**CONCISE REPORT**

Disseminated intravascular coagulation in catastrophic antiphospholipid syndrome: clinical and haematological characteristics of 23 patients

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**Background:** Disseminated intravascular coagulation (DIC) is an acquired syndrome characterised by formation of microthrombi and fibrin deposition in the microvasculature. The catastrophic antiphospholipid syndrome (APS) is characterised by multiorgan thrombosis, mainly involving small vessels. A broad spectrum of disorders may develop DIC features; however, the catastrophic APS has not previously been recognised as a cause of DIC.

**Objective:** To analyse the clinical and laboratory characteristics of catastrophic APS patients with DIC features.

**Methods:** The website based international registry of patients with catastrophic APS (CAPS registry) (http://www.med.ub.es/MIMMUN/FORUM/CAPS.HTM) was analysed and the cases with DIC features selected.

**Results:** In 173 patients with catastrophic APS, 23 (13%) were found with DIC features. The clinical and immunological characteristics were similar in catastrophic APS patients with and without DIC features; a significant difference was found only in the prevalence of thrombocytopenia (100% in patients with DIC features vs 59% in those without DIC features).

**Conclusions:** DIC features are not rare in catastrophic APS, supporting the need for systematic screening of antiphospholipid antibodies in all patients with DIC features without precipitating factors. The presence of DIC features in the context of an APS makes it imperative to rule out the catastrophic variant of this syndrome.

Disseminated intravascular coagulation (DIC) is an acquired syndrome characterised by the widespread activation of coagulation with occlusion of small and medium sized vessels. This condition may compromise the blood supply to organs and contribute to multiorgan failure. It is not a disease entity in itself but always occurs as a complication of an underlying disorder, the most common being infection, severe trauma, malignancy, and obstetric complications.²,³

The “catastrophic” variant of the antiphospholipid syndrome (APS) is an accelerated form of this syndrome resulting in multiorgan failure because of multiple small vessel occlusions.²,³ As with DIC, most of the catastrophic APS episodes are preceded by a precipitating event, such as infection, surgery or trauma, obstetric complications, and malignancies.²,³ Furthermore, laboratory features of DIC were reported in 19–28% of the largest catastrophic APS series.⁷,⁸

Our objective in the present study was to analyse the clinical and laboratory characteristics of catastrophic APS patients with DIC features to determine whether these patients form a special subset within the catastrophic APS population.

**METHODS**

We analysed the web site based international registry of patients with catastrophic APS (the CAPS registry; http://www.med.ub.es/MIMMUN/FORUM/CAPS.HTM). This registry was created in 2000 by the European Forum on Antiphospholipid Antibodies and compiles all the published reports as well as newly diagnosed cases from all over the world. Up to October 2003 it included 220 patients with this condition.⁹ We selected those patients who had some of the laboratory features of DIC (raised fibrin related markers, decreased fibrinogen concentrations, or both). Isolated thrombocytopenia and prolonged prothrombin time were not considered to be selection criteria as they can be manifestations of the APS.

According to the International Scientific Subcommittee for DIC, we calculated the DIC score as follows:

- platelets: >100 x 10⁹/l = 0; <100 x 10⁹/l = 1; <50 x 10⁹/l = 2;
- raised fibrin related markers (fibrin degradation products and D-dimers): no increase = 0; moderate increase = 2; marked increase = 3;
- prolonged prothrombin time: <3 seconds = 0, 3–6 seconds = 1; >6 seconds = 2;
- decreased fibrinogen: >1.0 g/l = 0; <1.0 g/l = 1.

Overt DIC was diagnosed when the total score was ≥5; a score <5 was considered suggestive of DIC.¹⁰

Fisher’s exact test (bilateral) was employed for the statistical analysis, using the SPSS 10.0 statistical program.

**RESULTS**

**General characteristics**

Of the 220 patients included in the CAPS registry, information on DIC features was not available in 34 cases and there were incomplete data for DIC scoring in 10 further patients. This left a total of 176 patients available for analysis. Of these, 23 (13%) had DIC features associated with catastrophic APS: 17 (74%) female and six (30%) male, mean (SD) age 39 (13) years (range 11 to 60). Ten (43%) suffered from primary APS, nine (39%) had systemic lupus erythematosus (SLE), three (13%) had lupus-like disease, and one (4%) had polychondritis.

**Abbreviations:** aCL, anticardiolipin antibodies; APS, antiphospholipid syndrome; DIC, disseminated intravascular coagulation
Clinical presentation and precipitating factors

Intra-abdominal involvement was identified in all 23 patients, mainly consisting of renal (78%), hepatic (48%), gastrointestinal (39%), splenic (17%), pancreatic (9%), and adrenal (9%) manifestations.

Pulmonary complications were reported in 16 patients (70%), mainly acute respiratory distress syndrome (ARDS) and confirmed pulmonary embolism, but occasionally intra-alveolar haemorrhage. Eleven patients (48%) had cardiac involvement, mainly cardiac failure and confirmed myocardial infarction or valve lesions. Fifteen patients (65%) had evidence of cerebrovascular complications, mainly encephalopathy and cerebrovascular accidents, but occasionally seizures, headache, or silent brain infarcts. Deep venous thrombosis was present in two patients (9%) and peripheral arterial occlusive disease in one (4%).

Skin manifestations were also frequent (78%) and consisted of livedo reticularis, ulcers, necrotic lesions, digital gangrene, purpura, microthrombosis of small vessels, splinter haemorrhages, and multiple ecchymoses.

Other lesions occasionally encountered were bone marrow necrosis, mononeuritis multiplex, and retinal involvement.

The most common precipitating conditions were infections (seven cases) and surgical procedures (four cases). Other cases were attributed to drug use and anticoagulation withdrawal (one each).

APS related laboratory findings

Thrombocytopenia (platelet count <150 x 10^9/l) was reported in all 23 patients and evidence of haemolytic anaemia in nine (41%), accompanied by schistocytes in five (23%). The IgG isotype of anticardiolipin antibodies (aCL) was detected in 19 patients (83%), IgM aCL in eight (38%), and lupus anticoagulant in 18 (82%).

DIC features

A platelet count of <100 x 10^9/l was reported in 20 patients (87%) (in one additional case, the count was reported only non-specifically, as “low platelet count”). Increased fibrin degradation products were reported in all 19 patients in whom they were recorded, and positivity for D-dimers in eight of nine cases (89%). A prolonged prothrombin time was reported in six of 10 patients (60%) and decreased fibrinogen levels were present in five of 13 cases (39%) (in two cases they were reported non-specifically as “low levels”). Table 1 shows in detail each case of catastrophic APS with DIC features. Eleven patients (48%) had a DIC score of 5 or above (compatible with overt DIC). The remaining 12 patients (52%) had a DIC score of 3 or 4 (suggestive of DIC).

Treatment and outcome

Most patients received a combination of treatments. Anticoagulation were used in 20 patients (87%), steroids in 19 (83%), plasma exchange in nine (39%), cyclophosphamide in eight (35%), intravenous gamma globulin in five (22%), and splenectomy in two (9%). Other treatments used were prednisolone and antithrombin concentrate (one case each).

Recovery occurred in 61% of catastrophic APS patients with DIC features and in 58% of those without DIC features (NS). Assessing the use of single treatments, recovery of DIC patients occurred in 58% of those treated with anticoagulants v 67% of those not treated with anticoagulants (NS); in 58% of those treated with steroids v 67% of those not (NS); in 38% of those treated with cyclophosphamide v 71% of those not (NS); in 50% of those treated with plasmapheresis v 64% of those not (NS); and in 83% of those treated with intravenous gamma globulin v 50% of those not (NS).

Comparison of catastrophic APS patients with and without DIC

The profiles of demographic characteristics (sex distribution and mean age), clinical features (severe organ involvement), and immunological findings (lupus anticoagulant, IgG aCL, and IgM aCL) were similar. Significant differences were found only in the prevalence of thrombocytopenia (100% in the DIC group v 59% in the catastrophic APS patients without

### Table 1: Laboratory variables according the scoring system for DIC

<table>
<thead>
<tr>
<th>Case</th>
<th>Platelets (&lt;10^9/l)</th>
<th>Score FDPs</th>
<th>D-dimers</th>
<th>Score PT (s)</th>
<th>Score Fibrinogen (g/l)</th>
<th>Score Total</th>
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<td>2</td>
<td>57</td>
<td>1</td>
<td>Increased</td>
<td>3</td>
<td>12.7</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>39</td>
<td>2</td>
<td>NR</td>
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<tr>
<td>4</td>
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<td>12</td>
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<tr>
<td>5</td>
<td>67</td>
<td>1</td>
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<td>2</td>
<td>12.7</td>
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<td>2</td>
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<td>23</td>
<td>35</td>
<td>2</td>
<td>Increased</td>
<td>2</td>
<td>Normal</td>
<td>0</td>
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</table>

Platelet count: >100 x 10^9/l = 0; <100 x 10^9/l = 1; <50 x 10^9/l = 2.

Raised fibrin related markers (FDPs and D-dimers): no increase = 0; moderate increase (raised but less than twice the normal level) = 2; marked increase (more than twice the normal level) = 3.

Prothrombin time: <3 seconds = 0; 3–6 seconds = 1; >6 seconds = 2.

Fibrinogen concentration: >1.0 g/l = 0; <1.0 g/l = 1.

Total score: >5, compatible with DIC; <5, suggestive of DIC.
DIC, p<0.001). Other characteristics typically encountered in other states causing DIC—such as renal failure, skin involvement, or ARDS—were not more frequent in catastrophic APS patients with DIC features than in those without.

**DISCUSSION**

We observed laboratory features of DIC in at least 13% of patients diagnosed as having the catastrophic variant of APS. However, it should be borne in mind that there were incomplete data for DIC scoring in 10 reported cases. Thus, under ideal circumstances where all the data were available, the incidence might turn out to be higher.

The pathophysiology of DIC and catastrophic APS is poorly understood, but the two conditions probably share some pathogenic mechanisms and triggering factors. In DIC, enhanced fibrin formation is caused by tissue factor mediated thrombin generation and simultaneous dysfunction of inhibitory mechanisms, such as the antithrombin system and the protein C and protein S system. In addition, fibrin removal is impaired because of fibrinolytic system depression, mainly caused by high circulating levels of plasminogen activator inhibitor type 1 (PAI-1). Conversely, catastrophic APS is associated with endothelial cell activation as a result of antigen–antibody reactions on the surface of endothelial cells or monocytes. Furthermore, inhibition of both protein C activation and the function of activated protein C have been observed in association with APS. Finally, increased plasma concentrations of PAI-1 characterise the hypofibrinolytic state in APS.

A link between DIC and catastrophic APS can be assumed from the original description of DIC by McKay in 1965. He described a 38 year old woman with SLE with some features suggestive of APS, such as chorea, mitral valve disease, and spontaneous abortion. A few days after an elective cholecystectomy she developed a sudden episode of multiorgan failure characterised by fever, severe congestive heart failure, ARDS, renal failure, and somnolence accompanied by features of DIC (low fibrinogen, thrombocytopenia, and a prolonged prothrombin time). Her clinical status deteriorated and she died three weeks after the surgical procedure. Necropsy showed microvascular thrombosis of the heart, adrenal glands, lungs, and bone marrow, in addition to a non-bacterial thrombotic endocarditis, all of these being typical features of catastrophic APS. At that time, however, APS was an unknown entity.

Infections associated with DIC were the most common precipitating factors in catastrophic APS in our series of patients. Molecular “mimicry” has been proposed as one of the major mechanisms responsible for the development of catastrophic APS following infections. Thus an infectious aetiology for the APS, especially its catastrophic variant, should perhaps be considered more frequently and appropriate antibiotic therapy instituted.

Another aspect to bear in mind is the differential diagnosis between DIC and disorders presenting with thrombotic microangiopathic haemolytic anaemia (TMHA). The clinical picture of DIC, TMHA, and APS may overlap and, if they coexist in the same patient, the diagnosis may be difficult at the time of presentation (table 2).

In conclusion, DIC features are not rare in catastrophic APS. This would support the need for systematic screening of antiphospholipid antibodies in all patients with DIC without precipitating factors. In addition, the presence of DIC in the context of an APS makes it mandatory to rule out the catastrophic variant of this syndrome.

**APPENDIX**

**THE CATASTROPHIC ANTIPHOSPHOLIPID SYNDROME REGISTRY PROJECT GROUP**

The members of the Catastrophic Antiphospholipid Syndrome Registry Project Group who contributed to the study are listed in the appendix.

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*The members of the Catastrophic Antiphospholipid Syndrome Registry Project Group who contributed to the study are listed in the appendix.*

**Table 2**

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Catastrophic APS</th>
<th>DIC</th>
<th>TMHA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haemorrhagic manifestations</td>
<td>±</td>
<td>+</td>
<td>±</td>
</tr>
<tr>
<td>Anaemia</td>
<td>±</td>
<td>±</td>
<td>+</td>
</tr>
<tr>
<td>Schistocytes</td>
<td>±</td>
<td>+</td>
<td>±</td>
</tr>
<tr>
<td>Thrombocytopenia</td>
<td>++</td>
<td>++</td>
<td>+++</td>
</tr>
<tr>
<td>Prolonged prothrombin time</td>
<td>−</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Prolonged activated partial thromboplastin time</td>
<td>±</td>
<td>±</td>
<td>±</td>
</tr>
<tr>
<td>Fibrinogen degradation products</td>
<td>±</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Antiphospholipid antibodies</td>
<td>++</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Plasma ADAMTS-13 activity</td>
<td>Normal or reduced</td>
<td>Moderately reduced</td>
<td>Absent or severely reduced</td>
</tr>
</tbody>
</table>

*In cases of familial thrombotic purpura (TTP), acquired idiopathic TIP, and pregnancy related TIP.* ADAMTS-13, von Willebrand factor cleaving protease; APS, antiphospholipid syndrome; DIC, disseminated intravascular coagulation; TMHA, thrombotic microangiopathic haemolytic anaemia.

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