Apoptopes and the Biliary Specificity of Primary Biliary Cirrhosis

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Primary biliary cirrhosis (PBC) is characterized by antimitochondrial antibodies (AMAs), directed to the E2 component of the pyruvate dehydrogenase complex (PDC-E2). Notwithstanding the presence of mitochondria in virtually all nucleated cells, the destruction in PBC is limited to small intrahepatic bile ducts. The reasons for this tissue specificity remain unknown, although biliary epithelial cells (BECs) uniquely preserve the PDC-E2 epitope following apoptosis. Notably, PBC recurs in an allogeneic transplanted liver, suggesting generic rather than host PBC–specific susceptibility of BEC. We used cultured human intrahepatic BECs (HIBECs) and other well-characterized cell lines, including, HeLa, CaCo-2 cells, and nontransformed human keratinocytes and bronchial epithelial cells, to determine the integrity and specific localization of PDC-E2 during induced apoptosis. All cell lines, both before and after apoptosis, were tested with sera from patients with PBC (n = 30), other autoimmune liver and rheumatic diseases (n = 20), and healthy individuals (n = 20) as well as with a mouse monoclonal antibody against PDC-E2 and AMA with an immunoglobulin A isotype. PDC-E2 was found to localize unmodified within apoptotic blebs of HIBECs, but not within blebs of various other cell lineages studied. The fact that AMA-containing sera reacted with PDC-E2 on apoptotic BECs without a requirement for permeabilization suggests that the autoantigen is accessible to the immune system during apoptosis. Conclusion: Our data indicate that the tissue (cholangiocyte) specificity of the autoimmune injury in PBC is a consequence of the unique characteristics of HIBECs during apoptosis and can be explained by exposure to the immune system of intact immunoreactive PDC-E2 within apoptotic blebs. (HEPATOLOGY 2009;49:871-879.)

Abbreviations: AB, apoptotic body; AIH, autoimmune hepatitis; AMA, antimitochondrial antibody; BEC, biliary epithelial cell; BrEpC, bronchial epithelial cells; CHC, chronic hepatitis C; DAPI, 4',6-diamidino-2-phenylindole; FBS, fetal bovine serum; GCDC, sodium glycochenodeoxycholate; HIBEC, human intrahepatic cholangiocellular epithelial cell; M, mitochondria; mAb, monoclonal antibody; PBC, primary biliary cirrhosis; PBS, phosphate-buffered saline; PBS-T, PBS with 0.5% Tween 20; PDC-E2, E2 component of the pyruvate dehydrogenase complex; PSC, primary sclerosing cholangitis; SLE, systemic lupus erythematosus; UVB, ultraviolet B light.

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Apoptotic cells are normally efficiently cleared after engulfment by “professional” phagocytes followed by an anti-inflammatory response.1,2 When such uptake is impaired, cell lysis can release intracellular components that are a potential source of autoantigenic stimulation3-6 and autoimmunity onset.7-9 The presence of intact autoantigens within apoptotic blebs,10 their participation in the processes involved in autoantigen presentation,11 and the activation of innate immunity through macrophage cytokine secretion in concert12 are likely links between apoptosis and autoimmunity. Of relevance to the autoimmune liver disease primary biliary cirrhosis (PBC), Odin and colleagues demonstrated that, following apoptosis of biliary epithelial cells (BECs), the autoantigenic E2 subunit of the pyruvate dehydrogenase complex (PDC-E2) remains immunologically intact and still recognizable as such by antimitochondrial autoantibodies (AMAs).13 It is reasoned that absence of glutathiolation13 may contribute to this unique feature of the BEC. We reasoned that immune-mediated BEC destruction would be accentuated if PDC-E2 were preserved in blebs during apoptosis. Indeed, this could lead to impaired clearance of such apoptotic cells, thus provoking an in-
nate immune response and even the autoimmune destruction of bile ducts. This scenario would help explain the recurrence of PBC following orthotopic liver transplantation, as well as the therapeutic failure of immunosuppressive agents. We report here the cellular topology of PDC-E2 during apoptosis of cultured human intrahepatic BECs (HIBECs) and other non-BEC cells. Our findings demonstrate that PDC-E2 is localized intact within blebs of apoptotic HIBECs and is thereby accessible to the immune system. We hypothesize that the unique HIBEC apoptotic features allow the exposure of a potent intracellular autoantigen to the PBC-associated multilineage autoimmune response that leads to the tissue-specific autoimmune injury.

**Patients and Methods**

**Human Sera and Antibodies.** After informed consent was given, serum samples were obtained from patients diagnosed with PBC (n = 30), systemic lupus erythematosus (SLE; n = 5), autoimmune hepatitis (AIH, n = 5), primary sclerosing cholangitis (PSC, n = 5), chronic hepatitis C (CHC, n = 5), and healthy individuals (n = 10). PBC sera included 20 randomly chosen AMA-positive cases and 10 well-defined patients who were AMA-negative; thus, the proportion of AMA-negative sera utilized in this study is significantly higher than the normally expected frequency because these were specifically sought for this study. The diagnosis of all cases was based on established criteria for PBC, SLE, specifically sought for this study. The diagnosis of all cases was based on established criteria for PBC, SLE, cspecific serum pAbs, LLEO ET AL. HEPATOLOGY, March 2009 as well as the therapeutic failure of immunosuppressive agents. We report here the cellular topology of PDC-E2 during apoptosis of cultured human intrahepatic BECs (HIBECs) and other non-BEC cells. Our findings demonstrate that PDC-E2 is localized intact within blebs of apoptotic HIBECs and is thereby accessible to the immune system. We hypothesize that the unique HIBEC apoptotic features allow the exposure of a potent intracellular autoantigen to the PBC-associated multilineage autoimmune response that leads to the tissue-specific autoimmune injury.

**Antibody Reagents.** Serum anti–PDC-E2 antibodies were tested using our well-defined assays with recombinant antigens, as explained below. We also utilized a previously described mouse monoclonal antibody (mAb) against PDC-E2, clone 2H-4C8. Secondary antibodies Cy3-labeled anti–human-immunoglobulin G (IgG), anti–human-IgA, and anti–mouse-IgG were purchased from Jackson ImmunoResearch (West Grove, PA). Normal mouse IgG was obtained from Invitrogen (Carlsbad, CA) and fluorescein isothiocyanate (FITC)-labeled annexin-V was obtained from BD Pharmingen (San Jose, CA). Monoclonal anti–human caspase-3 antibody was purchased from R&D Systems (Minneapolis, MN). Negative controls were used throughout and included sera from healthy individuals. Additionally, we purified IgA from a patient diagnosed with PBC and a monoclonal gammopathy with high levels of IgA-AMA using Jacalin-agarose beads (Pierce, Rockford, IL) following the manufacturer’s instructions. Serum IgA from a healthy subject was used as a control.

**Detection of AMA.** An established optimal amount (16 μg) of purified recombinant PDC-E2 was loaded onto a 10% mini protein gel (Bio-Rad Laboratories, Hercules, CA), fractionated at 170 V for 1 hour, and transferred overnight onto nitrocellulose membranes that were then cut into strips. AMA detection was performed as described. The samples to be analyzed included sera from patients with PBC at an optimal dilution of 1:2000, monoclonal AMA (10^−5 dilution), pooled AMA of IgA isotype (dilution 1:2000), and control IgA (1:2000). Blots were exposed on photographic membranes and images were digitized with a FluorTech 8900 gel doc system (Alpha Innotech, San Leandro, CA) equipped with a chemiluminescent filter. The absence of AMA was further confirmed in the sera of the patients who were AMA-negative, using our standardized enzyme-linked immunosorbent assay (ELISA) and pMIT-3 antigens.

**Cell Lines and Culture Conditions.** The cells studied were cultured HIBECs, two human epithelial nontransformed primary cell cultures (human keratinocytes and bronchial epithelial cells [BrEpCs]) purchased from ScienCell (San Diego, CA) and two human tumor-derived laboratory cell lines, HeLa cells and CaCo-2 cells purchased from American Type Culture Collection (Manassas, VA). HIBECs were cultured in sterile medium supplemented with 2% fetal bovine serum (FBS), epithelial cell growth supplement (ScienCell, San Diego, CA), and 1% penicillin in cell culture flask coated with poly-L-lysine (Sigma-Aldrich, St Louis, MO). The other two epithelial cells were cultured under the same conditions in absence of FBS, as recommended by the manufacturer. HeLa and CaCo-2 cells were cultured using low glucose Dulbecco’s modified Eagle medium, supplemented with 10% FBS for HeLa cells and 20% for CaCo-2 cells, gentamicin (6 μg/mL), sodium pyruvate (110 mg/L), and L-glutamine (2 mM). Cells were cultured at 37°C in a humidified 5% CO2 incubator.

HIBECs were isolated from human liver tissue by the supplier and cryopreserved immediately after purification. This primary cell culture was characterized by an immunofluorescence method with antibodies to cytokeratin 18, cytokeratin 19, and vimentin which stained more than 90% of the cells. All experiments on HIBECs were performed between cell passage 2 and 5.

**Induction of Apoptosis.** We used biliary salts (BS) to
induce apoptosis.26 To establish optimal conditions for the induction of apoptosis, we incubated all the cell types at 37°C for 1, 2, 3, and 4 hours using different concentrations (100 μM, 500 μM, 1 mM, and 2 mM) of sodium glycochenodeoxycholate (GCDC; Sigma-Aldrich) added to normal culture medium, and in absence of serum and growth factors.26 BS failed to induce any apoptotic effect in HeLa and Caco-2 cells so, in these transformed cell lines, apoptosis induction was performed by ultraviolet B light (UVB) irradiation, at 1650 J/m² for HeLa cells and 2200 J/m² for CaCo-2 cells, followed by incubation in fresh media for 8 (HeLa) or 16 (CaCo-2) hours.13 Apoptosis was also induced in HIBECs by UVB irradiation (1650 J/m²) followed by incubation in fresh medium for 6 hours.

Quantification of Apoptosis by Flow Cytometry. The cells to be analyzed for apoptosis were suspended in 200 μL of buffer containing 10 mM Hepes/NaOH (pH 7.4), 140 mM NaCl, and 2.5 mM CaCl₂. A total of 1 × 10⁶ cells were stained for 15 minutes at room temperature in the dark with FITC-labeled annexin-V and with propidium iodide (BD Pharmingen, San Jose, CA) to discriminate apoptotic from necrotic cells. The samples were immediately analyzed by flow cytometry, and at least 10,000 events were counted. Stained cells were assessed on a fluorescence-activated cell sorting (FACS) FACScan flow cytometer (BD Immunocytometry Systems, San Jose, CA) upgraded by Cytec Development (Fremont, CA). Acquired data were analyzed with CellQuest Pro (BD Immunocytometry Systems) and FlowJo (Tree Star Inc., Ashland, OR) software packages.

Immunostaining and Confocal Microscopy. Cells were washed twice with phosphate-buffered saline (PBS) and incubated with FITC-labeled annexin-V at 1:10 dilution for 15 minutes at room temperature (RT) in the dark. The samples were then fixed in 3.7% formaldehyde (5 minutes at RT), permeabilized with 0.2% Triton X-100 (5 minutes at RT) and blocked for 30 minutes at RT with 1 × Universal blocking solution (Bio-Genex). Immunofluorescence staining was performed with human sera diluted 1:40 or monoclonal antibody diluted 1:80 (overnight, 4°C), followed by Cy3-labeled secondary antibody diluted 1:500 for 1 hour at RT. The cells were stained with 4‘,6-diamidino-2-phenylindole (DAPI; Invitrogen, Carlsbad, CA) to visualize nuclear degeneration. All samples were stained when 80% confluence; samples with less than 100 cells were excluded. Identical settings were used for all samples. Controls consisting of incubation of cells with developing secondary antibody alone did not demonstrate any detectable staining under the same conditions.

Staining of nonapoptotic and apoptotic HIBECs was also performed in the absence of membrane-perforating agent. The same protocol previously described was followed but Triton X-100 was omitted. Immunofluorescence-labeled samples were examined using a Pascal Zeiss confocal laser scanning microscope with a 100X oil-immersion objective.

Isolation and Immunoblot Analysis of Apoptotic Bodies. After induction of apoptosis, subcellular fragments were isolated by filtration and ultracentrifugation as described elsewhere.10 Briefly, the cell culture supernatant fluid was collected after apoptosis induction. Two additional centrifugation steps (500 g for 5 minutes) were performed in order to remove the remaining cells. The supernatant fluid was then passed through a 1.2 μm non-pyrogenic, hydrophilic syringe filter. After centrifugation at 100,000 g for 30 minutes, the pellet containing apoptotic bodies (AB) was resuspended in lysis buffer containing a protease inhibitor cocktail (Roche Diagnostics, Indianapolis, IN). Lysis was performed for 30 minutes on ice.

Protein content of the samples was determined by the BCA (bicinconic acid) assay using a Nanodrop ND-1000 ultraviolet-visible Spectrophotometer (NanoDrop Technologies, Wilmington, DE). Each sample (20 μg) was diluted in loading buffer and subjected to standard sodium dodecyl sulfate polyacrylamide gel electrophoresis. After transfer to polyvinylidene fluoride membranes, PDC-E2 was detected using the monoclonal antibody previously described (clone 2H-4C8),23 and AMA-positive sera with two different AMA isotypes.

Results

Apoptosis in HIBECs. Apoptosis rates were evaluated in all cell lines by FACS (annexin V and propidium iodide double staining) and immunofluorescence. Apoptotic cells were identified with confocal microscopy by morphological criteria: high nuclear density, chromatin condensation, and nuclear fragmentation revealed with DAPI, and characteristic blebbing of the cell membrane revealed with annexin-V; on the basis of these characteristics, an apoptotic index was established as: (DAPI-apoptotic nuclei/total nuclei) × 100.

Bile salts (BS) accumulating in the human liver in the course of cholestatic conditions trigger liver injury and subsequent fibrosis, and it has been demonstrated that a constituent of the hydrophobic BS, glycochenodeoxycholate (GCDC), induces apoptosis of hepatocytes starting at a concentration of 50 μM.27 GCDC has also an apoptotic effect in cholangiocytes.26 The use of 1 mM GCDC in the absence of serum, or growth factors, induced apoptosis in 39% of HIBECs, whereas UVB irra-
Radiation (1650 J/m² followed by incubation in fresh medium at 37°C for 6 hours) induced apoptosis in just 11% of HIBECs; moreover, in our experience, a higher level of double-positive propidium iodide/annexinV cells, most probably referable to necrotic cells, was generated following UVB irradiation compared to BS in HIBECs (Fig. 1A). Under both sets of conditions, membrane blebbing and apoptotic fragments were observed in HIBECs using confocal microscopy (Fig. 1C). The levels of apoptosis in HeLa and CaCo-2 cells after UVB irradiation were similar to those reported in the literature, i.e., 45%, and 60%, respectively. GCDC induced apoptosis in 49% of BrEpCs and 51.7% of keratinocytes. On the other hand, no substantial level of apoptosis was observed in HeLa and CaCo-2 cells after incubation with BS.

In conclusion, GCDC at 1 mM in serum-free medium was chosen to induce apoptosis in HIBECs, keratinocytes, and BrEpCs, whereas apoptotic HeLa and CaCo-2 cells were generated by UVB irradiation. To exclude that a different method of apoptosis was the reason for different expression of PDC-E2, staining with mAb against PDC-E2 using UVB-irradiated HIBECs was performed and the same results were observed.

**PDC-E2 Is Not Altered in Apoptotic HIBECs and Localizes Within Blebs.** The localization of PDC-E2 was studied in apoptotic HIBECs by indirect immunofluorescence and compared with other human cell lines. Apoptotic cells were identified by morphological criteria; DAPI staining was preferred over the use of terminal deoxynucleotidyl transferase–mediated dUTP nick-end labeling (TUNEL) to detect apoptotic cells because of its greater specificity, since TUNEL identifies fragmented DNA in both apoptotic and necrotic cells and hence may overestimate the number of apoptotic cells. First, HIBEcs were stained with sera from patients with PBC (n = 20) before and after induction of apoptosis, using both IgG and IgA AMA isotypes. As expected, nonapoptotic cells presented the typical punctuate cytoplasmic mitochondrial immunofluorescence pattern (Fig. 2A), negative staining for annexin-V, and normal nuclear morphology. After staining was performed under the same conditions following induction of apoptosis with 1 mM GCDC in the absence of serum and growth factors, apoptotic HIBEcs expressed positive PDC-E2 staining that could be localized with annexin V within apoptotic blebs and fragments (Fig. 2B). No detectable staining was...
noted on cells treated with either the secondary antibody alone or with normal mouse IgG instead of the mAb (data not shown). These findings were also reproduced in HIBECs following UVB irradiation and incubated in fresh media at 37°C for 6 hours (data not shown).

Immunofluorescence with murine mAb confirmed the results obtained with the human sera, and apoptotic HIBECs retained the PDC-E2–specific staining within the apoptotic blebs. There were no differences in staining using each of the AMA-positive sera from the patients with PBC according to age, disease stage, or therapy with ursodeoxycholic acid, and the staining patterns were similar to those obtained with the AMA-IgA (Fig. 2C,D). Preincubation overnight with human recombinant PDC-E2 completely removed the capacity of the sera to stain normal and apoptotic cells (Fig. 2). Serum samples from patients with AMA-negative PBC (n = 10), AIH (n = 5), SLE (n = 5), PSC (n = 5), CHC (n = 5), and healthy controls (n = 10) failed to produce any staining of HIBECs (Table 1).

Localization of PDC-E2 Within Apoptotic Blebs Is Specific for HIBECs. PDC-E2 localization was investigated using the clone 2H-4C8 mAb and sera from patients with PBC and controls using a variety of different human cell lines (Table 1, Fig. 3). All experiments were performed before and after induction of apoptosis under the same conditions as specified above for HIBECs. Before apoptosis induction, each of the cell lines manifested the typical cytoplasmic staining of mitochondria when the clone 2H-4C8 mAb was used (Fig. 3, upper row). Following apoptosis induction, none of the control cell lines had PDC-E2 staining (Fig. 3, lower row). None of the cell lines used demonstrated detectable cytoplasmic staining when sera from patients with AMA-negative PBC, AIH, SLE, PSC, CHC, or healthy controls were used (data not shown).

These morphological observations were corroborated by western blot analysis wherein PDC-E2 was readily de-

![Fig. 2. Staining of nonapoptotic or apoptotic HIBECs using an AMA-positive PBC serum (A,B), serum from a patient with PBC with monoclonal production of IgA-AMA (C,D), and using the same AMA-positive serum after absorption with recombinant PDC-E2 (E,F). The immunofluorescence staining was performed with three fluorochromes: Cy3-conjugated secondary antibody (red, yellow-orange when costained with green), FITC-labeled annexin-V (green), and DAPI (blue) for apoptosis detection. Apoptotic cells (*) were identified by morphological criteria: high nuclear density, chromatin condensation and nuclear fragmentation revealed with DAPI (blue), and characteristic blebbing of the cell membrane revealed with annexin-V (green). Positive staining of blebs and apoptotic fragments (arrows) was observed using unabsorbed PBC sera and was virtually absent after absorption. Apoptosis was confirmed in all experiments (*). Scale bar represents 20 μm.](image)

### Table 1. Prevalence of staining for PDC-E2 in blebs of various cell types undergoing apoptosis using PBC and control sera. Of note, specific bleb staining was observed in virtually all HIBEC when AMA-positive PBC sera were used. This staining was not observed in non-apoptotic cells.

<table>
<thead>
<tr>
<th></th>
<th>HIBEC</th>
<th>Control cells*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBC AMA +ve</td>
<td>20/20</td>
<td>0/40**</td>
</tr>
<tr>
<td>PBC AMA -ve</td>
<td>0/10</td>
<td>0/20</td>
</tr>
<tr>
<td>Control sera***</td>
<td>0/30</td>
<td>0/60</td>
</tr>
</tbody>
</table>

* Control cells include in equal distribution, transformed cells (HeLa and CaCo-2 cells) and non-transformed human epithelial cells (keratinocytes and bronchial epithelial cells).

** p value < 0.0001 compared to HIBEC

*** One third of the control sera were from healthy subjects, and two thirds were from patients with systemic lupus erythematosus, autoimmune hepatitis, primary sclerosing cholangitis or chronic hepatitis C.
detected in lysates of isolated apoptotic bodies obtained from HIBECs, whereas no such reactivity was observed in lysates from HeLa cells and BrEpCs (Fig. 4).

**PDC-E2 in Apoptotic Blebs Is Accessible to Antibody Recognition.** AMA staining in normal cells requires the use of a perforating agent due to the conservation of membrane integrity, but we hypothesize that during apoptosis the mitochondrial antigen became accessible to the immune system. To determine whether the permeabilization agents were responsible for the access of autoantibodies to PDC-E2 within blebs, we stained HIBECs with the clone 2H-4C8 mAb in the absence of Triton X-100. Results obtained showed that while the absence of Triton X-100 did not modify the positive staining of blebs in apoptotic HIBECs (Fig. 5), no differences were observed in the other cell lines because all of them lose AMA staining during apoptosis. These results were obtained with 12 of 20 AMA-positive PBC samples whereas sera from patients with AMA-negative PBC (n = 5), AIH (n = 5), SLE (n = 5), PSC (n = 5), CHC (n = 5), or healthy controls (n = 5), or the use of secondary antibody alone failed to demonstrate any staining (Table 3).

**Table 2. Number of apoptotic cells in which blebs contain PDC-E2.** The apoptotic cells were stained as described with sera from PBC AMA positive patients (n=20). Values are expressed as mean ± SD. Apoptotic index was derived from DAPI-apoptotic nuclei/total nuclei x 100. An individual apoptotic cell was considered positive when PDC-E2 staining was observed in at least one of its blebs. Student’s t-test was used to calculate p values.

<table>
<thead>
<tr>
<th>Cell type</th>
<th>Apoptotic index</th>
<th>Cells with positive staining for PDC-E2 within blebs</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIBEC</td>
<td>38 ± 2.2</td>
<td>88 ± 0.7</td>
</tr>
<tr>
<td>Keratinocytes</td>
<td>51 ± 0.7</td>
<td>0 ± 1.3*</td>
</tr>
<tr>
<td>BrEpC</td>
<td>42 ± 1.2</td>
<td>0 ± 0.7*</td>
</tr>
<tr>
<td>HeLa</td>
<td>61 ± 3.5</td>
<td>0 ± 0.3*</td>
</tr>
<tr>
<td>CaCO-2</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* p value < 0.0001 compared to HIBEC

**Discussion**

We demonstrate in this report that PDC-E2, the major AMA autoantigen, is detectable in its antigenically reactive form within apoptotic blebs, specifically in cultured human BECs and not in other cell types, thus providing one explanation for the organ-specific pathology noted in human PBC. We thus suggest that the unique characteristics of BECs during apoptosis might constitute the pathogenic link between the ubiquitous distribution and high degree of conservation across species of the AMA autoantigen and the organ specificity of PBC pathology.

It has been previously reported that PDC-E2 remains intact and retains its immunogenicity during BEC apoptosis due to a cell lineage-specific lack of glutathiolation.13,29 This “apoptotic exposure” of PDC-E2 appears to be limited to BECs, and it may ultimately have a critical pathogenic relevance to both inductive and effector stages of PBC. The formation of apoptotic bodies and fragments is essential during apoptosis to limit the escape of intracellular content and preclude any ensuing immunological responses against intracellular autoantigens with inflammatory reactions.30,31 Nevertheless, apoptotic blebs and...
fragments can under some circumstances constitute a major source of immunogens in autoimmune diseases that involve the targeting of ubiquitous autoantigens. Thus, dysregulation of apoptosis or the ineffective removal of apoptotic cells has been documented in patients with SLE and the development of antibody-mediated myocarditis of infants born to mothers with anti-SSA/SSB/La. Also, Kupffer cell engulfment of apoptotic bodies from hepatocytes promotes inflammation and fibrogenesis.

Our observations may help seal several remaining gaps in the understanding of induction and perpetuation of PBC, albeit raising new questions. First, intact PDC-E2 in apoptotic fragments from BECs could be taken up by intrahepatic dendritic cells and transferred to regional lymph nodes for priming of cognate T cells, thus initiating PBC, but this attractive scenario still requires an explanation for the 
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in the first place for apoptosis and this is likely not PBC-specific. Second, the accessibility of PDC-E2 within apoptotic blebs to autoantibodies appears to support the pathogenic role of AMA as well as T cells in the perpetuation of BEC injury, even though antibody titers do not correlate with the clinical features or stages of PBC, and AMA-negative patients are clinically indistinguishable from their AMA-positive counterparts. Nevertheless, the appearance of serum AMA does often herald disease onset sometimes by several years. Third, we can propose that PDC-E2 within apoptotic blebs will also be recognized by major histocompatibility complex class I–restricted CD8 T cells; this point helps explain the BEC pathology in AMA-negative PBC. Interestingly, our lab has recently demonstrated the presence of autoreactive T cells to PDC-E2 in patients with AMA-negative PBC. These data are also of particular relevance in view of the major pathogenic role of these cells in producing PBC-like liver lesions in animal models. Fourth, our findings are consistent with the likelihood that PBC cholangiocytes do not manifest any unique features that make them the target of autoimmunity, noting the frequent recurrence of PBC following allogeneic liver transplantation. The latter two issues may ultimately be combined with the fact that the donor and recipient major histocompatibility complex class I alleles are major determinants of the allograft outcome.
Fifth and ultimately, the ensuing B and T cell autoreactive response may account for the perpetuation of the immune-mediated damage to BECs with a major role also played by elements of innate immunity which appears to be enhanced in PBC.40,41

Our data imply that the postapoptotic release of intact mitochondrial autoepitopes in small bile ducts is one contributor to this specificity. Indeed, we should note that, as previously reported, the overexpression of Bcl-2 (B cell lymphoma-2), specifically in apoptotic small BECs, inhibits PDC-E2 glutathiolation and prevents the loss of antigenicity.13,42 However, other factors have also been incriminated in playing a role in the selective destruction of small BECs. In particular, there are dramatic differences in expression of trefoils in small versus large bile ducts, suggesting not only an imbalance of homeostasis, but also a differential ability to repair or restitute cell damage.13

Our data also demonstrate that we are able to detect PDC-E2 without cell permeabilization. There are three explanations for this observation. First, PDC-E2 may leak out to the cell surface and is thus being detected on the cell membrane. Second, the cells undergoing apoptosis have holes in their cell membrane created by cellular proteases which allow passage into and localization of immunoglobulin in the bleb. Third, there may be a role for fragment crystallizable receptor–mediated uptake in the apoptotic cell. Future experiments will address these possibilities.

In conclusion, the evidence provided here leads to new scenarios in the pathogenesis of PBC and may constitute a credible link between the several convenient and inconvenient truths available thus far.37 However, it does not overcome all of the major challenges in PBC etiology, nor the need to ascertain the genetic basis of disease susceptibility and environmental triggers for cholangiocyte injury and apoptosis as an initial step in tolerance breakdown. Treatments that could modulate apoptosis44 should not be overlooked, and their assessment is warranted in recently established murine models for PBC.45-47

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References


Table 3. Percentage of apoptotic HIBEC with positive staining for PDC-E2 in the presence or absence of a permeabilizing agent during the staining as described in the materials and methods. None of the control sera showed any positive staining. Values are expressed as mean ± SD.

<table>
<thead>
<tr>
<th>Sera (n)</th>
<th>Proliferating HIBEC</th>
<th>Non permeabilized</th>
<th>Permeabilized</th>
</tr>
</thead>
<tbody>
<tr>
<td>mAb 2H-4C8</td>
<td>93 ± 7</td>
<td>37 ± 12</td>
<td>91 ± 5*</td>
</tr>
<tr>
<td>PBC AMA –ve (20)</td>
<td>88 ± 6</td>
<td>33 ± 7</td>
<td>88 ± 0.7*</td>
</tr>
<tr>
<td>PBC AMA –ve (10)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Control (30)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

* P value < 0.0001 compared to non permeabilized apoptotic HIBEC


