Antiphospholipid antibodies associated with alcoholic liver disease specifically recognise oxidised phospholipids

R Rolla, D Vay, E Mottaran, M Parodi, M Vidali, M Sartori, C Rigamonti, G Bellomo, E Albano

Abstract

Background—Circulating antiphospholipid antibodies (aPL) are often detected in patients with alcoholic liver disease (ALD) but little is known about the causes of their formation.

Aims—We have evaluated whether ethanol mediated oxidative injury might promote the development of aPL in ALD.

Patients and methods—IgG against β2 glycoprotein 1 (β2-GP1), cardiolipin, and human serum albumin (HSA) complexed with either oxidised arachidonic acid (HSA-APP) or malondialdehyde (HSA-MDA) were assayed by ELISA in heavy drinkers with or without ALD and in healthy subjects.

Results—Circulating IgG recognising cardiolipin were significantly higher in ALD patients than in controls. However, anti-cardiolipin reactivity of ALD sera was only evident using, as the antigen, oxidised cardiolipin, but not oxidation protected cardiolipin. In ALD patients, individual values of IgG antioxidised cardiolipin were associated with the titres of IgG antioxidised cardiolipin but not oxidation protected cardiolipin. In ALD patients, individual values of IgG antioxidised cardiolipin were associated with the titres of antibodies against HSA-MDA and HSA-APP (r=0.68 and 0.72, respectively; p<0.0001) used as markers of oxidative stress. ALD patients also displayed increased levels of antibodies against phospholipid binding protein β2-GP1, and individual reactivity towards oxidised cardiolipin and β2-GP1 were highly correlated (r=0.85; p<0.0001). IgG binding to oxidised cardiolipin, HSA-MDA, and HSA-APP was also significantly higher in β2-GP1 positive than in β2-GP1 negative sera. However, preadsorption of β2-GP1 positive sera on β2-GP1 coated ELISA plates reduced reactivity to oxidised cardiolipin by 80%, without affecting that to HSA-APP or HSA-MDA.

Conclusions—Ethanol induced oxidative injury is associated with the development of antibodies targeting complexes between oxidised cardiolipin and β2-GP1. These antibodies might account for high aPL titres observed in patients with severe ALD.


Keywords: oxidative stress; lipid peroxidation; β2 glycoprotein 1; ethanol; autoantibodies

Antiphospholipid antibodies (aPL) belong to a heterogeneous group of autoantibodies with apparent specificity for negatively charged phospholipids.1-3 The presence of these antibodies characterises primary antiphospholipid syndrome, or Hughes’ syndrome, consisting of recurrent vascular thrombosis, fetal losses, and thrombocytopenia.4 aPL and antiphospholipid syndrome are often associated with rheumatic and autoimmune diseases, such as systemic lupus erythematosus, rheumatoid arthritis, and systemic sclerosis.3-4 However, aPL are also present in patients with chronic infections (syphilis, malaria, human immunodeficiency virus 1), lymphoproliferative diseases, and sickle cell anaemia.1-4

The molecular targets of aPL have not yet been characterised in detail. aPL are usually identified by their ability to react with solid phase bound cardiolipin but other anionic phospholipids—that is, phosphatidylserine and phosphatidyethanolamine—can be recognised also.1-3 Studies by McNeil and colleagues5 and Galli and colleagues6 have shown that aPL binding to phospholipids is largely mediated by β2 glycoprotein 1 (β2-GP1) or apolipoprotein H, a plasmatic 50 kDa glycoprotein which readily binds to phospholipids as well as to other negatively charged substances.1 Conformational changes occurring in β2-GP1 following interaction with phospholipids render this protein antigenic to aPL. Indeed, β2-GP1 can be recognised by aPL also in the absence of phospholipids when attached to irradiated polystyrene or nitrocellulose membranes that are rich in negative charges.8 None the less, phospholipid complexes with prothrombin, protein C, and protein S can also be targets of aPL and this may contribute to the anticoagulant properties ascribed to aPL.4 Recently, Hörkkö et al have reported that epitopes originating during cardiolipin oxidation are recognised by aPL from women with primary antiphospholipid syndrome.9 They also suggested that the antigenic determinants might result from the reaction of β2-GP1 with products derived from the peroxidative breakdown of cardiolipin unsaturated fatty acids.9,10

Recent studies have shown that aPL can be detected in patients with chronic liver disease.
particularly alcoholic liver disease (ALD) and hepatitis C.11–16 High aPL titres are frequent in patients with alcoholic hepatitis or cirrhosis and the presence of aPL in these patients seems to reflect disease progression, showing significant correlation with disease severity.12 However, aPL can also be detected in alcoholics without liver damage.12 17 18 Furthermore, aPL are often associated with venous and arterial thrombosis: the presence of aPL in patients with chronic liver disease does not seem to be associated with thrombotic complications.19

Clinical and experimental evidence indicates that ethanol intoxication promotes free radical formation and oxidative liver injury.19 In particular, increased levels of lipid peroxidation products and protein carbonyls can be demonstrated in liver biopsies or in the sera obtained from patients with ALD.19–23 Furthermore, the formation of hydroxethyl free radicals during ethanol metabolism can also be documented in patients with alcohol abuse.24

The aim of this work was to evaluate whether the development of aPL in patients with ALD might reflect ethanol mediated oxidative injury.

**Patients and methods**

**PATIENTS AND CONTROL SUBJECT RECRUITMENT**

We enrolled 36 patients (27 men, nine women; mean age 51 years (range 31–73)) with alcoholic cirrhosis with or without hepatitis (ALD). The diagnosis was based on clinical, ecograpical, and laboratory criteria. Liver biopsy was available for 12 patients and showed the classical feature of micronodular cirrhosis. In three patients hepatocyte ballooning degeneration with Mallory’s bodies and inflammatory infiltrates were also noted. Evaluation of the clinical severity of liver injury according to Maddrey’s DF index or the Child-Turcotte classification25 26 revealed that 72% of patients in the ALD group had moderate or severe liver damage (DF values above 60) and 58% belonged to Child group B or C. Fifteen heavy drinkers (11 men, four women; mean age 44 years (range 25–66)) with normal liver blood tests and no clinical or ecograpical evidence of liver disease were also investigated. All patients with alcohol abuse had an estimated mean daily ethanol intake over the previous 12 months above 100 g (table 1) and none had a previous history of co-exposure to drugs or toxins. Patients with alcohol abuse were negative for serum markers of hepatitis B virus or for the presence of antibodies against hepatitis C virus (HCV), measured by a second generation enzyme linked immunosorbent assay (Abbott Laboratories, Chicago, Illinois, USA).

A group of 40 patients (29 men, 11 women; mean age 54 years (range 32–76)) suffering from HCV or hepatitis B virus chronic hepatitis or cirrhosis and without alcohol abuse were also investigated. In this group, 62% of patients were abstinent and the remaining had reported alcohol intake ranging from 10 to 50 g ethanol/day. The severity of liver injury in these patients was comparable with that in patients with alcoholic liver damage. Aspartate aminotransferase, alanine aminotransferase, gamma glutamyltransferase, bilirubin, and prothrombin time were assayed in all patients using routine laboratory procedures. The biological characteristics of the patients studied are reported in table 1.

Healthy controls were recruited among blood donors or university staff and consisted of 42 subjects (28 men, 14 women; mean age 47 years (range 33–66)). Alcohol intake among controls was below 20 g ethanol/day for women and was 40–60 g/day for men. At the time of blood testing, all subjects had abstained from alcohol for at least 48 hours. All subjects gave informed consent to the analysis and the study was planned according to the guidelines of the university ethics committee. Blood samples (5 ml) were taken after an overnight fast and serum was used for ELISA tests.

**ANTIPHOSPHOLIPID ANTIBODY DETERMINATION**

**TOWARDS OXIDISED PHOSPHOLIPIDS**

Phospholipid coated ELISA plates were prepared according to a standardised method proposed by Harris.27 Briefly, 30 µl of cardiolipin solution in ethanol (50 µg/ml) were applied to each well and ethanol evaporated overnight at 4°C in air. After two washes with phosphate buffered saline (PBS), 0.3 ml of coating buffer containing 3% bovine serum albumin (BSA) in PBS, pH 7.4, were added and the plates were further incubated for one hour at 37°C to block non-specific binding sites. The coated wells were washed three times with PBS. To avoid cardiolipin auto-oxidation in some preparations, 0.1 mmol/l diphenylphenylendiamine (DPPD) was added to cardiolipin ethanol solution and coating of the plates was carried out in sealed plastic bags filled with nitrogen. Cardiolipin coated plates were used immediately after preparation. For the assay, sera from patients and controls were diluted 1:50

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Alcohol consumption (g/day)</th>
<th>AST (U/l)</th>
<th>ALT (U/l)</th>
<th>GGT (U/l)</th>
<th>Bilirubin (µmol/l)</th>
<th>PT (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcoholic cirrhosis</td>
<td>36</td>
<td>146 (67)</td>
<td>74 (52)</td>
<td>50 (36)</td>
<td>143 (132)</td>
<td>1.68 (1.5)</td>
<td>72 (14)</td>
</tr>
<tr>
<td>Non-alcoholic cirrhosis</td>
<td>40</td>
<td>20 (14)</td>
<td>61 (36)</td>
<td>83 (58)</td>
<td>51 (24)</td>
<td>1.33 (0.6)</td>
<td>54 (4)</td>
</tr>
<tr>
<td>Heavy drinkers</td>
<td>15</td>
<td>190 (61)</td>
<td>28 (13)</td>
<td>22 (11)</td>
<td>37 (19)</td>
<td>0.62 (0.3)</td>
<td>101 (3)</td>
</tr>
</tbody>
</table>

AST, aspartate aminotransferase; ALT, alanine aminotransferase; GGT, gamma glutamyltransferase; PT, prothrombin time.

Values are mean (SD).
with the coating buffer and 0.2 ml aliquots were added in duplicate to the appropriate wells and incubated for one hour at 37°C. The plates were then washed three times with PBS 0.25% and further incubated with peroxidase linked goat antimouse IgG (dilution 1:6000) for 60 minutes at 37°C. Antibody binding was revealed by addition of 0.15 ml of a reaction mixture containing 0.4 mg/ml of L-phenylendiamine, 0.4 µl/ml hydrogen peroxide (30%), 5.1 mg/ml citric acid, and 6.1 mg/ml anhydrous NaH₂PO₄, pH 5.0. After 15 minutes the reaction was stopped by adding 50 µl of 2 N H₂SO₄ and absorbance was measured at 490 nm using a Bio-Rad microplate reader (Bio-Rad Laboratories Inc., Hercules, California, USA). The average optical densities (od) in duplicated wells were subtracted from non-specific background for each serum sample obtained by performing the test in plates coated with ethanol but lacking the phospholipid.

ASSESSMENT OF PHOSPHOLIPID OXIDATION
The occurrence of phospholipid peroxidation under the conditions used for ELISA plate coating was monitored by measuring conjugated diene absorbance. Briefly, 0.5 mg/ml of cardiolipin solution were placed in glass test tubes and allowed to evaporate from the solvent by overnight incubation at 4°C in air. Cardiolipin was then resolubilised in 2 ml hexane and conjugated diene absorbance was immediately measured at 234 nm using a Beckman DU650 UV visible spectrophotometer. Freshly prepared cardiolipin solution in hexane was used as reference.

PREPARATION OF PROTEIN ADDUCTS WITH LIPID PEROXIDATION PRODUCTS
Adducts between human serum albumin (HSA) and malondialdehyde (MDA) were prepared by reacting for two hours at 37°C, 1 mg/ml HSA with 50 mmol/l MDA, obtained by acid hydrolysis of malondialdehyde-bis-dimethylacetal. Unbound aldehyde was removed by overnight dialysis at 4°C against PBS, pH 7.4. The presence of MDA adducts was evaluated by measuring fluorescence intensity at 399/471 nm exc/em wavelength pairs according to Cominacini and colleagues.²⁷ HSA complexed with reactive products of fatty acid oxidation was generated by thermal auto-oxidation of arachidonic acid (HSA-APP) and linoleic acid (HSA-LPP) according to Palinski and colleagues.²⁸ Briefly, 10 µg arachidonic acid or linoleic acid were transferred in a glass vial open to air and kept at 37°C for 72 hours. The yellow-brown reaction products were dissolved in 50 µl methanol and suspended by vortexing in 1 ml PBS containing 10 µM EDTA, pH 7.4. Aliquots containing 3 mg of oxidised fatty acids were added to 1 mg HSA and incubated overnight at 20°C. Modified HSA was dialysed overnight at 4°C against PBS, pH 7.4.

MEASUREMENT OF ANTIBODIES AGAINST LIPID PEROXIDATION DERIVED ANTIGENS
Polystyrene microwell plates for ELISA (Immunolon IV; Nunc, Fisher Scientific, St Louis, Missouri, USA) were coated for four hours at 37°C with 0.05 mg/ml of HSA conjugated with either MDA, arachidonic acid oxidation products (HSA-APP), or with the unmodified protein solubilised in 0.1 M bicarbonate buffer, pH 9.6. After incubation, solutions were removed and replaced by 0.3 ml of coating buffer containing 3% BSA in PBS, pH 7.4. The plates were further incubated for one hour at 37°C to block non-specific binding sites. The coated wells were washed three times with PBS containing 0.25% Triton X-100. The sera of patients were diluted 1:50 with the coating buffer and added in duplicate as aliquots of 0.20 ml to the appropriate wells and incubated for one hour at 37°C. After washing three times with PBS-0.25% Triton X-100, IgG binding was revealed by peroxidase linked goat antihuman IgG as described above. The results were expressed by subtracting the spectrophotometric readings of each serum sample in the wells containing the unmodified HSA.

IMMUNOCOMPETITION EXPERIMENTS
Sera of ALD patients positive for anti-β₂-GP1 antibodies were diluted 1:50 with coating buffer and 300 µl aliquots placed in each well of microtitre plates coated with β₂-GP1 (IMTEC Immunodiagnostika GmbH, Berlin, Germany) or HSA. After overnight incubation at 4°C, 150 µl aliquots were added in duplicate to ELISA plates coated with oxidised cardiolipin, HSA-MDA, HSA-APP, or HSA-LPP and antibody binding was assayed as described above.

DATA ANALYSIS AND STATISTICAL CALCULATIONS
Data are expressed as mean (SD). Statistical analysis was performed by Instat-3 statistical software (GraphPad Software Inc, San Diego, California, USA) using one way ANOVA with Bonferroni’s correction for multiple comparisons when more than two groups were analysed. Pearson’s r values were used for estimation of correlation. Distribution normality of all groups was preliminary verified by the Komolgorov-Smirnov test. Significance was taken at the 5% level.

MATERIALS
Arachidonic acid, linoleic acid, L-phenylendiamine, cardiolipin, and fatty acid free human serum albumin (fraction V) were from Sigma Chemical Co. (St Louis, Missouri, USA). DPPD and malondialdehyde-bis-dimethylacetal were supplied by Fluka Chemie AG (Buchs, Switzerland).

Results
The use of commercial kits for the aPL assay based on binding to cardiolipin coated ELISA plates revealed that 20 of 36 (55%) patients with alcoholic cirrhosis with or without alcoholic hepatitis (ALD) had anticardiolipin IgG levels above the threshold value (15 GPL units/ml) while one healthy control was positive for anticardiolipin IgG. Mean IgG reactivity
towards cardiolipin among ALD patients was 22.7 (14) U/ml (range 11.8–71.1), significantly higher (p<0.001) than that in control subjects (11.8 (13) U/ml; range 10.9–17.6).

It has been shown that the unsaturated fatty acid component of cardiolipin rapidly auto-oxidises when exposed to air. Determination of conjugated diene absorbance at 234 nm, as an index of lipid peroxidation, revealed that overnight incubation at 4°C generally used for coating ELISA plates with phospholipids, appreciably increased the absorbance (od234 nm 0.276) of cardiolipin samples compared with freshly prepared solutions (od234 nm 0.110). Cardiolipin auto-oxidation was however completely prevented (od234 nm 0.112) by incubation under a nitrogen atmosphere in the presence of the antioxidant DPPD 100 µmol/l. When the antiphospholipid reactivity of sera was re-tested using ELISA plates coated with either oxidation protected cardiolipin or cardiolipin allowed to auto-oxidise in air, a dramatic difference was observed. As shown in fig 1, binding of IgG from ALD sera to oxidation protected cardiolipin (od490 nm 0.274 (0.184)) was significantly lower (p<0.0001) than that to oxidised cardiolipin (od490 nm 0.491 (0.340)). Reactivity of control sera with either form of cardiolipin was unchanged (od490 nm 0.205 (0.103) v 0.187 (0.111)) (fig 1). When tested with oxidised cardiolipin, 21 of 36 (58%) ALD sera showed IgG binding above the threshold value in controls given by mean od490nm values +2 SD (0.408). However, only eight of 36 (19%) ALD sera still displayed increased reactivity when tested with oxidation protected cardiolipin (cut off value 0.387) (fig 1). Furthermore, reactivity of ALD and control sera to oxidation protected cardiolipin was not statistically different. This was not due to interference of DPPD with the antibody reaction as addition of the antioxidant to already oxidised cardiolipin preparations did not affect IgG binding (not shown). No increase in reactivity towards oxidised cardiolipin was apparent in a group of 15 heavy drinkers without clinical or biochemical evidence of liver damage (0.153 (0.09)) or in 40 patients with cirrhosis unrelated to alcohol abuse (0.284 (0.10)).

Lipid hydroperoxides and aldehyde generated during lipid peroxidation are known to interact with proteins forming adducts that are immunogenic. We observed that ALD patients had enhanced level of circulating IgG against HSA modified by the reaction with malondialdehyde (HSA-MDA) or with oxidation products derived from arachidonic acid (HSA-APP) (od490 nm 0.79 (0.27) and 0.47 (0.25), respectively) v 0.45 (0.14) and 0.11 (0.08) in healthy controls; p<0.001). Individual levels of antibodies against oxidised cardiolipin were correlated with those of IgG directed against HSA-APP or HSA-MDA (r=0.67 and 0.60, respectively; p<0.0001). Moreover, among ALD patients, levels of antibodies against lipid peroxidation products were significantly higher in subjects positive for anti-oxidised cardiolipin antibodies than in the negative group (fig 2), indicating a relationship between the presence of an immune response against phospholipids and ethanol induced oxidative injury. On the other hand, titres of IgG against HSA-MDA and HSA-APP were not significantly increased in heavy drinkers without liver damage (od490 nm 0.450 (0.140) and 0.039 (0.030), respectively) or in non-alcoholic cirrhotics (od490 nm 0.590 (0.220) and 0.196 (0.151), respectively).

Antiphospholipid antibodies can also be detected by their ability to recognise complexes formed by β-GP1 and phospholipids. To avoid the confounding factor associated with cardiolipin oxidation state, antiphospholipid reactivity associated with ALD was re-evaluated by assaying anti-β-GP1 antibodies. IgG reactivity towards β-GP1 (46.1 (29.5) U/ml (range 13–150)) was significantly higher (p<0.001) in ALD patients than in control

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![Figure 1](http://gut.bmj.com/content/gut/49/6/852/F1.large.jpg)  
**Figure 1** Reactivity against native or oxidised cardiolipin (CL) of sera from 36 patients with alcoholic liver disease positive (Ox-CL) or negative (Ox-CL) for the presence of antibodies against oxidised cardiolipin. Human serum albumin (HSA) modified by the reaction with either malondialdehyde (HSA-MDA) or products from arachidonic acid peroxidation (HSA-APP) were used as antigens, as reported in the patients and methods section. Results are mean (SD) ELISA values. **p<0.01, ***p<0.0001 versus Ox-CL subjects.**
positive proteins with lipid peroxidation products of sera from patients with alcoholic liver disease (b2-GP1) were used to evaluate IgG binding. Results are mean (SD) ELISA values. **p<0.005, ***p<0.001 versus native cardiolipin.

**Figure 4** Reactivity with oxidised cardiolipin or human serum albumin (HSA) complexed with lipid peroxidation products following preabsorption with the b2-glycoprotein 1 (b2-GP1) of sera from alcoholic patients. Fifteen sera from patients with alcoholic liver disease displaying high titres of IgG against both cardiolipin and b2-GP1 were incubated overnight at 4°C on irradiated ELISA plates coated with b2-GP1 and tested for IgG binding to oxidised cardiolipin (Ox-CL) or HSA modified by reaction with malondialdehyde (HSA-MDA) or products of arachidonic acid peroxidation (HSA-APP). Results are mean (SD) ELISA values obtained with native sera or b2-GP1 preabsorbed sera. *p<0.0001 versus non-pre-absorbed sera.

Discussion

In recent years several reports have highlighted the association between alcoholic intoxication and the presence of circulating antiphospholipid antibodies (aPL). In these studies, the prevalence of aPL among patients with ALD ranged from 48% up to 81%. These values are in agreement with the prevalence of aPL (about 55%) observed in our group of patients with alcoholic liver cirrhosis. However, aPL have also been found in 31% of alcoholics and the presence of circulating antiphospholipid antibodies (aPL) has been correlated with severe ALD. Furthermore, circulating levels of IgG against oxidised cardiolipin were significantly higher (p<0.0002) in ALD patients with severe liver injury (Maddrey’s DF index >90) compared with patients with moderate or mild liver damage (Maddrey’s DF index <90).

**Figure 3** Reactivity with oxidised cardiolipin or with antigens derived from the reaction of proteins with lipid peroxidation products of sera from patients with alcoholic liver disease positive (b2-GP1; n=23) or negative (b2-GP1; n=13) for the presence of antibodies against b2-glycoprotein 1. ELISA plates coated with oxidation protected (CL) or oxidised (Ox-CL) cardiolipin, or with malondialdehyde (HSA-MDA) or products of arachidonic acid peroxidation (HSA-APP) were used to evaluate IgG binding. Results are mean (SD) ELISA values. **p<0.005, ***p<0.001 versus b2-GP1 negative subjects; †††p<0.001 versus native cardiolipin.

Subjects (22.5 (5.8) U/ml (range 10–32)) and 23 of 36 (64%) ALD patients were positive for anti-b2-GP1 IgG. b2-GP1 positive ALD sera showed titres of IgG recognising oxidised cardiolipin significantly higher (p<0.001) than b2-GP1 negative sera (fig 3), despite the fact that total circulating IgG levels were not significantly different between the two groups (15.89 (7.37) g/l v 13.26 (3.01) g/l). Furthermore, a strong positive correlation (r=0.85; p<0.0001) was evident when individual values of anti-b2-GP1 IgG were compared with those of antibodies directed against oxidised cardiolipin. Thus we postulated that aPL detected in ALD sera might recognise oxidatively modified cardiolipin complexed with b2-GP1. Indeed, preadsorption of ALD sera displaying both anticardiolipin and anti-b2-GP1 reactivity on ELISA plates coated with b2-GP1 reduced IgG binding to oxidised cardiolipin by about 80% (fig 4). ALD patients positive for b2-GP1 also showed reactivity towards HSA-MDA and HSA-AAP significantly higher (p<0.005) than that of b2-GP1 negative subjects (fig 3). To exclude the possibility that antibodies directed towards MDA or APP derived epitopes might account for the reactivity against oxidised phospholipids detected in the sera of ALD patients.

Patients with alcoholic liver disease (Child grades B and C). We have observed that individual reactivity against oxidised cardiolipin was positively correlated (r=0.65; p<0.0001) with Maddrey’s DF values. Furthermore, circulating levels of IgG against oxidised cardiolipin were significantly higher (p=0.0002) in ALD patients with severe liver injury (Maddrey’s DF index >90) compared with patients with moderate or mild liver damage (Maddrey’s DF index <90) (fig 5). A statistically significant difference (p<0.01) was also observed when the same patients were subgrouped according to the Child-Turcotte classification (fig 5). This indicated that the presence of antibodies against oxidised phospholipids might actually account for high aPL titres observed in association with severe ALD.

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Antioxidised phospholipid antibodies in alcoholics

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Little is known about the nature of the anti-

gens recognised by aPL associated with ALD. These antibodies have been detected by

binding to cardiolipin immobilised on the sur-

face of ELISA plates, but immune reactivity with other classes of phospholipids (that is, phosphatidyicholine, phosphatidylehol-

amine, phosphatidylserine, phosphatidylglycerol) have also been reported.$^{12}$ An elegant study by Hörkkö et al demonstrated that selected sera from women with antiphospho-

lipid syndrome bind to neo-epitopes generated when cardiolipin undergoes oxidation.$^{3}$ The fatty acid content of cardiolipin largely consists

(about 90%) of linoleic acid, and peroxidation of these fatty acids readily occurs when micro-

titre ELISA plates are exposed to air during the coating procedure.$^{4}$ We now report that anticardiolipin antibodies detected in sera of patients with ALD weakly react with native cardiolipin when it is protected from spontane-

ous auto-oxidation, but recognise instead anti-

gens present in oxidised cardiolipin.

Several lines of evidence indicate the involve-

ment of oxidative damage in alcoholic liver damage. Liver biopsies or sera from patients with ALD contain higher amounts of lipid per-

oxidation products (conjugated dienes, malondialdehyde, 4-hydroxynonenal, F$_{2}$-iso-

prostanates) and protein carbonyls than similar specimens from non-drinking subjects or patients with liver diseases unrelated to alcohol.$^{10-23}$ By reacting with proteins, free radicals, lipid hydroperoxides, and aldehydes generated from lipid peroxidation can produce a variety of adducts most of which are antigenic

and can stimulate the production of specific antibodies.$^{20,28}$ Increased levels of antibodies against hydroxyethyl free radicals and MDA-

protein adducts have previously been detected in patients with ALD.$^{24,10-32}$ We have observed that individual levels of IgG directed against adducts between serum albumin and either malonildialdehyde (HSA-MDA) or arachi-

donic acid oxidation (HSA-APP) products are closely associated with those of antioxidised cardiolipin antibodies. Furthermore, among patients with ALD, levels of antibodies against lipid peroxidation products are significantly higher in groups positive for antioxidised cardiolipin antibodies than in negative sub-

jects. Conversely, titres of IgG antioxidised cardiolipin in heavy drinkers without liver damage or in non-alcoholic cirrhosis that do not display evidence of oxidative injury are not significantly different from those in healthy controls. Thus we propose that the develop-

ment of antiphospholipid autoimmunity associ-

ated with ALD might be related to the pro-

oxidant conditions induced by ethanol. Such a hypothesis is consistent with experimental observations suggesting that oxidative modifications of phospholipids triggers the development of the antiphospholipid immune response.$^{10,33}$ Furthermore, Iuliano et al have shown that among patients affected by sys-

temic lupus erythematosus, those positive for aPL have higher urinary excretion of the lipid peroxidation product F$_{2}$-isoprostane than simi-

lar patients not displaying aPL reactivity.$^{14}$

It is now largely accepted that the plasma protein β$_{2}$-GP1 or apolipoprotein H plays a key role in aPL reactivity, serving as a cofactor for antibody binding to phospholipids.$^{4,24}$ In the liquid phase, β$_{2}$-GP1 is not recognised by aPL, but conformational changes occurring following interaction with phospholipids render β$_{2}$-GP1 antigenic for aPL.$^{7}$ We have observed that a high proportion (64%) of sera from patients with ALD displayed reactivity with β$_{2}$-GP1. A close correlation (r=0.85; p<0.0001) was also seen between individual levels of anti-β$_{2}$-GP1 IgG and those of antibodies directed against oxidised cardio-

lipin. Interestingly, only β$_{2}$-GP1 positive sera recognised antigens in oxidised cardiolipin and preadsorption of β$_{2}$-GP1 positive sera with β$_{2}$-GP1 attached to ELISA microtitre plates decreased antibody binding to oxidised cardi-

olipin antigens by 80%. This indicates that aPL

Figure 5 Reactivity against oxidised cardiolipin (Ox-CL) in relation to the severity of alcoholic liver injury. The clinical severity of hepatic damage was estimated according to Maddrey’s DF classification (A) or Child-Turcotte classification (B). Patients with DF values less than 90 (n=26) were considered to have mild or moderate hepatic injury and patients with DF values greater than 90 (n=10) represented the group with severe damage. Child A group consisted of 15 patients, while 21 subjects belonged to Child B-C groups. The optical density (od) values in each group were normally distributed, as evaluated by the Komologrov-Smirnov test. ***p<0.0002 versus DF <90; **p<0.01 versus Child A group.

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associated with alcoholic damage are directed against epitopes formed by the interaction of β$_2$-GP1 with oxidised phospholipids. Hörkkö et al have suggested that the neo-epitopes recognised by some aPL might consist of adducts formed between breakdown products of oxidised phospholipids and β$_2$-GP1. As mentioned above, circulating levels of antibodies directed towards lipid peroxidation related epitopes are increased in ALD patients. Furthermore, ALD patients displaying β$_2$-GP1 reactivity have higher serum levels of IgG recognising HSA-MDA or HSA-APP than β$_2$-GP1 negative subjects. However, preadsorption of ALD sera with β$_2$-GP1 that almost abolishes antibody binding to oxidised cardiop- lipin does not interfere with binding to epitopes in HSA complexed with MDA or with oxidised arachidonic or linoleic acids. We conclude that antibodies directed against protein adducts with lipid peroxidation products, such as MDA or lipid hydroperoxides, do not account for the immune reactivity towards oxidised cardiop- lipin. We propose that antiphospholipoid anti- bodies detected in patients with ALD rather recognise complexes between β$_2$-GP1 and oxid- ised cardiop- lipin.

Few data are available concerning the possible role of aPL in alcohol mediated liver injury. Antiphospholipoid immune reactivity can be detected in patients with alcoholic hepatitis or cirrhosis and particularly in non abstaining patients and in cirrhotics with severe liver damage (Child grades B and C). We have detected higher values of IgG against oxidised cardiop- lipin in ALD patients with severe liver damage. Moreover, a positive correlation (0.65; p<0.0001) was evident between individual antibody titers for oxidised cardiolipin in patients with alcoholic cirrhosis.

The clinical significance of aPL associated with alcoholic liver damage is at the moment unknown. The presence of aPL in patients with liver disease is not associated with an increased risk of thrombotic complications. Recent studies have proposed a possible link between aPL and cell death by apoptosis by showing that mice injected intravenously with syngenic monoclonal antibodies recognising oxidised cardiolipin die with liver injury. The epitopes for some antibodies detected in patients with ALD rather recognise complexes between β$_2$-GP1 and oxidised cardiolipin.

In conclusion, our results indicate that aPL detected in patients with ALD have a strict relationship with ethanol induced oxidative damage and preferentially target oxidised phospholipids complexed with β$_2$-GP1. These findings suggest a possible role for oxidative mechanisms in the development of aPL associated with alcoholic liver injury.

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10 Hörkkö S, Moller E, Branch DW, et al. The epitopes for some antiphospholipoid antibodies are adducts of oxidised phosphop- lipids and β$_2$-glycoprotein 1 (and other proteins).


