Altered cognitive function in systemic lupus erythematosus and associations with inflammation and structural brain changes

Michelle Barraclough,1,2 Shane McKie,3 Ben Parker,2,4 Alan Jackson,5 Philip Pemberton,6 Rebecca Elliott,7 Ian N Bruce1,2

ABSTRACT

Objectives Cognitive dysfunction (CD) is common in systemic lupus erythematosus (SLE) but the cause remains unclear and treatment options are limited. We aimed to compare cognitive function in SLE and healthy controls (HCs) using both behavioural and neuroimaging techniques.

Methods Patients with SLE with stable disease and HCs were recruited. Clinical and psychological data were collected along with a blood sample for relevant biomarkers. Neurocognitive function was assessed using tests from the Cambridge Neuropsychological Test Automated Battery (CANTAB) and functional magnetic resonance imaging (fMRI) was used to examine brain responses to working memory (WM) and emotional processing (facial emotional recognition task, FERT) tasks.

Results Compared with HCs (n=30), patients with SLE (n=36) scored higher on measures of depression, fatigue and had higher hsCRP (p=0.013), IL-6 (p=0.003) and B lymphocyte stimulator (p<0.001). Patients with SLE had poorer performance on a task of sustained attention (p=0.002) and had altered brain responses, particularly in default mode network (DMN) regions and the caudate, during the WM task. Higher organ damage and higher VCAM-1 were associated with less attenuation of the DMN (p=0.005 and p=0.01, respectively) and lower BOLD signal in the caudate areas (p=0.005 and p=0.001, respectively). Increased IL-6 was also associated with lower BOLD signal in caudate areas (p=0.032).

Conclusions Sustained attention was impaired in patients with SLE. Poor attenuation of the DMN may contribute to cognitive impairments in SLE and our data suggest that in addition to mood and fatigue inflammatory mechanisms and organ damage impact cognitive functioning in SLE. The multifaceted nature of CD in SLE means any therapeutic interventions should be individually tailored.

INTRODUCTION

Cognitive dysfunction (CD) is one of the most commonly reported neuropsychiatric symptoms in patients with systemic lupus erythematosus (SLE) and significantly affects quality of life. While it has been reported in up to 90% of patients, treatment options remain limited in large part due to uncertainty around the cause(s), the lack of a consistent measure and the observation that patients with SLE may perform similarly to healthy controls (HCs) on objective testing.

CD is common in other chronic conditions, such as inflammatory bowel disease and multiple sclerosis and factors associated with chronic disease such as mood disorders, medications and fatigue can all affect cognition. Specific SLE-factors are also hypothesised to play a role, with reported associations between autoantibodies and CD, particularly anti-N-methyl-D-aspartate, anti-dsDNA and anti-phospholipid (aPL) antibodies. Structural brain alterations in SLE such as cerebrovascular events, and the increased number of white matter hyperintensities also may contribute, although others have suggested that such structural changes are not directly associated with CD. As such, clinical and imaging biomarkers of CD in SLE remain elusive.
The American College of Rheumatology (ACR) established a recommended battery of cognitive tests but these require a trained professional to administer. An alternative is the Cambridge Neuropsychological Test Automated Battery (CANTAB), a tool that can assess changes in cognition over time, has been validated in many clinical settings, requires minimally trained administrators and has been successfully used in SLE.7 Another objective approach is functional magnetic resonance imaging (fMRI). fMRI provides a proxy measure of neuronal activation during cognitive testing. To date, only a few articles have reported fMRI findings in SLE but they suggest that patients with SLE may employ compensatory brain mechanisms within the brain to maintain adequate cognitive performance.8 Even fewer studies9,10 have examined cognition in SLE using a combination of behavioural, functional and structural assessments, although such an approach may better help identify causes and targets for therapy.

We aimed to compare cognitive function between patients with SLE with stable disease and HCs using CANTAB and fMRI. Variables that are known to affect cognition were also examined with differences between the two groups reported.

METHODS

Patients with SLE were recruited from Rheumatology departments at Manchester University NHS Foundation Trust Hospitals. HCs were recruited via study participants (e.g., friends) and social media. All SLE participants fulfilled ACR 199711 or Systemic Lupus International Collaborating Clinics (SLICC) criteria12 for SLE and were considered clinically stable if no change of treatment was required, and their Systemic Lupus Erythematosus Disease Activity Index-2000 (SLEDAI-2K) score was ≤4.13 Participants with a history of epilepsy, stroke, severe depression/psychiatric conditions or certain central nervous system (CNS)-acting medications were excluded. Severe depression was defined as currently receiving treatment and/or scoring ≥20 on the Montgomery Asberg Depression Rating Scale (MADRS). Participants on low-dose CNS-acting medications or who were taking no more than three such medications (and only if being used to treat conditions other than depression, such as fibromyalgia) were included.

Disease activity was assessed using the British Isles Lupus Assessment Group Index BILAG 200414 and SLEDAI-2K, and organ damage using the SLICC/ACR Damage Index (SDI).15 Specific biomarkers of the inflammatory response activation (B lymphocyte stimulator (BLyS), high sensitivity C reactive protein [hsCRP], interleukin 6 (IL-6) and vascular/endothelial (vascular cell adhesion molecule-1 (VCAM-1), vascular endothelial growth factor (VEGF)) were measured.

All participants completed validated questionnaires on depression, anxiety and fatigue: HADS: Hospital Anxiety and Depression Scale.16 BDI-II: Becks Depression Inventory-II.17 MADRS: Montgomery Asberg Depression Rating Scale.18 FSMC: Fatigue Scale for Motor and Cognitive Functions.19 After a literature review, we selected six CANTAB tests20 that assessed:

Immediate and delayed verbal memory (VRM: Verbal Recognition Memory).
Emotional processing (ERT: Emotional Recognition Test).
Sustained attention (RVP: Rapid Visual Information Processing).
Executive function (OTS: One Touch Stockings).
Spatial working memory (SWM: Spatial Working Memory).

Two functional scans were performed using a 3.0 Tesla Philips Gyroscan ACS NT (Philips, Best, NL) MR scanner while participants completed a WM task (n-back) and a facial emotional recognition task (FERT). Two structural scans were also performed: a fluid attenuated inversion recovery and T1-weighted magnetisation-prepared rapid gradient-echo.

All behavioural and assessment data were analysed using independent t-tests for parametric data, Mann-Whitney U tests for non-parametric data and χ² for proportional data in SPSS 22 and group region of interest analyses were undertaken for the fMRI data using SPM12.

The target number of participants recruited to the study was determined based on fMRI power guidance, where a sample size of between 16 and 32 is considered acceptable.21 To examine any possible associations between SLE and CD exploratory Pearson/Spearman’s correlations and χ² were undertaken using the SLE group only. These correlations were only conducted using the CANTAB tasks, structural brain abnormalities and fMRI results that were significantly different between the HC and SLE groups. These variables were assessed against factors proposed to affect cognition, including disease duration, disease activity, damage, medication use, aPL/LAC as well as measures of depression and fatigue.

Further details on all methods can be found in the online supplementary data.

RESULTS

Demographic and clinical findings

The SLE group were typical of a stable SLE cohort (table 1) and both groups were matched on age, gender, handedness and ethnicity; patients with SLE had fewer years in education and a lower mean IQ (table 2). The SLE group had higher depression scores (medians within the normal ranges). For each group, the percentages of participants that scored within the mild clinical ranges for depression/anxiety were: MADRS 0% HC, 6% SLE, BDI-II 6% HC, 33% SLE, HADS-D 3% HC, 39% SLE and HADS-A 20% HC, 39% SLE, despite excluding for clinical depression. The SLE group also had higher levels of motor and cognitive fatigue with median scores in the ‘severe’ (motor) and ‘moderate’ (cognitive) fatigue categories. Several biomarkers of inflammation and endothelial activation showed statistical difference between the two groups (table 2).

All measures of depression positively correlated with both cognitive and motor scores of fatigue (FMSMC) and negatively
with years in education (see online supplementary table S1). In addition, hsCRP positively correlated with HADS-depression score ($r=0.43$, $p=0.013$).

**CANTAB findings**
From the 66 participants (36 SLE and 30 HC) who underwent CANTAB testing, 2 SLE participants did not complete all tests due to fatigue.

The SLE group performed less well on the RVP task (a test of sustained attention) compared with the HC group (13 [12, 20] vs 20 [15.75, 22], $p=0.002$). Compared with the normative data available from CANTAB 33.3% of the SLE participants scored one or more SDs below the mean. The SLE group was also slower to identify emotions from the ERT task compared with the HC group ($t<0.001$). The differences between the SLE and HC group were more pronounced in the LSAT-R and LSAT-V conditions.

**Structural MRI findings**
Structural analysis was conducted on 53 participants (30 HC and 23 SLE). The SLE group had significantly more and larger regions where the BOLD signal reduced for both groups during the WM task. Significant results were found in the left transverse temporal gyrus (LTTG-WM) and right superior temporal gyrus (RSTG-WM) showed a more decreased BOLD response for the HC group compared with the SLE group. In the caudate the reverse was found, the SLE group had a more significant decrease in BOLD response compared with the HC group. Working memory condition (2-0back)
ROI analysis revealed significant results for the negative effect of the 2-back condition. This condition highlights regions where the BOLD signal reduced for both groups during the WM task. Significant results were found in the left transverse temporal gyrus ($t=2.12$, $p=0.039$), right superior temporal gyrus ($t=2.09$, $p=0.041$) and right caudate ($t=2.45$, $p=0.018$). The caudate reverse was also significant (SLE: $t=2.75$, $p=0.001$). The HC group responded correctly on the 1-back and 2-back levels (measures of WM, $p=0.019$ and $p=0.025$, respectively) compared with the HC group (see online supplementary table S5).

**Working memory condition (2-0back)**
ROI analysis revealed significant results for the negative effect of the 2-back condition. This condition highlights regions where the BOLD signal reduced for both groups during the WM task. Significant results were found in the left transverse temporal gyrus ($t=2.12$, $p=0.039$), right superior temporal gyrus ($t=2.09$, $p=0.041$) and right caudate ($t=2.45$, $p=0.018$). The caudate reverse was also significant (SLE: $t=2.75$, $p=0.001$). The HC group responded correctly on the 1-back and 2-back levels (measures of WM, $p=0.019$ and $p=0.025$, respectively) compared with the HC group (see online supplementary table S5).

**Structural MRI findings**
Structural analysis was conducted on 53 participants (30 HC and 23 SLE). The SLE group had significantly more and larger perivascular spaces (PVS) in the centrum semiovale (CSO-VRS), $\chi^2=15.50$, $p<0.001$. The differences between the SLE and HC group for the PVS in the basal ganglia (BG-VRS) did not reach significance, $\chi^2=8.96$, $p=0.077$ (see online supplementary tables S2-S4).

**Functional MRI findings**
Not all patients underwent an MRI scan due to scheduling, discomfort and artefact issues. Overall, 23 SLE and 29 HC participants had fMRI scan data available for analysis.

**n-back task results**
Patients with SLE performed worse than HC on the 0-back level (measure of attention, $p=0.008$) and were also slower to respond correctly on the 1-back and 2-back levels (measures of WM, $p=0.019$ and $p=0.025$, respectively) compared with the HC group (see online supplementary table S5).

---

### Table 1 Clinical and immunological characteristic of the SLE participants (n=36)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>n (%) or median (LQ, UQ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female sex</td>
<td>34 (94%)</td>
</tr>
<tr>
<td>Disease duration (years)</td>
<td>10.5 (5, 15)</td>
</tr>
<tr>
<td>ANA positive (ever)</td>
<td>34 (94.4)</td>
</tr>
<tr>
<td>Elevated IgG anti-dsDNA antibody*</td>
<td>9 (26)</td>
</tr>
<tr>
<td>Low C3 or C4*</td>
<td>12 (35)</td>
</tr>
<tr>
<td>Anticardiolipin (aCL) antibody-positive*</td>
<td>9 (26)</td>
</tr>
<tr>
<td>Lupus anticoagulant positive*</td>
<td>6 (18)</td>
</tr>
<tr>
<td>BILAG total score†</td>
<td>1 (0, 2)</td>
</tr>
<tr>
<td>SLEDAI-2K</td>
<td>2 (0, 2)</td>
</tr>
<tr>
<td>SDI</td>
<td>0 (0, 1)</td>
</tr>
<tr>
<td>Average daily corticosteroid dose (mg) (n=12)</td>
<td>8.75 (6.25, 11.25)</td>
</tr>
<tr>
<td>Current immunosuppressant use</td>
<td>15 (41.7)</td>
</tr>
<tr>
<td>Current antimalarial use</td>
<td>22 (61.1)</td>
</tr>
<tr>
<td>Biological medication</td>
<td>3 (8.3)</td>
</tr>
</tbody>
</table>

*At time of study.
†Score calculated as stated in Yee et al.2

### Table 2 Demographic, psychiatric, fatigue and biomarker characteristics across the participant groups

<table>
<thead>
<tr>
<th>Variable</th>
<th>SLE (n=36)</th>
<th>HC (n=30)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographic</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
<td>40 (32, 48.75)</td>
<td>32 (27, 46.5)</td>
<td>0.14</td>
</tr>
<tr>
<td>Gender (% female)</td>
<td>34 (94)</td>
<td>30 (100)</td>
<td>0.19</td>
</tr>
<tr>
<td>Handedness (% right-handed)</td>
<td>30 (83)</td>
<td>28 (93)</td>
<td>0.34</td>
</tr>
<tr>
<td>Years in education</td>
<td>16.11 (3.51)</td>
<td>17.97 (3.40)</td>
<td>0.034</td>
</tr>
<tr>
<td>WTAR (IQ)</td>
<td>102.5 (98.25, 108)</td>
<td>111 (105, 114)</td>
<td>0.001</td>
</tr>
<tr>
<td>Ethnic origin</td>
<td>P=0.132</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caucasian</td>
<td>24 (66.7)</td>
<td>24 (80)</td>
<td></td>
</tr>
<tr>
<td>Black Caribbean</td>
<td>4 (11.1)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Black African</td>
<td>3 (8.3)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Indian</td>
<td>1 (2.8)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Bangladesi</td>
<td>0</td>
<td>1 (3.3)</td>
<td></td>
</tr>
<tr>
<td>Chinese</td>
<td>1 (2.8)</td>
<td>1 (3.3)</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>3 (8.3)</td>
<td>4 (13.3)</td>
<td></td>
</tr>
<tr>
<td>Fatigue</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FSMC – Motor score</td>
<td>36 (22, 40.5)</td>
<td>114 (11.5, 18.5)</td>
<td>0.001</td>
</tr>
<tr>
<td>FSMC – Cognitive score</td>
<td>31 (22, 40)</td>
<td>114 (11.5, 18.5)</td>
<td>0.001</td>
</tr>
<tr>
<td>FSMC – total score</td>
<td>67.5 (44.75, 80.5)</td>
<td>27 (23, 37)</td>
<td>0.001</td>
</tr>
</tbody>
</table>
| Biomarkers of inflammation and endothelial activation

hsCRP (mg/L) $1.44 (0.66, 5.06)$ 0.88 (0.39, 1.39) $0.013$
IL-6 (ng/mL) $1.67 (0.50, 5.33)$ 0.50 (0.50, 1.32) $0.003$
VCAM-1 (ng/mL) $474.93 (194.30)$ 345.66 (53.79) $0.001$
VEGF (ng/mL) $66.04 (13.93, 139.60)$ 45.42 (6.04, 114.93) $0.027$
BLyS (ng/mL) $0.51 (0.35, 0.71)$ 0.34 (0.27, 0.39) $0.001$

Missing data: WTAR not included for 3 HCs, and 4 SLEs; these participants’ first language was not English and/or they had dyslexia, as such it was felt that the scale would not accurately measure IQ in these participants. MADRS=5 SLE, 2 HC; FSMC=2 SLE, 1 HC hsCRP, IL-6, VEGF, BLyS=2 HC, 2 SLE; VCAM1–2 SLE. P-values in bold are significant at $<0.05$. BDI-II, Becks Depression Inventory – II; BILAG, British Isles Lupus Assessment Group Index; C3, complement component 3; C4, complement component 4; SDI, The Systemic Lupus International Collaborating Clinics/American College of Rheumatology Damage Index; IgG ds-DNA, immunoglobulin G double-stranded DNA.

---

Table 3  Differences between the SLE and HC groups for each of the CANTAB outcome measures

<table>
<thead>
<tr>
<th>Variable*</th>
<th>Measurement</th>
<th>SLE, n=36</th>
<th>HC, n=30</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAL+ (visual memory and new learning)</td>
<td>Total errors (adjusted)</td>
<td>29.50 (19.00, 79.75)</td>
<td>24 (10.75, 48.75)</td>
<td>P=0.095</td>
</tr>
<tr>
<td>VRM (verbal memory)</td>
<td>Free recall – total correct</td>
<td>10 (8, 13)</td>
<td>10 (8.75, 14)</td>
<td>P=0.327</td>
</tr>
<tr>
<td></td>
<td>(Max.=18)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RVP (attention)</td>
<td>Total hits (Max.=27)</td>
<td>13 (12, 20)</td>
<td>20 (15.75, 22)</td>
<td>P=0.002</td>
</tr>
<tr>
<td>ERT (emotional processing)</td>
<td>Average percentage correct – total (%)</td>
<td>61.49 (8.85)</td>
<td>66.94 (9.36)</td>
<td>P=0.019</td>
</tr>
<tr>
<td></td>
<td>Overall mean response latency – total (ms)+</td>
<td>1626.10 (1411.71, 2274.22)</td>
<td>1343.15 (1152.27, 1744.23)</td>
<td>P=0.012</td>
</tr>
<tr>
<td>OTS+ (executive function)</td>
<td>Mean choices to correct</td>
<td>1.40 (1.27, 1.73)</td>
<td>1.33 (1.18, 1.62)</td>
<td>P=0.484</td>
</tr>
<tr>
<td>SWM+ (working memory)</td>
<td>Between errors</td>
<td>108.41 (57.96)</td>
<td>94.73 (52.36)</td>
<td>P=0.328</td>
</tr>
</tbody>
</table>

Missing data: VRM: 1 SLE, ERT: 1 SLE; RVP: 1 SLE; OTS: 1 SLE.
P-values in bold are significant at <0.05.
*Higher scores indicate better performance except where indicated with a “+”.
ERT, emotional recognition task; OTS, one touch stockings; Pal, paired associate learning; RVP, rapid information visual processing; SWM, spatial working memory; VRM, verbal recognition memory.

signal (figure 1). No significant differences between the groups were found for the positive effect of the 2-0back condition.

**Attention condition (0back-rest)**
Using ROI analysis, the SLE group had a greater decrease in signal in the lingual gyrus compared with the HC group (figure 2). No significant differences between the groups were found for the positive effect of the 0back-rest condition.

**FERT task**
Behaviourally, the SLE group was slower to correctly determine whether a face was female or male when displaying sadness (p=0.035) (see online supplementary table S6). In an ROI analysis, in the sadness-neutral condition, the positive effect of task for this condition showed that the SLE group had an increased BOLD response in frontal areas compared with the HC group (figure 3). There were no differences in the BOLD responses for the negative effect of task.

**Exploratory associations between SLE and cognitive function**
Improved performance on the attention task negatively correlated with the signal in a default mode network (DMN) region during the WM task (RVP associated with the RSTG-WM, r=-0.60, p=0.003). Better performance on the emotional processing task negatively correlated with the signal in a DMN area during a WM task (ERT average percentage correct associated with the RSTG-WM, r=-0.72, p<0.001). Also, the mean response latency for the identification of the emotions in the emotional processing task negatively correlated with the signal in a cognitive region.

Figure 1  Significantly different BOLD responses for the SLE and HC groups for the n-back task, 2-0back, negative effect of task. HC, healthy control; SLE, systemic lupus erythematosus.
during the WM task (ERT overall mean response latency associated with left caudate-WM, $r_p = -0.51$, $p = 0.013$), implying a greater response in the DMN when performing better and quicker on an emotional processing task.

Structurally, the enlarged PVS in the centrum semiovale (CSO-VRS) and basal ganglia (BG-VRS) were associated with years in education ($r_p = -0.48$, $p = 0.022$ and $r_p = -0.45$, $p = 0.035$, respectively). Neither was associated with vascular biomarkers including LAC or aCL positivity, or VCAM-1.

The DMN areas (left transverse temporal gyrus and right superior temporal gyrus), areas that are usually attenuated including LAC or aCL positivity, or VCAM-1. Neither was associated with vascular biomarkers respectively). Neither was associated with vascular biomarkers including LAC or aCL positivity, or VCAM-1.

The attention region of the n-back task, in the visual attention region of the lingual gyrus, positively correlated with the BILAG total score ($r_p = 0.45$, $p = 0.033$), IL-6 ($r_p = 0.44$, $p = 0.036$), current use of immunosuppressant ($r_p = 0.48$, $p = 0.019$) and anti-malarial medication ($r_p = 0.47$, $p = 0.028$) and negatively with BG-VRS score ($r_p = -0.46$, $p = 0.030$) suggesting that higher inflammatory disease activity increased responses in an attentional brain region during an attention task.

The BOLD response to sad faces from the FERT task in the left frontal cluster negatively correlated with the SDI score ($r_p = -0.57$, $p = 0.005$) and disease duration ($r_p = -0.43$, $0.47$).

**DISCUSSION**

We have identified structural, cognitive and fMRI differences in patients with SLE. While overall cognitive function was comparable between groups, the SLE group was less accurate on a test of sustained attention. Anatomically, we found increases in the PVS in the centrum semiovale in 43% of SLE participants and no controls. Using task-based fMRI, there was significant interference in emotional tasks and a reduced ability of patients with SLE to suppress the DMN during cognitive tasks.

Our data support previous work showing that attention is the most common cognitive problem in SLE. Such attention deficits can result in problems with other cognitive functions, such as WM although in this study we did not find any other non-emotional cognitive problem in the SLE group. A more detailed assessment of these relationships was limited due to time constraints with testing; however, follow-up studies focusing on these key inter-relationships are planned.

The 0-back-rest condition of the fMRI n-back task is a neuronal marker of sustained attention. Patients with SLE had a larger task-negative BOLD response in the lingual gyrus compared with the HC group. This region has been associated with visual attention, visual encoding/processing and WM. This may explain why our SLE group performed worse on the behavioural (CANTAB) attention task. The few studies published using fMRI in SLE have suggested that patients with SLE employ compensatory brain mechanisms to maintain cognitive function. Our cohort may have failed to recruit compensatory mechanisms on the challenging sustained attention task, resulting in performance deficits. Exploratory analyses also found that the response to attention in the lingual gyrus negatively correlated with an increase in PVS in the basal ganglia. Previous studies have found patients with SLE to have a greater number and larger PVS in the basal ganglia which did not reach significance in our study; we did however find differences between the HC and SLE in PVS in the centrum semiovale. PVS is an imaging marker for cerebral small vessel disease; however, we did not find any correlations between several serological markers of vascular disease (LAC, aCL antibodies and VCAM-1) and CSO-VRS, although we did exclude patients with severe vascular disease.

On the CANTAB battery, there were no differences between groups for the behavioural WM tasks. However, on fMRI patients with SLE had less task negative BOLD signals in the left transverse temporal gyrus and right superior temporal gyrus. These areas are part of the DMN, which is usually inactive during cognitive tasks and active during rest and internal processes, such as self-reflective processes and planning. The limited ability to reduce these signals in SLE implies an inability to inhibit self-reflective processes which can impede performance on cognitive tasks that do not usually have an emotional component, by allowing emotional interference from self-reflection and worries about task performance. In support of this, the FERT fMRI task found that patients with SLE had a greater response to viewing sad faces in frontal regions compared with the HC group. Such increased responses to sad expressions is also associated with depression. Similarly, patients with SLE were less accurate in correctly identifying emotions on CANTAB and showed evidence of reduced response latency implying a level of...
psychomotor slowing, both of which are associated with depression, and may contribute to some of the differences observed between groups. This is despite our groups scoring within normal ranges on the depression scales and that major depression was an exclusion criteria. We therefore cannot rule out the potential impact mood may have on cognitive performance in this SLE group even at the subtle end of the scale. Also, we recruited a low disease activity cohort and excluded NPSLE cases, so there may be different subtypes depending on CD severity.

We also noted differences in the caudate between the two groups. Patients with SLE had a larger task negative response compared with the HC group. The caudate, via the network linked to the dorsolateral prefrontal cortex, has been implicated in WM but usually as an area with an increased BOLD response during WM tasks. The attenuated response we found is in contrast to Mak et al who noted, using a different executive function task, an increased BOLD response in the caudate body from patients with SLE. It is therefore unclear if the differences we found are task specific and why our findings were in a task negative direction.

As noted, the SLE group had significantly higher scores on scales for depression and fatigue. While depression scores were still within the ‘normal range’ both motor and cognitive fatigue scores were higher in the SLE group. Both fatigue and mood can affect cognition and these symptoms are highly prevalent in SLE populations. In our study, mood and fatigue negatively impact neurocognition. It is increasingly recognised that inflammation and mood are closely interlinked and we found that VCAM-1, IL-6 and BILAG 2004 scores correlated with cognitive brain mechanisms, supporting the hypothesis that inflammation in SLE contributes to CD. We also noted associations with the SDI and disease duration, strengthening the suggestion that specific SLE factors directly impact on cognitive function over time.

We acknowledge some limitations to this study. For the non-fMRI analysis, the participant numbers are small; due to strict exclusion criteria and the use of fMRI as the main outcome. We also made no adjustments for multiple comparisons. However, many factors were closely correlated and in an exploratory study such as this, a Bonferroni correction would be too conservative. The HC group had a higher IQ and slightly more years in education than the SLE group. IQ can affect performance on cognitive tests but for many of the cognitive measures, no differences were seen so it is unlikely that this was the case. It was also impossible to recruit a SLE patient group on no medication. Patients on low dose psychoactive medications were included as well as those on corticosteroids. The correlations within the SLE group found no significant associations between corticosteroid dose and cognitive measures. Lastly, we chose to use the CANTAB battery as it is a sensitive measure of cognitive function that can assess changes over time and is easy to administer; however, some research has suggested that the tests measure overall cognition but cannot be divided into specific domains, such as executive function, so caution may be needed when interpreting individual test results.

In patients with SLE, we have noted impairments in sustained attention while other non-emotional cognitive functions remained unaffected. Poor attenuation of the DMN may contribute to CD in SLE, although prospective studies may be needed to confirm this, and our data suggest that in addition to mood and fatigue, inflammatory mechanisms and organ damage impact cognitive functioning in SLE.

Clinically, this study has implications when advising patients about CD in SLE. It has highlighted the multifaceted nature of CD in SLE and that future therapeutic approaches will need to be individually tailored to address the relevant drivers in individual patients.
Inflammatory mechanisms and organ damage impact cognitive functioning in people with Lupus.

**INTRODUCTION**

Systemic lupus erythematosus (often called Lupus or SLE) is an autoimmune disease. It typically starts in women between the ages of 15 and 45. Lupus causes immune cells in the body to become hyperactive and produce autoantibodies. An antibody is a protein that the immune system makes to attack foreign substances in the body, such as viruses or bacteria. In autoimmune disease, the body makes antibodies that attack its own tissues. These are called autoantibodies.

Lupus symptoms can vary from patient to patient. People with Lupus are often very tired (a symptom also called “fatigue”), have joint pain, and their skin may be sensitive to sunlight. Lupus can also cause cognitive problems with a person’s memory, concentration, attention and planning. People with the disease often refer to this as ‘brain fog’.

**WHAT DID THE AUTHORS HOPE TO FIND?**

The authors wanted to find out more about cognitive problems in people with Lupus.

**WHO WAS STUDIED?**

The study looked at 36 people with stable Lupus, and compared them to 30 healthy volunteers.

**HOW WAS THE STUDY CONDUCTED?**

All the people with Lupus were being treated in Manchester in the UK. The healthy volunteers were from the general population in the same area. Everyone had a one-off visit to the clinic, which lasted 4 hours. During this time, people in both groups told the researchers about their medical history, had blood samples taken, and completed questionnaires about their mood, fatigue, pain and sleep. They also did some puzzle games on a computer that tested their cognitive function. Finally, everyone had a special type of brain MRI scan (called a functional MRI, or fMRI), which allowed doctors to look at how areas of the brain responded when people were completing tasks to test their memory, attention and emotional processing.

**WHAT WERE THE MAIN FINDINGS OF THE REVIEW?**

Overall, the people with Lupus performed similarly to the healthy volunteers on all tasks except attention. However, the fMRI results showed that the people with Lupus had different brain responses while they were doing the tasks. This suggests that people with Lupus use different brain mechanisms to compensate for their disease. This allows them to maintain the same level of function as someone without the disease. This also means their brain has to work a bit harder to maintain normal cognitive function.

The authors also found subtle differences in the brain structure as well as differences in levels of mood, fatigue, and markers of inflammation between the people with Lupus and the healthy volunteers. These factors may all have an effect on cognitive function in Lupus.

**ARE THESE FINDINGS NEW?**

Yes and no! The use of this special kind of MRI in people with Lupus is relatively new, but there have already been some studies done with it that have suggested that people with Lupus use compensatory brain mechanisms to maintain cognitive function. So this part of the study supports previous findings.

However, this study is different from previous studies because it is bigger, and it looked at how someone performs on a cognitive task, what is happening in their brain whilst they are doing the task and if there are any structural issues in the brain. This study also excluded people with a form of Lupus called neuropsychiatric Lupus, where there is direct brain involvement. This is because the authors wanted to look at the milder cognitive problems reported by people with Lupus such as ‘brain fog’. Another new finding of this study is that...
Lupus-specific factors (such as inflammation and Lupus organ damage or disease duration) are associated with cognitive dysfunction as well as chronic disease factors such as fatigue, depression, pain, and medication. We know that people with chronic diseases are more likely to have cognitive problems, but these results suggest that it isn’t just the chronic disease factors that are affecting cognitive function, but that there is something specific to Lupus that is also having an effect.

WHAT ARE THE LIMITATIONS OF THE STUDY?
There are a few limitations because of the way the study was done. The strict exclusion criteria and the use of fMRI as the study measure meant that there were quite small numbers of people taking part.

There were also some slight differences between the two groups. Overall, the people in the healthy volunteer group had slightly higher IQs than the people with Lupus. IQ measures intelligence, which can affect cognitive function, but this difference does not seem to have affected the study. Certain medicines can also affect cognitive function, and some people in the Lupus group were taking medicine for their disease, but again the authors do not think this has affected the results.

WHAT DO THE AUTHORS PLAN ON DOING WITH THIS INFORMATION?
The authors hope to run another trial looking at treatments to try and improve cognitive function in people with Lupus. They are working on pilot studies to help get the design right. They will also share these results with other teams working in Lupus, as it may be important to consider these findings in other trials testing new medicines.

WHAT DOES THIS MEAN FOR ME?
If you have Lupus, this study should help raise awareness about the impact that the disease can have on your brain. You might find that although you can do the same tasks as your colleagues or friends, your brain gets tired more quickly. If you experience ‘brain fog’ it is worth bringing it up with your doctor, as it might be an important factor in deciding which treatment is best for you.

There are clinical trials going on in Lupus. If you are interested in being involved, you should speak to your doctor.

Disclaimer: This is a summary of a scientific article written by a medical professional (“the Original Article”). The Summary is written to assist non medically trained readers to understand general points of the Original Article. It is supplied “as is” without any warranty. You should note that the Original Article (and Summary) may not be fully relevant nor accurate as medical science is constantly changing and errors can occur. It is therefore very important that readers not rely on the content in the Summary and consult their medical professionals for all aspects of their health care and only rely on the Summary if directed to do so by their medical professional. Please view our full Website Terms and Conditions. http://www.bmj.com/company/legal-information/

Date prepared: June 2019

Summary based on research article published on: 12 April 2019


Copyright © 2019 BMJ Publishing Group Ltd & European League Against Rheumatism. Medical professionals may print copies for their and their patients and students non commercial use. Other individuals may print a single copy for their personal, non commercial use. For other uses please contact our Rights and Licensing Team.