Neuropsychiatric lupus erythematosus, cerebral infarctions, and anticardiolipin antibodies

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Abstract
Anticardiolipin antibody (aCL) has been associated with thromboembolic phenomena, including stroke, in certain patients with systemic lupus erythematosus (SLE); however, the relation between this antibody and the central nervous system manifestations of SLE is unknown. Serum samples and cerebrospinal fluid from five patients with SLE and acute central nervous system manifestations were assayed for the presence of aCL. Anticardiolipin antibody was identified in sera from four of the five patients but in none of the cerebrospinal fluid samples. Nuclear magnetic resonance imaging showed 'infarct-like' lesions in these four patients. This preliminary study suggests that a correlation between serum aCL and cerebral infarcts in central nervous system lupus may potentially exist. From this limited study it seems unlikely that aCL has a direct pathogenic role in the diffuse encephalopathy of acute central nervous system lupus.

Anticardiolipin antibodies (aCLs) are antiphospholipid antibodies with a related specificity to the lupus anticoagulant.1 2 Anticardiolipin antibody has been detected in patients with systemic lupus erythematosus (SLE)3-5 and other rheumatological and non-rheumatological entities.6-11 The presence of aCL is associated with thrombosis in patients with or without identifiable connective tissue disease.9 12 13 A hypercoagulable state promoted by aCL has been postulated, in which aCL interacts with endothelial surfaces and platelet membranes to reduce prostacyclin production and increase platelet adhesiveness.14 15 An association of aCL with cerebral infarction has been reported both in patients with SLE and in patients with other disorders.16-18

Central nervous system involvement in SLE is common19 and is manifested by headaches, psychiatric syndromes, organic brain syndrome, seizure disorder, cranial neuropathy, and movement disorder.20-23 Clinically, diffuse cerebritis (diffuse lupus encephalopathy) is the most common form of central nervous system lupus and is characterised by acute organic brain syndrome, psychosis, or seizures.24 Diffuse cerebritis is best defined as the presence of diffuse or shifting focal central nervous system manifestations not readily explained by the presence of infection, stroke, haemorrhage, steroid psychosis, or aseptic meningitis. Several reports have suggested that autoantibodies may play a part in the pathogenesis of central nervous system lupus25 26 and are most often associated with diffuse cerebritis.28 29 The presence of such antibodies would be manifested by the absence of acute infarction and by the presence of reversible oedema of neural tissue.30 Recently, aCLs have been shown to cross react with cephalin and sphingomyelin, which are components of neuronal tissue.31 The importance of anticardiolipin as an antineuronal antibody has not been studied, however.

In this study we assayed for aCLs in the sera and cerebrospinal fluid of five patients with neuropsychiatric SLE and attempted to relate the presence or absence of these antibodies to important clinical and laboratory measures of disease activity.

Patients and methods
PATIENTS
All five patients were under the care of the division of rheumatology and clinical immunology at the University of New Mexico School of Medicine. The diagnosis of SLE was established using American Rheumatism Association criteria.32 Serum and cerebrospinal fluid samples were obtained at admission. The diagnosis of diffuse encephalopathy and the presence of typical physical findings was confirmed by cranial magnetic resonance imaging (MRI) scan. Scans were obtained at the time of admission except where patients required ventilatory support on admission.

ANTICARDIOLIPIN ASSAY
Anticardiolipin antibody was measured by an enzyme linked immunosorbent assay (ELISA) similar to that described by Loizou et al.33 Immulon I microtiter ELISA plates (Dynatech) were coated with 30 μl (45 μg/ml) of cardiolipin (Sigma, St Louis, Mo). The plates were blocked for non-selective binding with a 1% gelatin solution. Serum and cerebrospinal fluid samples were diluted 1:100 in 10% fetal bovine serum; 200 μl of each sample was placed in duplicate wells and incubated for two hours. The plates were then washed with a phosphate buffered saline solution with 0.05% Tween and 10% fetal bovine serum and developed with a 1:1000 dilution of goat antihuman IgG or IgM conjugated to horseradish peroxidase and 2,2'-azino-di-(3-ethylbenzthiazoline) sulphonate. The optical density at 405 nm was read on a Dynatech microELISA reader. Known negative and positive control sera were used as standards to correct for plate to plate variability. Control serum samples from 543 normal blood donors were used to standardise the assay. A value at or
above the 98th centile was considered positive. In some cases serial titrations of sera were done and examined for the presence of aCL.

Magnetic resonance imaging showed a recent right occipital infarct. The patient improved with tapering of corticosteroids and discontinuance of antipsychotic drugs.

Case reports

Case 1
A 36 year old woman with a long history of SLE, nephritis, renal failure and chronic haemodialysis, positive antinuclear antibody test, lymphopenia, and arthritis was admitted with grand mal seizures and respiratory arrest. Lumbar puncture showed 36×10⁶ red blood cells/l and no white blood cells. A computed tomographic scan on admission showed mild ventricular enlargement and generalised atrophy. A clinical diagnosis of diffuse cerebritis was made and treatment with intravenous methylprednisolone was begun. Because of persistent need for continuing ventilatory support a cranial MRI scan was not obtained until eight days after admission. This showed punctate areas of abnormal increased intensity on intermediate and T2 weighted images in the subcortical and periventricular white matter, most marked in the left parietal region. There were patchy white matter changes in the subcortical white matter of the frontal and parietal lobes.

Case 2
A 25 year old woman with a one month history of SLE characterised by arthritis, positive antinuclear antibody test, positive anti-DNA antibody, and glomerulonephritis had been treated as an outpatient with prednisone for nephritis, but then became psychotic, necessitating large doses of corticosteroids and suppression of her psychosis with haloperidol and chlorpromazine. On admission, an electroencephalogram disclosed bilateral temporal slowing. The cerebrospinal fluid showed 155×10⁶ red blood cells/l, glucose 2·5 mmol/l, and protein 520 mg/l. C3 and C4 were normal.

Clinical and laboratory manifestations of patients with systemic lupus erythematosus and acute central nervous system lupus

<table>
<thead>
<tr>
<th>Patient No</th>
<th>Age</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>Clinical manifestations</th>
<th>Admission MRI*</th>
<th>Follow up MRI</th>
<th>Serum aCL*</th>
<th>CSF*</th>
<th>RPR*</th>
<th>PT*</th>
<th>PTT*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>36</td>
<td>F</td>
<td>Diffuse cerebritis</td>
<td>Seizures, coma</td>
<td>Diffuse and patchy white matter changes in left parietal and bilateral frontal and parietal lobes</td>
<td>None</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
<td>12</td>
<td>32</td>
</tr>
<tr>
<td>2</td>
<td>25</td>
<td>F</td>
<td>Diffuse cerebritis (possible steroid psychosis) and cerebral infarction</td>
<td>Psychosis</td>
<td>Right occipital infarct</td>
<td>None</td>
<td>(+)</td>
<td>(-)</td>
<td>(-)</td>
<td>12-1</td>
<td>25</td>
</tr>
<tr>
<td>3</td>
<td>61</td>
<td>F</td>
<td>Diffuse cerebritis syndrome and probable cerebral infarct</td>
<td>Seizures, organic brain syndrome</td>
<td>Multiple grey and white matter changes in cerebral hemispheres</td>
<td>Left pontine infarct, improvement in white matter changes</td>
<td>(+)</td>
<td>(-)</td>
<td>(-)</td>
<td>12-2</td>
<td>24</td>
</tr>
<tr>
<td>4</td>
<td>17</td>
<td>F</td>
<td>Diffuse cerebritis syndrome</td>
<td>Seizures, organic brain syndrome</td>
<td>Occipital, frontal, and left, parietal cortical lesions; bilateral cerebellar lesions</td>
<td>Persistent left cerebellar lesion</td>
<td>(+)</td>
<td>(-)</td>
<td>(-)</td>
<td>12-3</td>
<td>26</td>
</tr>
<tr>
<td>5</td>
<td>21</td>
<td>F</td>
<td>Acute stroke syndrome (cerebral infarct)</td>
<td>Left hemiparesis</td>
<td>Right internal capsule infarct, multiple punctate white matter changes in left frontal and parietal regions</td>
<td>Residual right internal capsule infarct only</td>
<td>(+)</td>
<td>(-)</td>
<td>(-)</td>
<td>11-4</td>
<td>27</td>
</tr>
</tbody>
</table>

*MRI=magnetic resonance imaging; aCL=anticardiolipin antibody; CSF=cerebrospinal fluid; RPR=rapid plasma reagin agglutination test; PT=prothrombin time; PTT=partial thromboplastin time.
cranial computed tomogram was normal. Cranial MRI showed extensive high intensity areas in cortex extending to the edge of the white matter, including both occipital and frontal lobes, and the left parietal lobe. Large discrete lesions were present in both cerebellar hemispheres. The patient improved after treatment with intravenous methylprednisolone. An MRI scan taken eight days after admission showed complete resolution of all cerebral lesions, with one persistent cerebellar lesion consistent with infarct.

**CASE 5**

A 21 year old woman with SLE characterised by serositis, leucopenia, Coombs' positive anaemia, malar rash, and arthritis was admitted owing to acute onset of left upper arm weakness. Physical examination showed a left hemiplegia and malar rash. The cerebrospinal fluid showed no evidence of infection or haemorrhage. Cranial MRI showed acute infarction in the posterior limb of the right internal capsule with multiple punctate white matter changes in the left frontal and parietal regions. A subsequent follow up MRI scan after discharge showed only a persistent right internal capsule infarct.

**Results**

Four of the five patients with acute central nervous system lupus were positive for serum aCL (table). In none of the patients was aCL detected in the cerebrospinal fluid. Three of the patients with a diagnosis of diffuse cerebritis were positive for serum aCL, but again none of these had detectable cerebrospinal fluid levels. The isotypes and titres of the sera of the four patients were as follows: patient 2, IgG=1/400, IgM=1/400; patient 3, IgG=1/100, IgM >1/1600; patient 4, IgG >1/1600, IgM >1/1600; patient 5, IgG >1/1600, IgM >1/1600.

Figures 1 and 2 show representative lesions seen with MRI of diffuse lupus encephalopathy and cerebral infarction.

**Discussion**

In none of our five cases of acute central nervous system lupus were aCLs detectable in the cerebrospinal fluid. Four of the patients (Nos 1–4) had symptoms not readily explainable on the basis of infarct and therefore were felt to have diffuse cerebritis. A diagnosis of steroid psychosis in patient 2 was also a possibility. In patients 3 and 4 diffuse cerebritis was suggested by the transient nature or marked improvement of MRI lesions with corticosteroids. Both these patients (Nos 3 and 4) had serum aCL but...
neither had detectable cerebrospinal fluid levels. Of note, the serum and cerebrospinal fluid from patient 1 were negative for aCL. A possible association of serum aCL with cerebral thrombosis was confirmed in our patients. Four of our five patients (Nos 2–5) had persistent lesions on MRI, consistent with infarct. It should be emphasised that many of these ‘infarct-like’ lesions are not visible on conventional computed tomographic scans but such lesions are seen by MRI in lupus patients; thus infarct-like lesions may be a more accurate designation at this time. All four of these patients had serum aCL. In all patients both IgG and IgM were found and the titres were similar for both isotypes.

The absence of aCL in cerebrospinal fluid makes a role for aCL as a direct antineuronal antibody in acute cerebritis unlikely. Of note, even in the patients in whom disruption of the blood-brain barrier might have been expected (Nos 2–5) with MRI documented infarct there was no detectable aCL, even though those four patients had serum aCL. To show central nervous system production of an antibody rather than leakage based on a disrupted blood-brain barrier it is necessary to show blood-brain barrier integrity by determining Q albumin and IgG index. The complete absence of this antibody in cerebrospinal fluid in our cases of definite diffuse cerebritis (Nos 3 and 4) made this analysis unnecessary, however.

Anticardiolipin may be a marker antibody for patients at risk for acute central nervous system lupus. Its absence in cerebrospinal fluid in our limited series seems to indicate that it does not play a part as a direct acting antineuronal antibody. Serum aCL was seen in all patients with infarcts, which suggests its presence may correlate strongly with lupus patients with cerebral infarcts. Cerebrospinal fluid determinations in a large number of patients would be useful to confirm our findings.

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