MPO-ANCA may produce a combination of PR-ANCA and atypical cytoplasmic ANCA indirect immunofluorescent patterns on certain ethanol fixed neutrophil substrates

The P-ANCA pattern is defined as perinuclear indirect immunofluorescent (IIF) staining on ethanol fixed normal human neutrophils. This pattern is an artefact of ethanol fixation, dependent on the redistribution of certain cationic neutrophil granule proteins (such as myeloperoxidase (MPO), lactoferrin, and lysozyme) around the negatively charged nuclear membrane. However, certain MPO-ANCA can produce cytoplasmic rather than perinuclear IIF staining, possibly related to a subpopulation of epitopes on MPO that do not redistribute with ethanol fixation. We now report that MPO-ANCA positive sera may produce a combination of P-ANCA and atypical cytoplasmic ANCA IIF patterns on certain ethanol fixed neutrophil substrates, potentially leading to interpretative and diagnostic difficulties.

Sera from six patients with biopsy confirmed microscopic polyangiitis (at different stages of disease activity) were selected because of initial difficulties in the interpretation of their IIF patterns on ethanol fixed neutrophil slides from Inova Diagnostics (San Diego, California, USA). All six sera were MPO-ANCA positive and proteinase 3-ANCA (PR3-ANCA) negative by the corresponding ORGenTec (Mainz, Germany) enzyme linked immunosorbent assay (ELISA). PR3-ANCA positive serum from a patient with biopsy confirmed Wegener’s granulomatosis was also tested. To establish whether other ANCA antigen specificities were present, all sera were tested on the ORGenTec ANCA Combi-kit® ELISA containing proteinase-3, MPO, lactoferrin, elastase, cathespine G, lysozyme, and bacterial/permelliberty increasing protein (BPI). IIF was then repeated on all sera on two separate occasions using in house (kindly supplied by the Division of Immunology, Royal Brisbane Hospital) and two commercial (Inova Diagnostics (different batch) and Medical and Biological Laboratories (MBL, Nagoya, Japan)) ethanol fixed neutrophil slides. The IIF staining patterns and end point titres were determined by consensus.

Table 1 summarised the results. In four of the six sera, no reactivity other than MPO-ANCA was detected using the ANCA Combi-kit ELISA. Of the other two sera, one also contained lactoferrin-ANCA and the other lysozyme-ANCA. Nevertheless, in addition to P-ANCA staining, atypical cytoplasmic staining was consistently produced by all six MPO-ANCA sera on the Inova slides, but not on the MBL or in house slides. These findings were reproducible on different batches of neutrophil slides from the former manufacturer.

Our small study demonstrates that sera containing MPO-ANCA may produce a combination of PR-ANCA and atypical cytoplastic ANCA IIF patterns on certain ethanol fixed neutrophil substrates. The recent International Consensus Statement recommends that such combined patterns be reported as "atypical ANCA". Because atypical ANCA are not strongly associated with microscopic polyangiitis or Wegener’s granulomatosis, an atypical ANCA IIF report on these sera could potentially erroneously lead the requesting clinician away from the correct diagnosis. However, in all six sera, the positive MPO-ANCA ELISA result would hopefully redirect attention towards a possible diagnosis of systemic necrotising vasculitis.

We have subsequently found that these combined IIF patterns do not occur with all MPO-ANCA positive sera on the Inova slides, and thus speculate that the phenomemon might be caused by factors in the ethanol fixation conditions of these slides resulting in the differential redistribution of different MPO epitopes. Therefore, we recommend that laboratories using this brand (and possibly other commercial brands) of ethanol fixed neutrophil slides be aware of this phenomenon, and consider repeating any sera producing such combined “atypical ANCA” IIF patterns on alternative ethanol fixed neutrophil substrates to clarify their “true” IIF pattern. Furthermore, antigen specific ELISA testing for MPO-ANCA and PR3-ANCA should also be performed on all such sera because combining IIF and ELISA in ANCA testing improves overall diagnostic specificity/precision value compared with using either test alone.


Table 1 MPO-ANCA and PR3-ANCA ELISA, ANCA Combi-kit® ELISA, and ANCA IIF results

<table>
<thead>
<tr>
<th>Sera</th>
<th>MPO-ANCA and PR3-ANCA IgG ELISA (U/ml)</th>
<th>ANCA Combi-kit IgG ELISA (OD ratio)</th>
<th>Inova Diagnostics</th>
<th>MBL</th>
<th>In house cytoplasm</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>MPO positive (58)</td>
<td>MPO (6.4)</td>
<td>P (1/160)</td>
<td>P (1/160)</td>
<td>P (1/160)</td>
</tr>
<tr>
<td>2</td>
<td>PR3 negative</td>
<td>MPO (2.9)</td>
<td>P (1/160)</td>
<td>P (1/160)</td>
<td>P (1/160)</td>
</tr>
<tr>
<td>3</td>
<td>MPO positive (8)</td>
<td>MPO (1.6)</td>
<td>AOGP (1/10)</td>
<td>P (1/40)</td>
<td>P (1/40)</td>
</tr>
<tr>
<td>4</td>
<td>PR3 negative</td>
<td>MPO (1.3)</td>
<td>AOGP (1/10)</td>
<td>P (1/40)</td>
<td>P (1/40)</td>
</tr>
<tr>
<td>5</td>
<td>MPO positive (&gt;100)</td>
<td>MPO (9.1)</td>
<td>AOGP (1/40)</td>
<td>P (1/160)</td>
<td>P (1/160)</td>
</tr>
<tr>
<td>6</td>
<td>PR3 negative</td>
<td>MPO (5.2)</td>
<td>AOGP (1/10)</td>
<td>P (1/160)</td>
<td>P (1/160)</td>
</tr>
<tr>
<td>7</td>
<td>MPO positive (53)</td>
<td>PR3 (6.44)</td>
<td>C (1/40)</td>
<td>C (1/40)</td>
<td>C (1/40)</td>
</tr>
</tbody>
</table>

ORGenTec MPO-ANCA and PR3-ANCA IgG ELISA: positive, >5 U/ml; negative, ≤5 U/ml.

ORGenTec ANCA Combi-kit® IgG ELISA OD ratio: positive, ≥1; negative, <1 (only positive results shown).

IIF staining pattern: C, classic granular cytoplasmic IIF staining with central/interlobular accentuation; P, perinuclear.

Sera 1-6 were from patients with biopsy confirmed microscopic polyangiitis. Serum 7 was from a patient with biopsy confirmed Wegener’s granulomatosis.

ANCA, antineutrophil cytoplasmic antibody; ELISA, enzyme linked immunosorbent assay; FITC, fluorescein isothiocyanate; IIF, indirect immunofluorescence; MPO, myeloperoxidase; OD, optical density; PR3, proteinase 3.

High prevalence of serum markers of coeliac disease in patients with chronic fatigue syndrome

There has been recent interest in the possibility that undiagnosed coeliac disease (CD) might be the cause of diverse clinical symptoms, most particularly “tired all the time”. A recent study reported a prevalence of three in 100 cases in a primary care environment in which samples were taken from patients with a range of symptoms and signs. The second most frequent symptom reported by the endomysial antibody (EMA) positive patients was “being tired all the time”. We decided to examine the prevalence of EMA in patients attending our tertiary referral centre with the diagnosis of chronic fatigue syndrome (CFS).

We tested serum from 100 consecutive patients (47 men, 53 women; median age, 40 years; range, 18–57) referred to our specialist clinic and satisfying the standard CDC criteria for a diagnosis of CFS, and from 100 healthy control subjects (45 men, 55 women; median age, 40 years; range, 18–68) who were blood donors at the South East Thames Blood Transfusion Service. The CFS samples had been stored as part of other studies, and were analysed retrospectively. EMA of the IgA class were detected by indirect immunofluorescence (IF) using cryostat sections of distal primate oesophagus as substrate (Binding Site, Birmingham, UK). Positive samples were confirmed using an enzyme linked immunosorbent assay (ELISA) for the detection of antitransglutaminase antibodies (Menarini Diagnostics, Wokingham, UK).
tissue transglutaminase being the autoantigen responsible for the IF pattern of EMA. To exclude selective IgA deficiency, serum IgA concentrations were measured by laser nephelometry using specific antisera according to the manufacturer’s instructions (Behring, Duesseldorf, Germany).

Two of the 100 CFS samples were positive for EMA using IF, and this was confirmed by ELISA, but none of the 100 control samples was positive. None of the subjects had selective IgA deficiency. Mean (SD) serum IgA concentrations among patients with CFS were 2.1 g/litre (0.98). Neither of the positive cases, both women aged 27 and 28, had reported symptoms typical of CD, although one had a history of constipation. Routine blood tests including serum proteins and full blood count were normal, and both had been seen by consultant physicians before referral. Both had histories of hypothyroidism, were taking long term thyroxine, and were currently euthyroid. Before the diagnosis of CD was made retrospectively, both had received cognitive behaviour therapy (CBT), a standard treatment for CFS. In both cases, CBT led to a substantial improvement in the quality of life and physical activity, but neither patient was symptom free at the end of treatment or at six months follow up. In both cases, CD was subsequently confirmed on jejunal biopsy after the retrospective identification.

However, there is now evidence from primary care of a surprisingly high frequency of unsuspected positive EMA tests in people with non-specific symptoms and a suggestion that a higher index of suspicion is needed when assessing such patients. We now extend that observation to our CFS clinic. Indeed, given our prevalence of 2%, and the fact that there is a treatment for CD, we now suggest that screening for CD should be added to the relatively short list of mandatory investigations in suspected cases of CFS.

Correction


C Visser (Department of Cardiology, Free University Hospital, Amsterdam, The Netherlands) was mistakenly omitted from the list of authors of this paper. The journal apologises for any inconvenience that this may have caused.

Calendar of events

Diagnostic Pathology of Breast Disease
23–27 April 2001, Hammersmith Hospital (Imperial School of Medicine), London, UK

Further details: Wolfson Conference Centre, Hammersmith Hospital, Du Cane Road, London W12 ONN, UK. (Tel +44 020 8383 3171/3227/3245; fax +44 020 8383 2428; email wcc@ic.ac.uk)

Gynecologic and Obstetric Pathology
26–29 April 2001, Fairmont Copley Plaza, Boston, Massachusetts, USA

Further details: Department of Continuing Education, Harvard Medical School, 25 Shattuck Street, Boston, MA 02115, USA. (Tel +1 617 432 1525; fax +1 617 432 1562; email hms-cme@hms.harvard.edu)

BSCC London Spring Tutorial: Lung and Pleural Cavitary Fluid Cytology
27 April 2001, Guy’s Hospital, London, UK

Further details: BSCC Office, PO Box 352, Uxbridge UB10 9TX, UK. (Tel +44 01895 274 020; fax +44 01895 274 080; email lesley.couch@psilink.co.uk)

International Consultation on the Diagnosis of Noninvasive Urothelial Neoplasms
11–12 May 2001, University of Ancona School of Medicine, Torrette, Ancona, Italy

Further details: R Montironi, Ancona Italy (email r.montironi@popsci.unian.it), DG Bostwick, Richmond, VA, USA (email bostwick@bostwicklaboratories.com), P-F Bassi, Padua, Italy (email bassipf@ux1.unipd.it), M Droller, New York, USA (email michael_droller@smtplink.mssm.edu), or D Waters, Seattle, WA, USA (email waters@vet.vet.purdue.edu)

Human Adverse Drug Reactions
30 May 2001, Royal College of Pathologists, London, UK

Further details: Michelle Casey, Academic Activities Coordinator, 2 Carlton House Terrace, London SW1Y 5AF, UK. (Tel +44 020 7451 6700; fax +44 020 7451 6701; www.rcpath.org)

Professional Standards of Pathologists in a Modern NHS Pathology Service
7 June 2001, Royal College of Pathologists, London, UK

Further details: Michelle Casey, Academic Activities Coordinator, 2 Carlton House Terrace, London SW1Y 5AF, UK. (Tel +44 020 7451 6700; fax +44 020 7451 6701; www.rcpath.org)

Infectious Hazards of Donated Organs
28 June 2001, Royal College of Pathologists, London, UK

Further details: Michelle Casey, Academic Activities Coordinator, 2 Carlton House Terrace, London SW1Y 5AF, UK. (Tel +44 020 7451 6700; fax +44 020 7451 6701; www.rcpath.org)

Recent Advances in Genetics
5 July 2001, Royal College of Pathologists, London, UK

Further details: Michelle Casey, Academic Activities Coordinator, 2 Carlton House Terrace, London SW1Y 5AF, UK. (Tel +44 020 7451 6700; fax +44 020 7451 6701; www.rcpath.org)

BSCC Annual Scientific Meeting
9–11 September 2001, Majestic Hotel, Harrogate, UK

Further details: BSCC Office, PO Box 352, Uxbridge UB10 9TX, UK. (Tel +44 01895 274020; fax +44 01895 274080; email lesley.couch@psilink.co.uk)
High prevalence of serum markers of coeliac disease in patients with chronic fatigue syndrome
A Skowera, M Peakman, A Cleare, E Davies, A Deale and S Wessely

*J Clin Pathol* 2001 54: 335-336
doi: 10.1136/jcp.54.4.335-a

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