Fibrin(ogen)olysis in polymyalgia rheumatica and temporal arteritis: preliminary findings on association with disease activity

RAFAEL G. GRAU, STUART S. KASSAN, JOHN J. FRANKS, HERBERT KAPLAN, STROTHEN H. WALKER, AND ENG M. TAN*

From the University of Colorado School of Medicine and the Denver VA Medical Center, Denver, CO, USA

SUMMARY Postulating an increased production of fibrin(ogen)olytic degradation products (FDP) and an abnormality of fibrinogen metabolism in polymyalgia rheumatica (PMR) and temporal arteritis (TA), we studied 16 PMR/TA patients and 10 control subjects using sensitive radioimmunoassay for a specific type of FDP, namely, fibrinogen-related D-antigen. Median serum D-antigen levels were increased five-fold in those 11 PMR/TA patients who were untreated compared with control subjects. In the five PMR/TA patients who were treated with prednisone the median D-antigen levels were not significantly different from those of the healthy controls. D-antigen concentration correlated significantly (r=0.83) with ESR in the seven untreated PMR patients. In PMR patients prednisone therapy was followed by a reduction of serum D-antigen levels.

Polymyalgia rheumatica (PMR) and temporal arteritis (TA) are interrelated clinical syndromes which are often accompanied by giant cell arteritis and a raised erythrocyte sedimentation rate (ESR). The affected vessels often show thrombosis and luminal narrowing. These findings in conjunction with elevated fibrinogen levels, suggest that abnormalities of fibrinogen metabolism may exist, either of pathogenetic significance or as a reflection of the underlying process.

In health most if not all fibrinogen is catabolised by unknown processes, with minimal conversion by thrombin to fibrin or by plasmin to fibrinogen degradation products. In certain pathological states conversion of fibrinogen to fibrin and proteolysis of fibrin and fibrinogen to fibrin(ogen)olitic degradation products (FDP) occur at a sufficiently rapid rate for specific FDP to be identified in the serum or plasma. A variety of FDP can be isolated. Among them are two D-antigens, D₁ (a specific proteolytic product of fibrinogen or fibrin monomer) and D dimer (a specific cross-linked fibrin product). We report the use of a sensitive fibrinogen-related D-antigen radioimmunoassay capable of estimating serum levels of FDP containing D₁ and D dimer in a group of PMR or TA patients, either untreated or treated with prednisone, and in normal control subjects.

Materials and methods

The patients and control subjects were seen in the Arthritis Clinic at the University of Colorado Health Sciences Center. Included were 11 patients with PMR, five with TA, and 10 non-rheumatic control subjects, four of whom had mild osteoarthritis and two mild hypertension.

All PMR and TA patients fulfilled previously outlined criteria. No PMR patient had historical or clinical evidence of temporal artery inflammation.

Blood samples for fibrinogen-related D antigen measurements were collected in tubes containing 3670 NF units of soybean trypsin inhibitor and 20 NIH units of bovine thrombin. Each sample was incubated for two hours at 37°C and centrifuged. Serum was removed and stored at 4°C for one week or less before assay. Fibrinogen-related D antigen was measured by radioimmunoassay by a double antibody technique described previously.

Statistical methods. Groups were compared by the
Kruskal-Wallis test. If the result of this test was significant, pairwise comparisons were made by the rank-sum test. Results were confirmed by a one-way analysis of variance on a log transformation of the data, followed by Duncan's k ratio procedure for multiple comparisons. Association between D antigen concentration and ESR was determined by the Spearman rank correlation coefficient as well as by unranked regression, with comparable results.

Results

Sixteen patients, 11 PMR and 5 TA (12 females and 4 males), with a median age of 69 (range, 50 to 86), were studied. The median age of the control subjects was 63 (range, 50 to 74) with 9 females and 1 male (Table 1). The median disease duration was 3 months (range 1 to 24 months). Three TA patients had the diagnosis confirmed by biopsy, while two were diagnosed clinically. Four TA patients had a mild preceding or concurrent myalgic syndrome. Five patients, 4 PMR and 1 TA, were receiving prednisone when first studied (8-40 mg/day). The median ESR of the untreated PMR (n=7) and TA (n=4) patients was 86 mm/h (range, 63-127) and 81 mm/h (range, 58-119) respectively. These values differed significantly (p<0.02) from the median ESR of treated PMR/TA patients, which was 19 mm/h (range 5 to 41), and from the control subjects' median, which was 20 mm/h (range 6 to 40).

Fig. 1 shows logarithmic plots of individual measurements of D antigen concentration in the various study groups: untreated PMR or TA patients, treated PMR or TA patients, and control subjects. D antigen levels in seven untreated PMR patients (median 3.54 µg/ml) and four untreated TA patients (median 1.28 µg/ml) were elevated compared with controls (median 0.49 µg/ml) (p<0.01). D-antigen levels were higher in untreated PMR than in untreated TA (p<0.03). (SI conversion: µg/ml=mg/l.)

Median concentrations (µg/ml serum) for the combined 11 untreated PMR and TA patients was 2.45 and for the five treated PMR and TA patients was 0.77, compared with 0.49 for the controls. Differences between untreated patients and either treated patients or control subjects were significant (p<0.01), but treated patients did not differ significantly from controls.

Table 1  Clinical and therapeutic data on 16 patients with polymyalgia rheumatica and temporal arteritis

<table>
<thead>
<tr>
<th>Diagnosis and treatment studies</th>
<th>Number</th>
<th>Female/male ratio</th>
<th>Median age</th>
<th>Median ESR</th>
<th>Median prednisone dose (mg 4 times daily)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMR treated</td>
<td>4</td>
<td>1/3</td>
<td>72</td>
<td>23</td>
<td>11</td>
</tr>
<tr>
<td>untreated</td>
<td>7</td>
<td>6/1</td>
<td>68</td>
<td>86</td>
<td>0</td>
</tr>
<tr>
<td>TA treated</td>
<td>1</td>
<td>1/0</td>
<td>73</td>
<td>5</td>
<td>40</td>
</tr>
<tr>
<td>untreated</td>
<td>4</td>
<td>4/0</td>
<td>62</td>
<td>81</td>
<td>0</td>
</tr>
<tr>
<td>Total patients</td>
<td>16</td>
<td>12/4</td>
<td>69</td>
<td>64</td>
<td>20</td>
</tr>
<tr>
<td>Control subjects</td>
<td>10</td>
<td>9/1</td>
<td>63</td>
<td>63</td>
<td>20</td>
</tr>
</tbody>
</table>

PMR=Polymyalgia rheumatica. TA=Temporal arteritis.
Fibrinogen/fibrin deposits have been found in the capillary walls of deltoid muscle biopsies in PMR patients. In addition thrombi have been noted in temporal artery biopsies with proved giant cell arteritis. It is conceivable, therefore, that PMR and TA are associated with local fibrin formation and lysis within affected vessels, perhaps because of release of activators from the altered vascular endothelium. Indeed, in-vitro cytotoxicity of sera from PMR and TA patients to human endothelial cells has been described. The raised levels of FDP may indicate enhanced thrombin or plasmin (or, less likely, other proteolytic enzyme) activity. Whether the elevated D antigen concentrations represent a primary pathogenetic process or merely reflect vascular inflammation and subsequent fibrin formation and lysis is unclear.

Three PMR patients were studied before and after treatment with prednisone for one to three months. In each of these patients treatment resulted in a drop in D antigen concentration toward normal, the initial median level being 2.26 μg/ml and the subsequent one 1.21 μg/ml.

Significant correlation was observed between D antigen levels and ESR in the seven untreated PMR patients (r=0.83, p<0.01). No such correlation was found in the four untreated TA patients (Fig. 2).

Discussion

The fibrinogen/fibrin-related D-antigen measured by radioimmunoassay in our patients with PMR or TA cannot definitely be identified as fibrin fragment D dimer, because anti-D dimer rabbit globulin reacts with the D domain of fibrinogen and of fibrinogen fragments X, Y, and Z as well. However, antibody affinity for these latter substances is only about 10% that of an equivalent amount of D dimer. In addition the assay dilution curves of patient (and control) sera were parallel to the D dimer standard curves. Parallelism of assay dilution curves is a necessary, though not a sufficient, condition to establish identity. The evidence suggests that most of the FDP found with our assay in patients with PMR or TA is fibrin fragment D dimer, either free or complexed with other fibrinogen moieties.

D antigen concentrations in our healthy subjects and patients varied from about 0.4 to 10 μg/ml. Our results correspond closely with values seen in control and postoperative subjects (0.86 to 10.1 μg/ml) using a different fibrinogen fragment D assay. These values are lower than those obtained by other tests, in which FDP levels ranging from 12 to over 1000 μg/ml were found in a variety of acute and subacute conditions. However, those patients presented with a manifest defibrination syndrome and might be expected to produce larger amounts of FDP than patients with PMR or TA.

While fibrinogen levels are increased in untreated PMR or TA patients, it seems unlikely that their high D dimer concentrations merely reflect the fibrinogen increment. In fact the converse may be operating. Rats injected with homologous D1 or isolated rat livers perfused with blood from traumatized rats (which contain high levels of D antigens) respond by increasing their rates of fibrinogen synthesis.

Fibrinogen/fibrin deposits have been found in the capillary walls of deltoid muscle biopsies in PMR patients. In addition thrombi have been noted in temporal artery biopsies with proved giant cell arteritis. It is conceivable, therefore, that PMR and TA are associated with local fibrin formation and lysis within affected vessels, perhaps because of release of activators from the altered vascular endothelium. Indeed, in-vitro cytotoxicity of sera from PMR and TA patients to human endothelial cells has been described. The raised levels of FDP may indicate enhanced thrombin or plasmin (or, less likely, other proteolytic enzyme) activity. Whether the elevated D antigen concentrations represent a primary pathogenetic process or merely reflect vascular inflammation and subsequent fibrin formation and lysis is unclear.

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References


