupus Science & Medicine

Raghunath S, et al. Lupus Science & Medicine 2021;8:e000580. doi:10.1136/lupus-2021-000580

Finally, we used ROCs to assess diagnostic accuracy of different MoCA cut-off scores. Comparing AUC for various MoCA cut-offs against the three cognitive impairment definitions suggested that <27 was the cut-off with the maximum diagnostic accuracy (figure 2). This threshold had the highest AUC for three definitions of cognitive impairment: two domains >1.5 SD below the HC group mean, one cognitive domain ≥2 SD below and all three definitions pooled (AUC 0.78, 0.76 and 0.75, respectively). For the most severe definition of cognitive impairment, two domains >2 SD below the HC group mean, the AUCs for <26 and <27 were similar (0.783 and 0.776, respectively). A cut-off of <28 had a similar AUC of 0.70–0.74, but superior sensitivity (table 4) and positive predictive value (table 5).  

**DISCUSSION**

Cognitive dysfunction is an important cause of functional impairment and disability in patients with SLE. Because formal diagnosis involves detailed neuropsychological testing that requires trained personnel, cognitive dysfunction in SLE is under-recognised. We sought here to assess the validity of a simple cognitive screening test in the identification of patients with SLE with cognitive impairment. We found a strong correlation between MoCA scores and formal cognitive test performance across a wide range of cognitive domains as defined by the ACR-recommended cognitive test battery, consistent with previous studies. 10 The cognitive domain of psychomotor speed was the only cognitive endpoint where there was no correlation with the MoCA score, however, this domain was the least affected in the cohort studied. Using a variety of definitions for cognitive dysfunction in order to capture the broad spectrum of involvement, we found that MoCA score had significant associations with all four definitions of cognitive impairment.

The MoCA had good diagnostic accuracy and adequate sensitivity for use as a screening test for cognitive dysfunction in SLE. These findings are consistent with previous preliminary studies in this area.10–12 However, our

| MoCA cut-off | Cognitive impairment definitions* | Sensitivity | Specificity |
|--------------|-----------------------------------|-------------|-------------|
| <24          | 2 domains 1.5 SD below | 23.1% | 98.2% |
| <25          | 1 domain 2 SD below | 38.5% | 94.6% |
| <26          | 2 domains 2 SD below | 72.2% | 84.4% |
| <27          | All 3 definitions pooled | 88.9% | 66.2% |

*Impairment defined by number of cognitive domains either 1.5 or 2 SD below healthy control group mean, all three definitions pooled into fourth cognitive impairment category.

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**DISCUSSION**

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| MoCA cut-off | Sensitivity | Specificity |
|--------------|-------------|-------------|
| <24          | 21.3%       | 100.0%      |
| <25          | 36.2%       | 97.9%       |
| <26          | 46.8%       | 93.8%       |
| <27          | 72.3%       | 83.3%       |
| <28          | 83.0%       | 58.3%       |
| <29          | 91.5%       | 35.4%       |
| <30          | 95.7%       | 20.8%       |

Table 4

**Table 5**

Positive and negative predictive value of Montreal Cognitive Assessment (MoCA) for cognitive impairment in SLE

| MoCA cut-off | 2 domains 1.5 SD below | 1 domain 2 SD below | 2 domains 2 SD below | All 3 definitions pooled |
|--------------|------------------------|---------------------|----------------------|-------------------------|
|              | PPV | NPV | PPV | NPV | PPV | NPV | PPV | NPV |
| <24          | 100.0% | 56.5% | 90.0% | 64.7% | 70.0% | 87.1% | 100.0% | 54.1% |
| <25          | 94.4% | 61.0% | 83.3% | 68.8% | 61.1% | 70.0% | 94.4% | 58.4% |
| <26          | 88.0% | 64.3% | 76.0% | 71.4% | 52.0% | 92.9% | 88.0% | 61.4% |
| <27          | 81.0% | 75.5% | 69.1% | 81.1% | 38.1% | 96.2% | 81.0% | 71.7% |
| <28          | 66.1% | 77.8% | 57.6% | 86.1% | 28.8% | 97.2% | 67.8% | 75.0% |
| <29          | 58.1% | 81.0% | 48.7% | 85.7% | 19.0% | 95.2% | 59.5% | 79.2% |
| <30          | 54.2% | 83.3% | 45.8% | 91.7% | 21.7% | 100.0% | 55.4% | 75.0% |

*Impairment defined by number of cognitive domains either 1.5 or 2 SD below healthy control group mean, all three definitions pooled into fourth cognitive impairment category.

NPV, negative predictive value; PPV, positive predictive value.
The cut-off specificity as the MoCA threshold decreased, and this was suggested <26 as having the highest AUC. Our study is in contrast to previous studies using ROC analyses which reported in previous studies in SLE. We saw a progression of cognitive dysfunction, which is lower than those ranging from 44.5% to 72.2% depending on the definition of cognitive impairment category.

The MoCA has established utility as a sensitive screening tool for detection of mild cognitive impairment in the elderly. For this population, the cut-off of 26 has a sensitivity of 90% and a specificity of 87%. In contrast, the sensitivity of the MoCA cut-off of <26 in our SLE cohort ranged from 44.5% to 72.2% depending on the definition of cognitive dysfunction, which is lower than those reported in previous studies in SLE. We saw a progressive decrease in sensitivity and corresponding increase in specificity as the MoCA threshold decreased, and this was observed across all definitions of cognitive impairment. The cut-off score of <27 was associated with the highest AUC across the four cognitive impairment definitions, in contrast to previous studies using ROC analyses which suggested <26 as having the highest AUC. Our study is the first to include patients with SLE with previous neuropsychiatric involvement and examine the performance of the MoCA screening tool against a range of cognitive impairment definitions.

There are multiple considerations when determining the optimal MoCA cut-off to use for screening purposes. For the purposes of screening, high sensitivity is favoured over high AUC or diagnostic accuracy, given that the primary aim is to detect all potential cases and follow these up with a more specific diagnostic test, namely conventional cognitive testing. However, higher sensitivity comes at the expense of specificity and therefore more false positives. Although in this context there are no high-risk follow-up tests such as biopsies, patients may be distressed by positive screening test results and the triggering of formal cognitive testing. In addition, access to neuropsychologists for formal testing may be limited in some settings, resulting in delay in moving from screening to formal result, contributing further to patient stress.

Keeping these considerations in mind, our findings suggest that the optimal cut-off MoCA score in screening for cognitive dysfunction in SLE population is <28, which had a sensitivity and specificity of 85%–94% and 45%–59%, respectively, and AUC of 0.70–0.74. For comparison, the most commonly used screening tool in the diagnosis of SLE is the ANA test, that has a similar sensitivity and specificity of 93% and 57%, respectively, and is considered to be an excellent screening tool.

Screening tools such as MoCA serve in the initial assessment of patients, providing a quick and highly sensitive assessment for identifying potential cognitive dysfunction, but in clinical practice patient symptoms and functional impact are also important considerations in determining whether further cognitive assessment is required. Patient-reported outcome tools to measure cognitive symptoms such as the Cognitive Symptoms Inventory or Perceived Deficits Questionnaire should also potentially be incorporated into the preliminary assessment, before recommendation of appropriate follow-up which may include further investigations such as neuroimaging. In addition, MoCA score was highly collinear with premorbid IQ and hence the effects of education level and premorbid IQ must also be considered in its interpretation in individual patients. Therefore, in individual patients with above or below average education levels or premorbid IQ, the change in MoCA score over time and patient-reported functional impact are particularly important considerations.

There are several limitations to the current study. The optimal MoCA cut-off may vary according to age, ethnicity, premorbid IQ and educational level. Our sample was highly educated, with 66% having completed some form of tertiary education. This may limit the ability to extrapolate our findings to populations with lower education levels. However, this effect was mitigated by a well-matched control group and the use of standard scoring guidelines for the MoCA, which include a one-additional point adjustment for those who have completed less than 12 years of education. In addition, in this multiethnic cohort, some participants were not native English speakers, but cognitive assessments were performed in English hence participants were excluded if they had not completed at least part of their secondary schooling in English. The MoCA has been translated into 36 languages and it is unclear how using multiple language versions of the MoCA in the same cohort may affect its validity as a screening tool. Finally, in this cross-sectional study, we did not explore the utility of repeated MoCA testing for long-term monitoring in SLE. The utility of serial MoCA in SLE for monitoring is not yet determined,

![Figure 2 Receiver operating curves (ROCs) for different Montreal Cognitive Assessment (MoCA) cut-offs by cognitive dysfunction (CD).](http://lupus.bmj.com/)

Co-morbidities

Raghunath S, et al. Lupus Science & Medicine 2021;8:e000580. doi:10.1136/lupus-2021-000580
even though in other disease states there is a suggestion of a value for serial testing.33

In summary, the MoCA provides a brief, freely available, practical screening tool for cognitive dysfunction, which our findings confirm has adequate sensitivity and specificity for use in SLE with a cut-off of ≤28. Given that cognitive dysfunction is commonly reported by patients with SLE, screening will improve detection in clinical practice, enabling future research to focus on addressing unmet needs in this domain.

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REFERENCES
1 Brey RL, Holliday SL, Saklad AR, et al. Neuropsychiatric syndromes in lupus: prevalence using standardized definitions. Neurology. 2002;58:1214–20.
2 Rayes HA, Tani C, Kwan A, et al. What is the prevalence of cognitive impairment in lupus and which instruments are used to measure it? A systematic review and meta-analysis. Semin Arthritis Rheum 2018;48:240–55.
3 Arnts CN, Wildman P, Gross D. Lupus: patient voices. Joint report by lupus research alliance, lupus Foundation of America and lupus and allied diseases association for the CDER and FDA. 2016 6 March 2018. Available: http://www.lupusfoundation.org/LupusPDDRReportCover.pdf
4 Panopoulos P, Julian L, Yazdany J, et al. Impact of memory impairment on employment status in persons with systemic lupus erythematosus. Arthritis Rheum 2007;57:1453–60.
5 Appenzeller S, Cendes F, Costallat LTL. Cognitive impairment and employment status in systemic lupus erythematosus: a prospective longitudinal study. Arthritis Rheum 2009;61:860–7.
6 Vogel A, Bhattacharya S, Larsen JL, et al. Do subjective cognitive complaints correlate with objective cognitive function in systemic lupus erythematosus? A Danish outpatient study. Lupus 2011;20:35–43.
7 Hanly JG, Su L, Omsiade A, et al. Screening for cognitive impairment in systemic lupus erythematosus. J Rheumatol 2012;39:1371–7.
8 Zabala A, Salgueiro M, Saez-Abuakkarro O, et al. Cognitive impairment in patients with neuropsychiatric and Non-neuropsychiatric systemic lupus erythematosus: a systematic review and meta-analysis. J Int Neuropsychol Soc 2018;24:629–39.
9 Nasreddine ZS, Phillips NA, Bedirian V, et al. The Montreal cognitive assessment, MoCA: a brief screening tool for mild cognitive impairment. J Am Geriatr Soc 2005;53:695–9.
10 Paez-Venegas N, Jordan-Estrada B, Chavarria-Avila E, et al. The Montreal cognitive assessment test: a useful tool in screening of cognitive impairment in patients with systemic lupus erythematosus. J Clin Rheumatol 2019;25:325–9.
11 Nantes SG, Su J, Dhalwai A, et al. Performance of screening tests for cognitive impairment in systemic lupus erythematosus. J Rheumatol 2017;44:1583–9.
12 Adhikari T, Piatti A, Luggen M. Cognitive dysfunction in SLE: development of a screening tool. Lupus 2011;20:1142–6.
13 Petri M, Naigbuddin M, Carson KA, et al. Depression and cognitive impairment in newly diagnosed systemic lupus erythematosus. J Rheumatol 2010;37:2032–8.
14 Kozora E, Ellison MC, West S. Reliability and validity of the proposed American College of rheumatology neuropsychological battery for systemic lupus erythematosus. Arthritis Rheum 2004;51:810–8.
15 Tayer-Shifman OE, Green R, Beaton DE, et al. Validity evidence for the use of automated neuropsychological assessment metrics as a screening tool for cognitive impairment in systemic lupus erythematosus. Arthritis Care Res 2020;72:1809–19.
16 O’Neill S, Morand EF, Hol A. The Australian lupus Registry and Biobank: a timely initiative. Med J Aust 2017;206:194–5.
17 Hochberg MC. Updating the American College of rheumatoid revised criteria for the classification of systemic lupus erythematosus. Arthritis Rheum 1997:40:1725.
18 Petri M, Orbai A-M, Alarcon GS, et al. Derivation and validation of the systemic lupus international collaborating clinics classification criteria for systemic lupus erythematosus. Arthritis Rheum 2012;64:2677–86.
19 Kompauer R, Németh HT, Niikura M, et al. Employment status in systemic lupus erythematosus patients with overt central nervous system disease. Arthritis Rheum 1996;39:2035–45.
20 Rey A. L’examen psychologique dans les cas d’encéphalopathie traumatique. (Les problems.) [The psychological examination in cases of traumatic encephalopathy, Problems]. Archives de Psychologie 1941;28:215–85.
21 Delis DC, Kramer JH, Kaplan E. California verbal learning Test-3. 3rd edn. San Antonio: TTP 2017.
22 Benton AL. Development of a multilingual aphasia battery. progress and problems. J Neurol Sci 1969;9:39–48.
Co-morbidities

26 Wechsler D. Wechsler adult intelligence scale. 4th edn (WAIS–IV), San Antonio, TX: NCS Pearson, 2008: 22. 1.
27 Bowie CR, Harvey PD. Administration and interpretation of the TRAIL making test. Nat Protoc 2006;1:2277–81.
28 Strauss E, Sherman EM, Spreen O. A compendium of neuropsychological tests: administration, norms, and commentary. American Chemical Society, 2006.
29 Wechsler D. The test of premorbid function (TOPF). San Antonio, TX: Corporation. TP, 2011.
30 Ad Hoc Committee on Lupus Response Criteria: Cognition Sub-committee, Mikdashi JA, Esdaile JM, et al. Proposed response criteria for neurocognitive impairment in systemic lupus erythematosus clinical trials. Lupus 2007;16:418–25.
31 Solomon DH, Kavanaugh AJ, Schur PH, et al. Evidence-Based guidelines for the use of immunologic tests: antinuclear antibody testing. Arthritis Rheum 2002;47:434–44.
32 Milani SA, Marsiske M, Cottler LB, et al. Optimal cutoffs for the Montreal cognitive assessment vary by race and ethnicity. Alzheimers Dement 2018;10:773–81.
33 Sivakumar L, Kate M, Jeerakathil T, et al. Serial Montreal cognitive assessments demonstrate reversible cognitive impairment in patients with acute transient ischemic attack and minor stroke. Stroke 2014;45:1709–15.