Characterisation of anticytoplasmic antibodies and their clinical associations

Wei-Howe Koh, Juliet Dunphy, Jean Whyte, Jonathan Dixey, Neil John McHugh

Abstract

Objectives—To characterise the cytoplasmic staining patterns identified by indirect immunofluorescence (IF) of human epithelial (HEP-2) cells and the antigens recognised using additional serological techniques. To define the disease associations of anticytoplasmic antibodies.

Methods—Sera from 1173 patients were screened for cytoplasmic IF staining on HEP-2 cells and the patterns characterised. The presence of antimitochondrial antibodies (AMA) was evaluated by a sensitive anti-pyruvate dehydrogenase complex enzyme linked immunosorbent assay (ELISA) (IgG) and by immunoblotting. Detection of antibodies to extractable nuclear antigens (ENA) was performed by double immunodiffusion and the presence of anti-ribosomal P antibodies was determined by immunoblotting.

Results—Cytoplasmic IF staining was demonstrated in 75 sera (6.4%). Six different patterns were recognised: coarse granular filamentous speckles (AMA, n=9); condensed large speckles (anti-golgi apparatus antibodies, n=3); cytoskeletal (n=9); centriolar (n=4); diffuse coarse speckles (n=33); and fine speckles (n=17). Of the nine sera with an AMA pattern, the presence of these antibodies was confirmed in seven by the ELISA (n=6) and on immunoblotting (n=7). One of the seven patients had primary biliary cirrhosis, and two had scleroderma. Two patients with anti-golgi antibodies had rheumatoid arthritis and two with ant centriolar antibodies had scleroderma. Of 33 sera that had cytoplasmic staining and were ANA negative, three were positive for anti-Ro and two were positive for anti-Jo-1 antibodies.

Conclusions—In general, defined cytoplasmic IF patterns have no specific disease associations. However, the finding of cytoplasmic fluorescence should not be ignored, as it may indicate the presence of antibodies to ENA in the absence of nuclear staining.

IMMUNOFLUORESCENCE

The sera were diluted 1/40 in phosphate buffered saline (PBS) and incubated on HEP-2 cells (Biodiagnostic Ltd, Worcs, UK) for 30 minutes at room temperature. The slides were then washed in PBS and incubated with fluorescein isothiocyanate (FITC) conjugated polyvalent goat anti-human immunoglobulin (1/150 dilution) for 30 minutes at room temperature. The observed cytoplasmic IF patterns were then characterised.
IMMUNODIFFUSION
Precipitating antibodies against extractable nuclear antigens (ENA) were detected by means of double immunodiffusion in Ouchterlony plates containing 1% agarose. Saline extracts of calf thymus prepared as previously described, fresh extracts of rabbit thymus powder (Pel-Freeze, Rogers, AR, USA) and commercially produced Ro/La extract (Bradshaw Biologicals Ltd, Leics, UK) were used as sources of antigen. Antibody specificity was defined using a variety of prototype sera, including those with antibodies to U1RNP, Sm, Ro(SS-A), La(SS-B) and Jo-1 antigens.

PDHC ELISA AND IMMUNOBLOTTING
We tested all sera which produced cytoplasmic fluorescence for antimitochondrial antibodies (AMA) by a sensitive anti-pyruvate dehydrogenase complex (PDHC) enzyme linked immunosorbent assay (ELISA) as previously described. Those sera that had coarse granular filamentous speckled IF patterns characteristic of AMA, or increased AMA on the ELISA were further evaluated by immunoblotting which has been previously described. Purified human brain mitochondria was used as a source of antigen for immunoblotting. All sera that produced a cytoplasmic fluorescence were also tested for the recognition of ribosomal P antigen by immunoblotting using a whole cell extract prepared from K562 cells.

Results
IMMUNOFLUORESCENCE
Of 1173 sera tested, 75 (6.4%) had cytoplasmic fluorescence staining. Figure 1A–F shows the six patterns that were identified. The commonest pattern was the diffuse coarse speckled (A), which occurred in 44% (n = 33) of the sera that tested positive for cytoplasmic IF (table). Cytoplasmic fluorescence patterns that were infrequently encountered were those characteristic for the anti-golgi antibodies (C) (n = 3), and anticientriolar antibodies (B) (n = 4). Altogether, 33 of the 75 sera were ANA negative.

IMMUNODIFFUSION
Autoantibodies to ENA were detected in 10 of the sera with anticytoplasmic antibodies, five of which had a diffuse coarse speckled pattern. Five had anti-Ro (SS-A) antibodies, four had anti-Jo-1 antibodies and another had anti-U1 ribonucleoprotein (RNP) antibodies. Of the five sera that had anti-Ro antibodies, one also had anti-La antibodies and another had both AMA and anti-La antibodies. Five of the patients who had autoantibodies to ENA (anti-Ro, n = 3; anti-Jo-1, n = 2) were ANA negative.

PDHC ELISA
Of the nine sera with AMA IF pattern, six had antibodies detected by the sensitive anti-PDHC ELISA (IgG). Figure 2 shows the results of the anti-PDHC ELISA of the 75 test sera and of 38 healthy controls. A cut off of 2 SD from the mean of the healthy control was used to determine a positive result. Three patients who did not have an AMA IF pattern were tested positive for antibodies to PDHC by the ELISA.

IMMUNOBLOTTING
Seven of the nine sera that produced an AMA IF recognised both a 70 kDa and a 52 kDa polypeptide on immunoblotting (fig 3). Of these seven patients, six had antibodies to PDHC detected on the ELISA. None of the three patients who had a negative AMA IF but were positive for antibodies to PDHC by ELISA recognised a 70 kDa mitochondrial antigen on immunoblotting. No sera had antibodies against the cytoplasmic antigen ribosomal P detected by immunoblotting.

CLINICAL ASSOCIATIONS
The table lists the various cytoplasmic IF patterns identified and the associated clinical diagnoses. Amongst those patients who had a coarse granular filamentous speckled pattern characteristic of the presence of AMA, one patient had primary biliary cirrhosis (PBC) and two had limited systemic sclerosis. Another patient with generalised osteoarthritis had increased serum alkaline phosphatase without other clinical evidence of PBC or scleroderma.

Two of the three patients with anti-golgi antibodies had seropositive RA, whilst two patients with anticientriolar antibodies had scleroderma. The diffuse coarse or fine speckled and cytoskeletal patterns were identified in patients with a variety of rheumatic conditions (table) and no disease specificities were noted.

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### Cytoplasmic immunofluorescence patterns and associated diagnoses

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<th>Pattern</th>
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<th>Diagnosis</th>
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<tr>
<td></td>
<td></td>
<td>RA</td>
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<td></td>
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<td>Polymyositis</td>
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<td></td>
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<td>PMR</td>
<td>2</td>
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<td></td>
<td></td>
<td>Limited SS</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Diffuse SS</td>
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</tr>
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<td></td>
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<td>Others</td>
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<tr>
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<td>5</td>
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<td></td>
<td></td>
<td>Sjögren’s disease</td>
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<td></td>
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<td>Diffuse SS</td>
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<td></td>
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<td>Others</td>
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<td>2</td>
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<tr>
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<td>SLE</td>
<td>2</td>
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<tr>
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<td>Diffuse SS</td>
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<td>Dermatomyositis</td>
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<td></td>
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<td>PBC</td>
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<tr>
<td>(anti-golgi antibodies)</td>
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<td>Others</td>
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</table>

SLE = Systemic lupus erythematosus; RA = rheumatoid arthritis; PMR = polymyalgia rheumatica; SS = systemic sclerosis; PBC = primary biliary cirrhosis; Others include undifferentiated connective tissue disease, seronegative arthritis, and vasculitis.
Characterisation of anticytoplasmic antibodies and their clinical associations

Discussion

Our results show that cytoplasmic fluorescence is not an infrequent finding when ANA is tested for by indirect IF of HEp-2 cells. About one in 20 sera (6.4%) produced a cytoplasmic IF stain. We have identified six cytoplasmic IF patterns, including some well defined patterns like those characteristic of AMA, anti-golgi, anticentriolar, and anticytoskeletal antibodies. The most common pattern found was a diffuse coarse speckled pattern, which occurred in almost half the sera which had a cytoplasmic stain. Diffuse fine speckled and cytoskeletal patterns occurred less frequently. These patterns were produced by sera from patients with a spectrum of rheumatic diseases.

By using an indirect IF method with PTK 2 cells fixed with paraformaldehyde and digitonin as substrate, Senécal et al found a high frequency of antibodies to intermediate filaments (one of three classes of cytoskeleton complex network of filaments) in patients with PM/DM, RA, systemic sclerosis, and SLE. In contrast to Senécal et al, who were unable to demonstrate the presence of these antibodies using HEp-2 cells as substrate, we were able to detect anticytoskeletal antibodies in patients with a similar profile of autoimmune diseases (table). Similarly, Osung et al using HEp-2 cells as substrate, detected antibodies against intermediate filaments of the cytoskeleton in patients with RA.

Figure 1  Cytoplasmic fluorescence patterns on human epithelial (HEp-2) cells. A: Diffuse cytoplasmic coarse speckled pattern. B: Anticentriolar antibodies pattern. C: Condensed large speckles (anti-golgi antibodies). D: Diffuse cytoplasmic fine speckled pattern. E: Cytoskeletal pattern. F: Coarse granular filamentous speckles (antimitochondrial antibodies). Bars represent 100 μm.
In the nine sera which showed a characteristic AMA pattern, the presence of the antibodies was detected by the sensitive anti-PDHC ELISA in six and was confirmed by immunoblotting in seven using human brain mitochondrial antigen. Apart from the 70 kDa polypeptide which is the major mitochondrial antigen in PBC, the seven sera that were positive for AMA on immunoblotting also recognised a 52 kDa polypeptide which could correspond to the protein X of the PDHC, or the branched chain keto acid dehydrogenase complex (BCODH). Of the nine sera with an AMA IF pattern, two were negative for antibodies to PDHC on the ELISA and did not recognise either the 72 kDa or the 52 kDa polypeptide on immunoblotting. Although the two sera demonstrated a characteristic AMA IF pattern, the nature of the cytoplasmic antigens was not determined.

The presence of autoantibodies to mitochondrial antigens has been described in the majority of patients with PBC and up to 25% of patients with systemic sclerosis. Patients with other autoimmune diseases may also have AMA. In our study, of those patients who had AMA IF staining patterns, two had limited systemic sclerosis and one had PBC. The remainder of the patients with AMA IF pattern had other rheumatic diseases, including seronegative polyarthritis and SLE, but no overt liver disease; nevertheless, the patients may have had subclinical liver disease.

A condensed large speckled cytoplasmic IF pattern characteristic for anti-golgi antibodies was not a pattern commonly detected. Only three of the 1173 sera tested exhibited this pattern—a prevalence rate similar to that reported in other studies. The antibody was first reported by Rodriguez et al in a patient with Sjögren’s syndrome and lymphoma. Subsequently, the antibody was found to be associated with other rheumatic diseases, including RA. Of our three patients with the antibody, two had seropositive RA, supporting the report by Hong et al concerning the association of anti-golgi antibodies with RA.

The IF pattern typical of anticentriolar antibodies was detected in two of our patients with scleroderma—an association described previously. However, the pattern is not specific to the disease, as it was also seen in a patient with polymyalgia rheumatica and another with mixed connective tissue disease, both of whom did not have features of scleroderma.

We were able to detect autoantibodies to ENA (anti-Ro, anti-La, anti-Jo-1 and anti-U1 RNP) in some of the sera with cytoplasmic fluorescence. Ro and La antigens shuttle between cellular compartments and may be found in the cytoplasm in association with RNA complexes. These antigens have been termed soluble cytoplasmic RNP (scRNP) and may have been responsible for some of the cytoplasmic fluorescence detected. The antigen Jo-1, which is a cytoplasmic enzyme histidyl tRNA synthetase, can also give rise to a cytoplasmic IF stain on HEp-2 cells. We found that the majority of sera with antibodies to ENA tended to be associated with a diffuse coarse cytoplasmic IF pattern. Of interest, five of the sera with these antibodies were negative for ANA on IF. Thus it is important not to disregard cytoplasmic fluorescence in the absence of nuclear staining, as this may indicate the presence of antibodies to ENA.

Antibodies to the cytoplasmic antigen ribosomal P protein occur predominantly in patients with SLE and it is thought that they often account for cytoplasmic IF. Using the specific Western blot technique, we did not detect anti-ribosomal P antibodies in any of the sera.
our patients, some of whom had SLE. This may have resulted from their relatively low prevalence which varies between 5 and 20% of patients with SLE. Furthermore, the antibody titre may vary with time and is higher in those patients with active disease—a feature not necessarily present in our patients.

In conclusion, although, in general, anti-cytoplasmic antibodies are not disease specific, certain characteristic patterns on IF are of more diagnostic help than others. Furthermore, a negative ANA result may sometimes be misleading if the cytoplasmic IF is ignored.

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