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**Natural antibodies with the T15 idiootype may act in atherosclerosis, apoptotic clearance, and protective immunity**

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The immune response to oxidized LDL (OxLDL) may play an important role in atherogenesis. Working with apoE-deficient mice, we isolated a panel of OxLDL-specific B-cell lines that secrete IgM Abs that specifically bind to oxidized phospholipids such as 1-palmitoyl-2-(5-oxovaleroyl)-sn-glycero-3-phosphorylcholine (POVPC). These Abs block uptake of OxLDL by macrophages, recognize similar oxidation-specific epitopes on apoptotic cells, and are deposited in atherosclerotic lesions. The Abs were found to be structurally and functionally identical to classic “natural” T15 anti-PC Abs that are of B-1 cell origin and are reported to provide optimal protection from virulent pneumococcal infection. These findings suggest that there has been natural selection for B-1 cells secreting oxidation-specific/T15 antibodies, both for their role in natural immune defense and for housekeeping roles against oxidation-dependent neodeterminants in health and disease.


**Introduction**

Atherosclerosis is a chronic inflammatory disease that results from hyperlipidemia and a complex interplay of environmental, metabolic, and genetic risk factors. There is much evidence that the oxidation of LDL plays a central, if not obligatory, role in the atherogenic process (1). In addition, oxidation of LDL generates a variety of oxidatively modified lipids and proteins that represent highly immunogenic neodeterminants for the immune system. In murine models of atherosclerosis, such as apoE-deficient (ApoE−/−) mice, atherosclerosis is correlated with the development of high titers of autoantibodies to various oxidation-specific neoepitopes of oxidized LDL (OxLDL) (2, 3). The consequences of such cellular and humoral responses are still poorly understood, but under certain conditions they can clearly modify the natural history of the disease. For example, immunization of hypercholesterolemic rabbits or mice with homologous LDL containing epitopes of OxLDL ameliorates atherosclerosis (4, 5).

To investigate the origins of these atherosclerosis-associated autoantibodies, we recently isolated a panel of B-cell hybridomas with specificity for OxLDL neodeterminants from the spleens of diseased ApoE−/− mice that had not received experimental immunization or in vitro stimulation (3). Reflecting the disease-associated immune response, a minimum of 32% of 1,536 Ig-secreting hybridomas expressed Abs that bound to one or more epitopes of OxLDL, of which 17 hybridoma lines were cloned and their secreted Abs termed “EO” Abs. All were IgMs that bound strongly to either a model epitope of Ox-LDL, malondialdehyde-modified LDL (MDA-LDL), or to LDL that was oxidized by exposure to copper (Cu-OxLDL) (3). Each of the Abs selected for binding to Cu-OxLDL was later shown to bind to oxidized phospholipids, and specifically to 1-palmitoyl-2-(5-oxovaleroyl)-sn-glycero-3-phosphorylcholine (POVPC). (Please note that in this article PC stands for phosphorylcholine, not for phosphatidylcholine, an intact phospholipid containing two fatty acid chains and the phosphorylcholine headgroup.) Of considerable interest is the fact that all of these POVPC-specific autoantibodies recognized the lipid moiety of OxLDL, as well as the delipidated modified apoB (the protein moiety of OxLDL), but not native LDL. Thus, these Abs bind to oxidized phospholipids or their protein adducts (e.g., with apoB or BSA). In addition, each of these Abs was able to block macrophage scavenger receptor–mediated binding and uptake of OxLDL. POVPC (as a BSA adduct) also blocked the uptake of OxLDL (6). Macrophage scavenger receptors involved in uptake of OxLDL are also believed to be responsible for the recognition of apoptotic cells, as well as dam-
aged or senescent cells. We demonstrated that the POVPc-specific EO Abs also bound to apoptotic cells and inhibited their phagocytosis by elicited peritoneal macrophages. Similarly, POVPc-BSA also inhibited the phagocytosis of apoptotic cells by macrophages (7). These data provide compel[i ng evidence that these naturally arising autoantibodies bind to common oxidation-specific epitopes on both OxLDL and apoptotic cells that mediate macrophage recognition.

To gain a better understanding of the roles these IgM Abs play in health and disease, we investigated their genetic and structural origin. Our studies suggest that these anti-OxLDL autoantibodies represent highly conserved natural Abs that arise spontaneously in the perinatal period, and later become further selected in ApoE−/− mice in response to the extensive burden of oxidation-specific antigens that occur with the development of atherosclerosis.

Methods

Antigens and ligands. Phosphorylcholine chloride (PC-Cl) was obtained from Sigma Chemical Co. (St. Louis, Missouri, USA). PC-conjugated keyhole limpet hemocyanin (PC-KLH) was from Biosearch Technologies Inc. (Novato, California, USA). PC-BSA and PC-histone were kind gifts of Norman Klinman (The Scripps Research Institute, La Jolla, California, USA) and Marvin Rittenberg (Oregon Health Sciences University, Portland, Oregon, USA), respectively. Purified pneumococcal cell-wall polysaccharide (C-PS) was obtained from the Staatsen serum Institute (Copenhagen, Denmark). POVPc was synthesized, and POVPc-BSA adducts were generated in the presence of cyanoborohydride, as described (6). LDL was prepared in the presence of EDTA, and copper-oxidized LDL (Cu-OxLDL) and MDA-LDL were prepared as described (3).

Hybridomas and mAb’s. Monoclonal autoantibodies directed against oxidation-specific epitopes of LDL were produced from hybridomas generated from apoE-deficient mice, which have very high titers of autoantibodies to Cu-OxLDL (3). In brief, B lymphocytes from the spleens of two apoE-deficient mice that had not been immunized exogenously were fused with the P3 × 63Ag8.653.1 myeloma cell line. Hybridomas were screened for binding to model epitopes of OxLDL, including Cu-OxLDL and MDA-LDL. Seventeen hybridoma cell lines were isolated by limiting dilution, and the Abs expressed were designated EO autoantibodies EO1–EO17. All were isolated as IgM and characterized as described (3, 6). Abs used in this report were purified from ascites fluid by fast protein liquid chromatography (FPLC). The T(EPC)15 murine monoclonal IgA anti-PC Ab (referred to hereafter as T15) was the kind gift of H. Kohler (University of Kentucky, Lexington, Kentucky, USA). Anti-idiotype Abs Tc54.6 and T139.2 were kind gifts of M. Scharff (Albert Einstein College of Medicine, Bronx, New York, USA). AB1-2, an anti-T15 idiotype Ab, was provided by J. Kearney (University of Alabama at Birmingham, Birmingham, Alabama, USA).

Chemiluminescent immunoassay. To coat wells, protein antigens were diluted in PBS containing 0.27 mM EDTA; 50 μL per well was applied to 96-well white round-bottomed MicroFluor microtiter plates (Dynex Technologies, Chantilly, Virginia, USA) and left overnight at 4°C. The wells were washed three times with PBS, and blocked with PBS containing 1% BSA (BSA-PBS) for 30 minutes. Fifty microliters of primary Abs, diluted with BSA-PBS, were added to wells in the presence and absence of competitors, and then incubated for 1 hour at room temperature. The wells were washed three times with PBS. Binding of primary Ab was detected with isotype-specific alkaline phosphatase-labeled (AP-labeled) goat anti-mouse Ig secondary Abs (Sigma Chemical Co.) in Tris-buffered saline (TBS) buffer containing 150 mM NaCl, 50 mM Tris base, 0.27 mM EDTA, and 1% BSA. This was followed by a rinse with water and the addition of 25 μL of 50% LumiPhos 530 solution (Lumigen Inc., Southfield, Michigan, USA). In some experiments, the mAb EO6 was biotinylated as described previously (6) and added to antigen-coated wells, followed by the addition of AP-labeled NeutrAvidin (Pierce Chemical Co., Rockford, Illinois, USA). The light emissions were measured as relative light units (RLU) over 100 ms using a Dynex Luminometer (Dynex Technologies) (6).

Isolation and amplification of Ab genes. Total RNA was extracted from 107 hybridoma cells using RNA STAT-60 (Tel-Test Inc., Friendswood, Texas, USA) using the manufacturer’s protocol. A Superscript II cDNA synthesis kit (Gibco BRL, Gaithersburg, Maryland, USA) was then used to synthesize cDNA with oligo(dT) primer. Each PCR reaction contained 0.5 μg of cDNA template and 1 μg of each oligonucleotide primer. The reactions were supplemented with dNTP, Taq polymerase (Pharmacia, Peapack, New Jersey, USA), and reaction buffer, and were amplified under optimized thermal cycler conditions. The oligonucleotide primers for PCR reactions were synthesized in the Molecular Biology Core facility of the La Jolla Specialized Center of Research for Molecular Medicine and Atherosclerosis. For the first round of PCR, heavy-chain variable (VH) genes were amplified in seven different reaction tubes, each containing one of a panel of seven sense primers targeted for the VH framework region 1 (FR1) subdomain, paired with an antisense primer specific for the Jh constant region (CH1) (Table 1). The oligonucleotides were designed to amplify more than 80% of all expressed VH gene rearrangements (8). In preliminary studies, an immunoassay using anti-κ specific Ab determined that each of the reported EO Abs uses a κ light chain (not shown). Therefore, Vκ gene rearrangements were amplified in six separate PCR reactions that included one of two degenerate VC κ gene FR1 sense primers, and one of three anti-Jκ sense primers that each target distinct Jκ gene sequences (Table 1).

Sequence analysis of variable-region genes of EO hybridomas. The PCR products of the murine hybridoma Ig VH and Vκ gene rearrangements were purified after elec-
trophoretic separation on a 2% agarose gel and then cloned into the TA cloning vector pCR2.1 (Invitrogen Corp., Carlsbad, California, USA), according to the manufacturer’s protocol. Plasmid DNA from individual clones was purified using QIAprep columns (QIAGEN Inc., Chatsworth, California, USA), and the nucleotide sequences were determined with an ABI PRISM automated sequence analyzer (Perkin-Elmer Applied Biosystems, Foster City, California, USA) using T7 and M13 primers. The sequences were analyzed using software packages from the Genetics Computer Group (Madison, Wisconsin, USA) in combination with the EMBL/GenBank databases. Sequence alignments of the identified V_H and V_L genes were also determined with BLAST tools of the National Center of Biotechnology Information (Bethesda, Maryland, USA).

**Induction of apoptosis and flow cytometry analysis.** Porcine aortic endothelial cells (PAECs) from wild-type swine were cultured in Medium 199 containing Earl’s salt solution and 15% FCS in a CO_2-enriched atmosphere, as described elsewhere (7). Apoptosis was induced during the sixth passage by culturing PAECs for 16 hours in serum-free media, using Teflon® membrane-coated culture flasks (Sarstedt, Newton, New Jersey, USA) to inhibit anchorage-dependent cell spreading. Cells were then harvested and washed in ice-cold PBS with 1% BSA. To determine Ab binding, 10^6 cells were incubated with 50 μg/mL of T15 or isotype control murine IgA (from a myeloma cell line) at 4°C for 20 minutes; cells were then washed and incubated for another 20 minutes with 1 μg/mL of fluorescein-conjugated anti-mouse IgA (PharMingen, San Diego, California, USA). Cells were again washed and then incubated for 10 minutes with 1 μg/mL of PI. Data were immediately acquired with a FACScan instrument (Becton Dickinson and Co., Franklin Lakes, New Jersey, USA), and analyzed using CellQuest software.

**Immunohistochemistry.** Murine aortas were perfusion fixed, and segments containing large atherosclerotic lesions were embedded in paraffin. Blocks were sectioned, and sections were immunostained for oxidation-specific epitopes and macrophages (9). The presence of mouse IgG in lesions was detected by staining with antisera to mouse IgG and IgM (9), using biotinylated mouse Ig in lesions was detected by staining with antisera to mouse IgM and IgA (PharMingen, San Diego, California, USA). Double staining with murine autoantibodies in lesions, to assess the presence of EO/T15 idiotype Abs (0.25 μg/mL). As a positive control, adjacent sections were stained with the same protein concentration of an irrelevant biotinylated mouse IgG. Abs bound to the tissue were then detected with an avidin-biotin–AP complex and phosphatase substrate. Endogenous tissue AP activity was blocked by 15 minutes of incubation with 5 mM levamisole (Sigma Chemical Co.).

**Genomic DNA PCR to determine the origin of EO hybridomas.** PCR primers were designed to cover V_H framework region 3 (FR3) (FR3S, 5’-CTCGAYATTTTGATGACNCAR-3’ and the downstream JH4 gene 3’ flanking region (JH4AS, 5’-AAC TCC ATA ACA AAG GTT AAA AAT AAA G). Independent PCR reactions were performed with 0.5 µM of each primer and 2 µg of genomic DNA template from each hybridoma in the optimized condition for 32 cycles. Genomic DNA extracted from mouse fibroblasts was used as a negative control. The PCR products were analyzed on 1% agarose gel and subcloned in a TA cloning vector (Invitrogen Corp.) for later sequence determination.

To further analyze the amplified fragments from these EO Ab–expressing clones, the PCR products were purified with QIAprep micro spinning columns (QIAGEN Inc.). The DNA was then digested with restriction endonuclease HindIII or BstNI, followed by 1% agarose gel electrophoresis to compare the fingerprint patterns of the digested products.

**Results**

**Genetic origins of EO Abs.** Preliminary data documented that each of the EO B-cell hybridomas expressed an IgM-k Ab (data not shown). Using a strategy designed to enable the unbiased cloning of the vast majority of rearrangements among V_H and V_L genes (Table 1), DNA was extracted first from the EO6 cell line, the best characterized of the POVPC-specific EO Ab–expressing lines. In four of seven separate PCR reactions, each of which targeted different V_H FR1 sequences, amplifiers of the expected size (~380 bp) were obtained. After

### Table 1

<table>
<thead>
<tr>
<th><strong>Primers designed to amplify V_H and V_L gene rearrangements of autoantibodies cloned from apoE−/− mice</strong></th>
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<tbody>
<tr>
<td><strong>V_H FR1–specific sense primers</strong></td>
</tr>
<tr>
<td>CLIII-FRIA</td>
</tr>
<tr>
<td>CLIII-FRI</td>
</tr>
<tr>
<td>CLI-FRIB</td>
</tr>
<tr>
<td>CLI-FRIA</td>
</tr>
<tr>
<td><strong>V_L-degenerate sense primers</strong></td>
</tr>
<tr>
<td>V_A-degA</td>
</tr>
<tr>
<td>V_A-degB</td>
</tr>
<tr>
<td><strong>JL-specific sense primers</strong></td>
</tr>
<tr>
<td>C(3)J1,2</td>
</tr>
<tr>
<td>C(3)L4</td>
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</tbody>
</table>

V_H primers include seven sense primers specific to FR1 of repertoires, and an antisense primer specific to the CH1 domain of the IgH gene. V_L primers include two degenerate (deg) sense primers, which will be primed to FR1 of most V_L genes, and three antisense primers specific to J_2, J_4, and J_5 junction regions.
independent cloning and DNA sequence determinations, each of the cloned amplimers was found to represent the same unique in-frame Vh-μ gene rearrangement, derived from the S107 family. This gene rearrangement was 100% identical to a previously reported nonmutated, functional, canonical VHS107.1-DPL16.1-JH1 rearrangement that is also expressed in the T(EPC)15 IgA-κ plasmacytoma (10) (Figure 1). Amplimers of the expected size (~350 bp) were obtained in two of six separate Vh gene amplification reactions of cDNA from EO6 cells (Table 1). Sequence determination of plasmids containing these independently cloned inserts yielded the same in-frame nonmutated, canonical Vh22-J5 rearrangement that was also previously reported for the T(EPC)15 plasmacytoma.

To independently confirm the genetic origin of the Vh gene expressed by the EO6 cell line, separate RT-PCR reactions were performed using primers specific to unique upstream leader-specific sequences that can discriminate between the V1 gene and the V11/13 functional genes of the S107 family. Amplification was obtained only for the V1-specific primer, and subsequent sequence analysis of the cloned product confirmed the exact nonmutated Ab gene sequence identified in the earlier Vh gene cloning studies.

Equivalent studies were also performed with cDNA from three other OxLDL-reactive POVPC-specific cell lines. As with EO6, DNA sequence analysis of the VH and Vh genes of the EO3, EO4, and EO7 cell lines indicated that they were each 100% homologous to the classical T15 Ab. Figure 2 shows the deduced amino acid sequences for these Abs; the Ab gene usage for each of these EO cell lines and their binding reactivities (6) are compiled in Table 2.

**EO autoantibodies are recognized by T15-specific anti-idiotypes.** To characterize the Vh and Vl regions expressed in different EO Abs and determine their relationship to the classic T15 Ab, we tested the recognition of these Abs by three well-characterized T15-specific anti-idiotypic Abs. These were T139.2, directed against the unique T15-specific Vh22-J5 light chain (11); Tc54.6, directed against the unique T15-specific VHS107.1 usage (11, 12); and AB1-2, directed against a conformational determinant requiring both the heavy- and light-chain CDR3 regions of T15 (13). As shown in Table 2, each of the POVPC-specific, anti-OxLDL Abs for which the VH and Vl region sequences were determined (EO3, EO4, EO6, and EO7) also expressed the T15 idiotype (i.e., they are recognized by each of three anti-idiotypic Abs). In addition, three other POVPC-specific autoantibodies, EO12 and EO14, which bind MDA-LDL (3), did not express any of the T15 markers.

**Binding of EO6/T15 to OxLDL and PC.** T15 is a classic Ab to PC, the “headgroup” moiety [-PO4-(CH2)2-N-(CH3)3] of many phospholipids, including PAPC. Because the genetic and idiotypic analyses indicated that these EO autoantibodies to OxLDL used Vh1 and Vl regions identical to those of T15, we systematically compared the antigen-binding properties of T15 (an IgA) with a representative EO Ab, EO6 (an IgM). As expected, EO6 bound

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**Table 2**

Summary of gene usage and immunological properties of T15 and autoantibodies from apoE-deficient mice

<table>
<thead>
<tr>
<th>Clone name</th>
<th>Vh usage</th>
<th>Vl usage</th>
<th>AB1-2 (S107.1/k22) idiotype</th>
<th>T139.2 (k22) idiotype</th>
<th>Tc54.6 (S107.1) idiotype</th>
<th>Binding to Cu-OxLDL</th>
<th>Binding to POVPC-BSA</th>
<th>Binding to MDA-LDL</th>
<th>Binding to PC-KLH</th>
<th>Binding to C-PS</th>
</tr>
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<tbody>
<tr>
<td>EO0</td>
<td>S107.1</td>
<td>k22</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>–</td>
<td>++++</td>
</tr>
<tr>
<td>EO1</td>
<td>S107.1</td>
<td>k22</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>–</td>
<td>++++</td>
</tr>
<tr>
<td>EO6</td>
<td>S107.1</td>
<td>k22</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>–</td>
<td>++++</td>
</tr>
<tr>
<td>EO7</td>
<td>S107.1</td>
<td>k22</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>–</td>
<td>++++</td>
</tr>
<tr>
<td>EO12</td>
<td>ND</td>
<td>ND</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
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<tr>
<td>EO14</td>
<td>ND</td>
<td>ND</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
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</tr>
<tr>
<td>T15</td>
<td>S107.1</td>
<td>k22</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
<td>++++</td>
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Clones secreting EO Abs were sequenced and analyzed for their germline gene usage, and compared with the T15 Ab. Their recognition by anti-T15 idiotypic Abs (AB1-2, T139.2, and Tc54.6) and their binding to oxidation epitopes (Cu-OxLDL, MDA-LDL, and POVPC-BSA) and classical PC epitopes (PC-KLH, C-PS) are compared. Note the identical patterns of EO Abs and T15. ND, not determined.
to Cu-OxLDL and POVP-BSA, but not to native LDL or MDA-LDL (6) (Figure 3a). As reported previously, T15 bound to its known antigen, PC, derivatized onto the carrier proteins BSA and KLH (data not shown) (14). Like EO6, T15 also bound to Cu-OxLDL and POVP-BSA, but not to native LDL or MDA-LDL (Figure 3b). In addition, with reactivity similar to that of T15, EO6 bound to the PC conjugates PC-BSA and PC-KLH. Neither EO6 nor T15 bound to the native, unoxidized phospholipid PAPC (data not shown).

To further investigate the fine binding specificity of EO6 to PC, a competition immunoassay was performed. Figure 4 shows that the binding of EO6 to Cu-OxLDL was totally inhibited by soluble PC (PC-Cl) and by PC-KLH. In analogous experiments, the binding of T15 to Cu-OxLDL was also inhibited by PC-Cl in a dose-dependent manner (data not shown). In control experiments, the binding of another oxidation-specific IgM monoclonal autoantibody, EO14, to its antigen, MDA-LDL, was unaffected by PC-Cl (data not shown).

During the oxidation of LDL, a large number of reactive lipid peroxidation products are generated that can serve as epitopes for Ab recognition. Therefore, we directly tested the ability of T15 to compete with biotinylated EO6 for binding to OxLDL. The inset in Figure 4 demonstrates that T15 competed effectively, whereas a control murine IgA did not. This suggests that these two mAb’s are directed against an immunologically closely related (if not identical) epitope on OxLDL. In addition, we demonstrated that the binding of EO6 to PC-KLH was inhibited by both PC-KLH and Cu-OxLDL in a dose-dependent manner (data not shown). In addition, the demonstration that the anti-idiotypic T139.2 competed very effectively with EO6 for binding to PC-KLH provided further confirmation that the binding of EO6 to PC uses the same antigen-binding site used by the classic T15 Ab (data not shown).

**EO6 and T15 bind to bacterial C-PS.** PC is the immunodominant determinant in the teichoic acid–containing C-PS of *Streptococcus pneumoniae* (15), and both EO6 and T15 bound to C-PS, whereas EO14 (an MDA-LDL–specific Ab) did not (Figure 5a). Furthermore, the binding of both EO6 and T15 to OxLDL was dose-dependently inhibited by the addition of C-PS (Figure 5b). These data indicate that both EO6 and T15 recognize a common
epitope present on OxLDL as well as on C-PS.

T15 Abs recognize apoptotic but not normal endothelial cells. We have previously shown that EO6 bound to apoptotic cells and inhibited their phagocytosis by macrophages (7). To test whether this was a common feature of all T15 Abs regardless of isotype, we examined the capacity of the classic IgA T15 to bind to apoptotic PAECs by flow cytometric analysis. In these studies, serum deprivation was used to induce apoptosis in PAECs. As illustrated in Figure 6, T15 preferentially bound to PAECs that were in advanced stages of apoptosis (Figure 6c), but not to normal cells or those in very early stages of apoptosis (Figure 6b).

Abs with the T15 idiotype are present in atherosclerotic lesions. We previously demonstrated that EO6 immunostains OxLDL-associated determinants in atherosclerotic lesions from rabbits and humans (3). We have also shown that autoantibodies to OxLDL are present in lesions, in part complexed to OxLDL (16). To determine whether some of the anti-OxLDL Abs deposited in the atherosclerotic lesions of mice consist of the EO6 and/or T15 idiotype, we immunostained advanced lesions from cholesterol-fed LDLR−/− mice for the presence of Abs expressing the T15/EO6 idiotype. Figure 7a demonstrates the marked deposition of Ig in murine lesions, as reported previously (9). In Figure 7b, an adjacent section was stained with the biotinylated AB1-2 anti-T15 idiotype, which yielded the same pattern of staining depicted for total Ig deposition. An adjacent section stained with an equivalent concentration of a biotinylated isotype control did not reveal staining (Figure 7c). In aortic sections from other diseased mice, specific staining with this anti-idiotype reagent was also demonstrated, although the extent of T15 immunostaining varied in different atherosclerotic lesions (not shown). Because staining with AB1-2 recognizes a T15-specific idiotope that is independent of the associated isotype or isotypes (e.g., IgA, IgM, or IgG), these studies cannot distinguish the isotypes of the deposited T15 idiotype-bearing Ig's. Nevertheless, these studies document that T15 and EO6 Abs are specifically deposited in lesions of murine atherosclerosis.

OxLDL- and POVPC-reactive EO cell lines display highly restricted clonal heterogeneity. To evaluate their clonal heterogeneity, a PCR-based strategy was developed to characterize the genomic VH rearrangements in OxLDL- and POVPC-reactive EO cell lines, using conditions wherein nonrearranged genomic templates do not yield an amplimer product (see Methods). In previous stud-
ies, we demonstrated that EO6 cells expressed transcripts for the productive T15 V_{H} gene (S107.1-D_{FL16.1}-J_{H}1) and also a nonproductive, frame-shifted S107.13-D_{S92.2}-J_{H}3 rearrangement. Confirming these findings, amplification of the genomic DNA of EO6 cells yielded two bands, with a larger, 1,500-bp product shown by DNA sequence analysis to represent the productive T15 V_{H} gene and flanking sequence, and a smaller 800-bp product that represents the nonproductive S107 rearrangement on another allele. Significantly, the other OxLDL- or POVPC-specific cell lines, EO3, EO4, and EO7, also yielded only bands of these exact same sizes, which were shown to be identical by sequence analysis and HindIII and BstNI fingerprint patterns (data not shown). Notably, amplification of the genomic DNA of the fusion partner used to create these cell lines yielded only the smaller (800-bp) band, indicating that in the EO cell lines this band was a remnant of the fusion event, and not a marker of the rescued B cells. These data provide compelling evidence that each of these OxLDL- or POVPC-reactive EO hybridomas are genetically identical, likely arising from the same clonal set of B lymphocytes that are specific for an oxidation-associated neodeterminant.

Discussion

Natural autoantibodies are predominately IgMs that constitute a significant fraction of serum Ig’s at birth and may increase with age. Many of these Abs are encoded by nonmutated germline V_{H} or V_{L} genes (17), and they react with a variety of self-determinants, such as carbohydrates and glycolipids, and often cross-react with bacterial or oncofetal antigens to provide a fundamental tier of the adaptive immune system. The developmentally regulated processes responsible for the spontaneous production of natural autoantibodies are not clearly understood. We now report that a set of antibodies expressed by a disease-associated dominant B-cell clonal set, directed against oxidation-specific epitopes present on OxLDL and apoptotic cells, is identical to classic T15 natural autoantibodies that dominate anti-PC responses and provide optimal protection from bacterial infection from virulent S. pneumoniae (18).

We documented previously that Ab titers to oxidation-specific epitopes of OxLDL, including those of the IgM isotype, increase in parallel with the development of atherosclerosis in mice (9). Examination of the immune response occurring in atherosclerotic ApoE^{−/−} mice demonstrated that at least 32% of 1,536 viable, Ig-secret-
ers. Moreover, beginning from about day 6 of life, T15 Abs dominate in vivo responses to PC-containing immunogens (21), and they are the most protective of all anti-PC Abs to systemic infection with pneumococci (18). However, despite their importance and the large amount of investigative interest they have generated, the mechanism or mechanisms responsible for their in vivo dominance have remained a mystery, as they are highly represented even in germ-free mice (22). Therefore, the current findings are important because they identify previously unsuspected relevant “neo-self” ligands for the T15 clonal set that appear to contribute to the selection and maintenance of an important component of the host’s preformed immune defense from microbial attack.

Adoptive transfer studies have rigorously documented that T15 B cells exist predominantly (or solely) as part of the CD5+ B-1 pool (23). Although B-1 cells normally constitute only a minor fraction of peripheral lymphoid tissue (such as spleen), they represent a major fraction of B cells found in the peritoneal cavity of mice, and are the source of most natural IgMs that spontaneously arise in naive hosts without specific immunization (24). Hence, it is speculated that these primordial B cells represent evolutionarily selected Ab genes and binding specificities that dominate the repertoire expressed during early ontogeny, in part to serve housekeeping functions of removing everyday “self” debris required for maintaining homeostasis. Our findings that classical T15 Abs bind to oxidation-specific neodeterminants on both OxLDL and apoptotic cells may therefore represent the first demonstration of a defined autoreactivity by the T15 B-cell set.

These novel observations demonstrate the endogenous antigens that could be responsible for positive selection, resulting in the progressive in vivo dominance of the T15 anti-PC B-cell set.

Negative selection has long been thought to be the dominant mode of regulation of B-cell development in health, though the presence of natural autoantibodies is in seeming conflict with this theory (25). Although a positive role for self-reactivity in B-cell development has been previously considered, several recent reports have now provided firm evidence that mature B cells can be affected by receptor-mediated positive selection based on their autoreactivity. Hayakawa and colleagues (26) generated a murine transgenic model in which coexpression of an unmutated VH rearrangement encoding for self-specificity for Thy-1 glycoprotein (CD90), along with its natural self-ligand on T cells, resulted in the positive selection of the Tg B-1 cell population and enhanced serum autoantibody levels. Their important findings modify the view, based on most earlier Tg-Ig systems, that autoreactive binding of high-affinity or high-avidity determinants, constitutively expressed throughout immune maturation, results in B-cell clonal anergy or deletion. Several other recent reports also support the notion of positive antigen selection of B-1 cells (27, 28). Although these reported systems have similarities to the T15 B-1 subset, our findings are novel in that they strongly implicate B-1 cells in immune response to oxidation-specific epitopes that are linked to a pathologic state, atherosclerosis.

The binding of T15/EO6 Abs is highly specific for oxidized PAPC, and they do not bind to unoxidized PAPC, a prominent phospholipid of LDL and cells. Because

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Figure 7
Presence of T15 idiotypic Abs in atherosclerotic lesions. Atherosclerotic segments of aorta of LDLR–/– mice were prepared and stained as described in Methods. Epitopes recognized are indicated by a red color; the nuclei are counterstained with methyl green. (a) Section immunostained with a combined anti-mouse IgG/IgM antisera, indicating the presence of Ig in lesion. (b) An adjacent section immunostained with biotinylated AB1-2 (anti-T15 idiotype). A similar pattern demonstrates that some Ig deposited in the lesion represents Abs of the T15 idiotype. (c) An adjacent section stained with biotinylated nonimmune IgG was free of specific staining.
the PC headgroup is present in many phospholipids, including PAPC, and the T15/EO6 Abs do not bind native PAPC, it must be that oxidation alters the configuration or accessibility of PC so that it is able to fit into the antigen-binding site. These findings are potentially important in light of the extensive structural observations that have been made for PC-binding Abs. Crystallography studies of the hapten-binding site of a Group I anti-PC Ab similar to T15 revealed that PC occupied only a small part of the antigen-binding pocket, with the choline moiety in the interior (29). We speculate that changes to other parts of the phosphatidylcholine molecule caused by oxidation, or by formation of covalent adducts between the core aldehyde of the oxidized sn2 fatty acid and a lysine group of a protein (6), can make the PC headgroup in phosphatidylcholine accessible for T15 and EO6 binding. Alternatively (or additionally), it is possible that both the sn2 oxoaleryl group and the sn3 PC fit into the antigen-binding site simultaneously, with the sn1 hydrophobic saturated fatty acid remaining outside the antigen-binding pocket. In either case, the “true antigen” appears to be an oxidized phospholipid, such as POVPc, that is predicted to have a higher binding affinity. This would suggest that other oxidized phospholipids containing the PC headgroup might also fit into the same pocket. Similarly, we speculate that PC and perhaps other similar oxidized phospholipid moieties become expressed on OxLDL and the surface of apoptotic cells as they undergo oxidative changes known to accompany apoptosis.

The EO Abs bearing the T15 idiotype, such as EO6, have the capacity to block the uptake of OxLDL by macrophages (6). Furthermore, the epitopes recognized by EO6 are present on minimally oxidized LDL found in the circulation (3). Thus, we speculate that these Abs could affect atherosclerosis both by preventing foam-cell formation and by leading to enhanced removal from plasma of such minimally oxidized LDL, preventing their deposition in the artery wall. Similarly, these Abs might also influence the disposition of apoptotic cells (7). The observation that the same Abs also bind to C-PS, by its PC epitopes, raises the fascinating possibility that pneumococcal (or other) infections, or conceivably even vaccination, could increase the titers of T15 idiotypic Abs; this in turn could impact atherogenesis, as noted above. Obviously, extensive in vivo studies will be needed to test these speculative observations.

In this discussion we have referred to oxidation-specific epitopes generated on OxLDL and apoptotic cells as “neodeterminants,” and to the T15 and EO6 Abs that recognize them as “autoantibodies.” It is a matter of conjecture whether these are truly self antigens and autoantibodies in the classical sense. Indeed, we hypothesize that many disease-associated autoantibodies are in fact directed against subtle modifications of self proteins or lipids that have not been previously recognized. For example, we demonstrated previously that some of the so-called “anti-phospholipid autoantibodies,” found in patients with anti-phospholipid Ab syndrome, in fact bound to oxidized phospholipids but not to native phospholipids (30, 31). Based on the current findings, we speculate that other B-1 cell clones that secrete natural housekeeping autoantibodies will also be found to be specific for a variety of neo-self epitopes generated by common modifications, such as other oxidation-dependent epitopes, and enzymatic and nonenzymatic glycation.

These data suggest a new immunologic paradigm whereby immune recognition (both humoral and cell-mediated) of oxidation-specific epitopes contributes to physiologic homeostasis of apoptotic and necrotic cells, as well as senescent cells (such as erythrocytes) and lipoproteins (and similar lipid-rich membranes) that have undergone oxidative changes. This process is integral to normal physiology, because we exist in an aerobic environment. Indeed, we have recently demonstrated the presence of OxLDL even in fetal arteries (32). Furthermore, apoptosis is a normal developmental and homeostatic prerequisite. Hence, we hypothesize that neo-self oxidized phospholipids have served as an evolutionary pressure contributing to the selection of this primordial B-cell set. Under conditions of enhanced oxidative stress, as occurs in the ApoE<sup>−/−</sup> mouse with hypercholesterolemia and extensive atherosclerosis, we speculate that there is an in vivo expansion of the T15 B-1 cell set. Our data support the hypothesis that natural selection has contributed to the conservation and recurrent expression of the T15 B-cell clone during early immune development, presumably both for its housekeeping roles in health and for the advantages it provides for natural immunity from certain common infectious agents. The specific role or roles of these T15 and EO Abs in the development (or amelioration) of atherosclerosis require further elucidation.

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