Increased Mucosal B-Lymphocyte Apoptosis During Polymicrobial Sepsis Is a Fas Ligand But Not an Endotoxin-Mediated Process

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Sepsis is reported to induce an increase in the rate of apoptosis (A0), in immature lymphoid cells residing in hematopoietic tissues such as the thymus and bone marrow. Alternatively, secondary lymphoid tissue, such as the spleen exhibit little innate (unstimulated) A0. However, it is unknown whether or not polymicrobial sepsis has any effects on the frequency of A0 in mucosal lymphoid tissue and what, if any, are the functional consequences of such a change. To assess this, Peyer's patch cells were harvested from C3H/HeN (endotoxin-sensitive) mice killed 12 or 24 hours after the onset of polymicrobial sepsis (cecal ligation and puncture [CLP]). The results indicate that the percentage of cells that were A0+ as determined by flow cytometry were markedly increased at 24 hours, but not at 12 hours post-CLP. This correlates well with evidence of increased DNA fragmentation as well as histological changes observed both at a light and transmission electron microscopic level of the Peyer's patch A0. Phenotypically, these changes were restricted to the bone marrow exhibited evidence of increased B-cell A0, not observed in the splenic B cells. Whereas this may in part be explained by variation in the maturational state of these lymphocytes, it may also reflect differences in local tissue exposure to mediators and/or antigens released during polymicrobial sepsis. Inasmuch, one would speculate that lymphoid cells resident in the intestinal wall, such as Peyer's patch cells, would be exposed to increased enteric antigenic burden because studies indicate that after traumatic injury, shock, and/or sepsis, gut barrier permeability is increased. Such exposure should lead directly or indirectly to the stimulation of local mucosal lymphoid (T- and/or B-cell) tissue, which contributes to local inflammatory response. These mediators in tum may contribute to the suppression of B- and T-cell responses that have been observed after trauma, shock, and sepsis. The Peyer's patches are also the major site in which mucosal B cells ultimately become committed to IgA production. As the secretion of IgA is an important component in maintaining gut mucosal epithelia immune function, and the production of IgA is an outcome of the activation of the B cell, alterations in this response may provide insight into the ongoing immune response developed in response to a polymicrobial septic challenge. In this respect, B-lymphocyte activation (as evidenced by nuclear factors activation and subsequent IgA release) itself is associated with increased A0, as a possible autoregulatory mechanism with pathologic potential. We, therefore, hypothesize that among the many factors that may play an important role in the modulation of mucosal immunity during sepsis, is the induction of A0.

The aim of this study was to determine: (1) whether or not mucosal lymphoid cells derived from the Peyer's patches show evidence of increased A0 after polymicrobial sepsis; (2) if such changes are detected, are they associated with alterations in mucosal lymphocyte function, such as IgA production and/or nuclear factor activation; and (3) if such alterations are mediated by endotoxin and/or Fas ligand-Fas antigen (Apo-1/CD95) interaction.

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MATERIALS AND METHODS

Cecal ligation and puncture (CLP).

Male inbred C3H/HeN mice (endotoxin-sensitive) (Charles River, Portage MI), C3H/HeJ (endotoxin-tolerant), or C3H/HeJ-Fas-/- mice (endotoxin-tolerant/Fas ligand [Fasl] deficient) (Jackson Laboratory, Bar Harbor, ME) 6 to 8 weeks of age were lightly anesthetized with metofane [Methoxyflurane 2.2-Dichloro-1,1 difluoroethyl methyl ether. B.H.T. (Butylated hydroxytoluene, 0.01% w/v)] and a midline incision (1.5 to 2 cm) was made just caudal to the diaphragm, to expose the internal organs. The cecum was isolated, ligated just below the ileocecal valve, and punctured (CLP) in two places to induce sepsis. For the controls (CLP-shams), the cecum was isolated but neither ligated nor punctured. The muscle layer and epidermal layer were sutured in layers and xylocaine applied to the areas of the incision. Lactated Ringer’s Solution (0.6 mL) was administered subcutaneously in these and the sham animals.

The studies performed here were all carried out in accordance with the National Institutes of Health Guidelines on Laboratory Animals and were approved by the Rhode Island Hospital Committee on Animal Use and Care.

Peyer’s patch lymphocyte isolation. The Peyer’s patches (4 to 6 patches per mouse) were excised aseptically from the exposed small intestine of methoxyflurane-killed mice at 12 or 24 hours post-CLP or sham-CLP, then placed into petri dishes (60 x 55 mL) containing 5 mL Hank’s Balanced Salts Solution (HBSS). The Peyer’s patches were gently glass ground, then transferred into a 15 mL conical centrifuge tube, washed twice with Dulbecco’s Modified Eagle’s Medium (DMEM), and centrifuged at 400 x g for 10 to 15 minutes at room temperature. The pellet was disrupted, resuspended in 8 mL of DMEM, layered over pH 7.4 if the extent of Ao was to be determined. Samples were stored in DAPI staining reagent [0.1% Triton X-100, 0.1 mmol/L EDTA disodium salt, 0.05 mg/mL RNase A (50 units/mg), 1 µg/mL DAPI, in PBS pH 7.4] if the extent of Ao was to be determined. Samples were stored in the dark at 4°C until analysis was carried out usually within 24 hours. Isotopic controls, i.e., 1 µg of either FITC-, PE-, or Cy-Chrome–conjugated rat IgG2a or FITC-conjugated hamster IgG (Pharmingen Inc)/10⁶ cells, were included so as to allow us to assess the nonspecific antibody staining and gate this out appropriately.

For multicolor analysis, DAPI was excited with a Coherent ANOVA-70S Spectrum laser (Spectrum Products Div, Palo Alto, CA) set to 360 nm and fluorescent emission detected with a 424 ± 22 nm blue reflecting dichroic filter. Alternatively, FITC, PE, and Cy-Chrome conjugates were excited with an argon laser set at 488 nm, but detection of FITC–fluorescent emission was with a 530 ± 15 nm wideband pass filter, whereas PE emission was detected with a 575 ± 13 nm dichroic filter and Cy-Chrome detected with a 670 nm long pass filter. FITC, PE, Cy-Chrome, and/or DAPI emission overlap was corrected by electronic compensation. FITC or PE single-positive as well as double-positive and double-negative DAPI cell cycle analysis was determined after gating of cell-debris and doublets for no less then 20,000 cells/sample. By using PC-Lysis Version 1.0 software (Becton Dickinson, San Jose, CA), FITC, PE, or Cy-Chrome–positive as opposed to fluorochrome–negative cells were established based on the fluorescent emission of the nonspecific FITC/PE isotopic antibody controls with two-color fluorescent cytograms. Histograms of the regionalized cells were then produced of cell number versus DAPI stain intensity (DAPI fluorescent emission) from which the percentage of cells residing in various stages of the cell cycle could be determined.

Alternatively, after the determination of B220⁺ versus B220⁻ negative regions, as described above, histograms of the regionalized cells were then produced of cell number versus anti-Fas–FITC stain intensity (mean channel fluorescence) from which not only the percent- age of cells that were Fas positive (Fas⁺) could be determined based on the isotopic control but also the intensity of Fas antigen expression (see Results section for the results of typical two-color data for DAPI-cell cycle analysis and Fas antigen expression) could be determined.

Light and electron microscopic examination. For histologic exami- nation, Peyer’s patches were excised and fixed immediately by submersion in 4% buffered glutaraldehyde. After 1 to 3 hours of fixation, the buffered glutaraldehyde was then drawn off and the tissues were rinsed with 0.1 mol/L phosphate buffer. The specimens were then postfixed with osmium tetroxide for 1 hour, rinsed in phosphate buffer, and stained en bloc with 2% uranyl acetate. Samples were then dehydrated through a graded ethanol series followed by propylene oxide. The tissue was then infiltrated overnight in a 1:1 mixture of propylene oxide and (Polybed-Araldite) epoxy resin (Polysciences Inc, Warrington, PA). The following day, samples are infiltrated for 8 more hours in Polybed- Araldite resin after which they are embedded in fresh resin. Tissue blocks were polymerized for 2 days at approximately 74°C and then 1 µm sections were prepared with an LKB Ultratome and stained with 1% toluidine blue for examination by light microscopy. Areas selected for ultramicrotomy were sectioned at 70 to 90 nm with a diamond knife and placed on 300-mesh copper grids. Sections were stained with uranyl acetate for 1 hour and lead citrate for 2 to 3 minutes. All sections were examined at 60 kV on a Philips Model 301 transmission electron microscope (Philips Inc, Mahwah, NJ).

IgA ELISpot assay. The number of Peyer’s patch cells secreting IgA was determined by an Enzyme-linked Immunoslot (ELISpot) assay. Cells were cultured (10⁵/well) overnight on monolonal antimouse IgA (5 µg/mL; Biosource Inc, Camarillo, CA) precoated (overnight at 4°C), sterile Multiscreen-HA 96-well nitrocellulose filtration plates (Milli- pore Corp, Bedford, MA). After washing and incubation (overnight at 4°C) with a secondary biotinylated monoclonal antimouse IgA (2.5 µg/mL; Pharmingen Inc, San Diego, CA), antibody color was developed with avidin-peroxidase-H₂O₂–ACE (3-amino-9-ethylcarbazole in 0.1 mol/L sodium acetate buffer [pH 5.0]). The number of spots present in the well was determined on a Mocha Image analysis system (Jandel Scientific, Corte Madera, CA).

Assessment of B-cell nuclear expression of Rel(c-Rel and RelB)/ nuclear factor κB (NF-κB) factors by western blot analysis. The nuclear appearance (expression) of Rel/NF-κB factors, c-Rel, and RelB, as an index of the potential nature of the in vivo B-cell stimulant, was assessed by Western immunoblot analysis of the nuclear extracts of
B220+ cells harvested from the Peyer’s patches of animals 24 hours after sham-CLP operation or CLP.

B220+ cells were selected (enriched) from 1 × 10^7 Peyer’s patch cells, as previously described in our laboratory, by binding to 4 × 10^7 antimouse B220 coated/goat antirat IgG magnetic M-450 Dynabeads (Dynal, Inc, Great Neck, NY) per 5 mL and then gently stirring with a rotary blood mixer (Robbins Scientific Corp, Sunnyvale, CA) for 30 minutes at 4°C. (4 × 10^7 antimouse B220 coated/goat antirat IgG magnetic M-450 Dynabeads were created when approximately 1.5 × 10^7 Dynabeads in 10 mL of RPMI 1640 were coated overnight at 4°C with monoclonal rat antimouse B220 [Pharmingen, San Diego, CA] 10 µg/mL, and washed three times before use.) The bead-bound cells (B220+) were then concentrated by using a MCP-1 magnetic particle concentrator (Dynal, Inc) for 1 minute and the unbound (B220−) cells discarded. The cells that remained were washed one more time, magnetically concentrated, and resuspended in RPMI 1640 medium.

Typical cell yield after positive magnetic selection was approximately 4 × 10^6 cells total.

Nuclear fraction was then prepared by using a modification of the methods of Sikora et al. Briefly, the nuclear fraction of 2 × 10^7 cells was prepared by the addition of two volumes of 0.2% Nonidet P-40 in cell lysis buffer (10 mMol/L, HEPES pH 7.9, 0.2 mMol/L, EDTA, 1 mMol/L DDT, 0.5 mMol/L PMSF, and 10 mg aprotinin/mL) to the washed cell pellet that was then incubated for 10 minutes on ice. The nuclei were pelleted by centrifugation at 3,300 g for 15 minutes and the nuclear proteins extracted from the pellet by resuspension in half the volume of extraction buffer (20 mMol/L HEPES pH 7.9, 0.4 mMol/L NaCl, 0.2 mMol/L EDTA, 1 mMol/L DDT, 0.5 mMol/L PMSF, and 10 mg aprotinin/mL), incubated for 30 minutes at 4°C, aliquoted, and stored at −70°C until required for further assay.

The protein content of nuclear extract was determined fluorometrically by using the NanoOrange Protein Kit (Molecular Probes Inc, Eugene, OR). The assays were carried out in a 96-well microculture dish (Corning Scientific, Corning, NY) at a final reaction mixture volume of 250 µL and the change in fluorescence measured on a Bio-Tek FL 500 model, fluorescent plate reader (Bio-Tek Instruments, Winooski, VT).

Ten µg of nuclear extract protein (diluted in Tricine-gel sample buffer) was separated on a precast 10% to 20% Tricine-sodium dodecyl sulfate (SDS) polyacrylamide gel (Novex Experimental Technology, San Diego, CA) electrophoretically by using a Profile minigel electrophoresis system (Schleicher & Schuell, Keene, NH) with EC 105 power supply (approximately 90 mA, 70 V for approximately 1.5 hours; E-C Apparatus Corp, St. Petersburg, FL). The separated proteins were then electroblotted (by using a Profile miniblotter [Schleicher & Schuell, Keene, NH] approximately 280 mA, approximately 30 V, for approximately 1.5 hours) out of the gel onto nitrocellulose (Novex Exp Tech). The blots were rinsed in Tris-buffered saline containing 0.05% Tween-20 (TBS-Tween 20) and blocked for 2 to 4 hours with TBS-Tween 20 including 5% nonfat dry milk. Blots were then washed 4 to 5 times (15 minutes per wash), incubated (4°C) with 3 to 4 mL of primary rabbit IgG polyclonal antibody directed against either c-Rel or RelB (sc-070 and sc-226, respectively; Santa Cruz Biotech, Santa Cruz, CA), or normal rabbit IgG control polyclonal nonimmune sera (Santa Cruz Biotech) at concentrations of 3µg/mL for 4 hours at 4°C in a 50 mL polyethylene centrifuge tube gently rotated on a Robbins blood mixer (Robbins Sci Corp, Mountain View, CA). After washing the blots 4 to 5 times, they were incubated at room temperature for 1 hour with secondary polyclonal goat antirabbit IgG antibody conjugated to alkaline phosphatase (1:1000; Santa Cruz Biotech). Finally, after 4 to 5 washings the blots were developed in alkaline phosphatase substrate buffer (100 mMol/L Tris, 100 mMol/L NaCl, 50 mMol/L MgCl_2; pH 9.5) containing 330 µg nitroblue-tetrazolium and 168 µg 5-bromo-4-chloro-3-indolyl phosphate/mL for 45 minutes. Molecular weight (kD) of the proteins of interest was determined by comparison with low molecular weight standards included in each gel. To quantify the differences present in various nuclear extract samples, the band intensities were assessed densitometrically by using a Mocha Image Analysis/SigmaGel work station (Jandel Scientific, Corte Madera, CA).

**Qualitative assessment of internucleosomal DNA fragmentation by gel electrophoresis.** After washing, 3 × 10^6 total Peyer’s patch cells were extracted for genomic DNA by 10 minute lysis in 20 mMol/L Tris-HCl, pH 7.4, containing 10 mMol/L EDTA, and 0.2% Triton-X100. After removal of cell debris the supernatant was incubated overnight with Proteinase K (100 µg/mL, 50°C). The DNA was then chloroform/phenol extracted and precipitated by using isopropanol (overnight, −20°C). After centrifugation, the pellet was dried in a SpeedVac Plus (Savant Instrument Inc, Farmingdale, NY).

Genomic DNA was dissolved in 25 µL Tris-EDTA buffer containing 1 µg RNase/mL and incubated for 1 hour (37°C). Five µL of 6 × TBE sample buffer was then added to each DNA sample, heated (65°C, 10 minutes) and then 15 µL applied to precast TBE 4% to 20% polyacrylamide gel (Novex Experimental Technology). The samples and standard (Lambda-DNA HindIII) were then electrophoresed for 90 minutes (90 to 100 V), stained with ethidium bromide, rinsed, visualized under UV light, and documented with Polaroid film (Polaroid Corp, Cambridge, CA).

**Presentation of data and statistical analysis.** The data are presented as a mean and SE of the mean for each group. Differences for total viable cell yields were considered to be significant if P < .05, as determined by ANOVA (Spectrum Products Div) and a Tukey’s test, was applied for multiple comparison. Differences in percentile data (ie, percentage of A c+) were considered to be significant if P < .05, as determined by using the Mann-Whitney U test.

**RESULTS**

**Polymicrobial sepsis induces a marked increase in the frequency of A c detected in Peyer’s patches.** Figure 1A illustrates that a small but statistically significant decline in total viable Peyer’s patch cell yield from endotoxin-sensitive C3H/HeN mice was detected at 24 hours but not 12 hours after CLP. Alternatively, at 24 hours but not 12 hours, a significant increase in the frequency of A c was evident as detected by hypoploid DAPI DNA staining (Fig 1B). Evidence of endogenous endonuclease activity, as detected by the fragmentation of genomic DNA, was also observed in the DNA obtained from the septic-mouse Peyer’s patch cells harvested at 24 hours post-CLP (Fig 2).

To determine whether the alterations observed in the ex vivo isolated cells represented in vivo changes at the tissue level, we examined the Peyer’s patches harvested from C3H/HeN mice at both a light and electron microscopic level for evidence of A c. Figure 3A and C are representative samples taken from a 24-hour postsham-operated mouse that exhibited typical Peyer’s patch morphology. However, Peyer’s patches taken from mice 24 hours after onset of polymicrobial sepsis showed a marked change in morphology (Fig 3B). With respect to the septic mouse Peyer’s patches, the majority of the apoptotic cells appear to be clustered in the germinal center and not in the T-cell region localized more proximal to the mucosal epithelial layer. Closer inspection of these apoptotic clusters on an electronic microscopic level revealed karyorrhectic cells with condensed nuclei and what appear to be apoptotic bodies (Fig 3D).
Increased endogenous Ao detected in septic mouse Peyer’s patch cells is restricted to cells of the B- but not T-cell lineage. In an effort to delineate the population of C3H/HeN mouse Peyer’s patch cells undergoing Ao after the onset of sepsis, cells were concomitantly stained with the nonspecific murine B-cell marker B220 (CD45) and the DNA dye DAPI.

Figure 4A through H illustrates the results of typical two-color flow cytometric analyses of cell suspensions obtained from the Peyer’s patch 24 hours after sham-CLP (sham) or CLP (simultaneously stained and analyzed). The contour plots in Fig 4A and B show the typical cell cycle progression for sham or CLP-mouse Peyer’s patch cells and the primary region (R1) established around these populations. In Fig 4C and D, the cell cycle histograms of cell number versus DNA content generated from the R1 region in Fig 4A and B are shown. By using the sham, the histogram was divided into two subregions (M1 = apoptotic [Ao+] cell cycle region and M2 = which includes G0/G1, S, and G2/M cell cycle region, respectively). These regions were kept consistent for all samples analyzed in a given experiment. With respect to the sham, it can be observed that Peyer’s patch cells extracted from a septic mouse at 24 hours exhibited a population of cells tailing off towards the lower left-hand corner with lower DNA content from the G0/G1 peak (Fig 4A and B). This is reflected by the increased percentage of apoptotic cells detected in the M1 region of the histogram (Fig 4C and D). By using a rat IgG-PE control, B220+ cells were discriminated from B220− cells as depicted in Fig 4E and F and then cell cycle histograms were produced (Fig 4G and H). Cell cycle histograms of DNA content produced from each of these gated phenotypic populations illustrate that the majority of the increase in apoptotic cells was typically present in the B220+− stained cell population (Fig 4G and H).

Figure 5 depicts the summary of the data from no less than six independent animals. Peyer’s patch samples were examined in each group. Although a small decline in the percentage of cells that are B220+ was observed in the CLP mice, this was not statistically significant (Fig 5A). Alternatively, there was a marked increase in the percentage of cells that were apoptotic in septic mice, as compared with sham mice, as determined by cell cycle analysis in the population of cells that were B220+ (Fig 5B). No similar change was evident in the B220− cell population. Although not shown, combined staining with antibody against the T helper cell marker CD4 and DAPI did not show evidence of an increased frequency of Ao in this cell population (data not shown).

Polymicrobial sepsis induces an increase in the number of Peyer’s patch B cells that are secreting IgA. To determine the potential functional significance of the increase in B-lymphocyte Ao during sepsis, the ability of cells harvested from these animals to secret IgA was determined by using an ELISpot method. The result in Fig 6 indicates that septic C3H/HeN...
mouse Peyer’s patch cells exhibited a marked increase in the number of cells secreting IgA.

Polymicrobial sepsis induces an increase in the nuclear expression of c-Rel but not RelB in Peyer’s patch B cells. Because recent studies indicate that mature B-cell activation through the B-cell (Ig) receptor induces a marked increase in the nuclear translocation/activation of the c-Rel component of the Rel family of NF-κB factors, as opposed to stimulants such as endotoxin (ie, lipopolysaccharide) or CD40 ligand, which appear to activate alternative components, such as RelB, we thought it should be possible to obtain indirect insight as to the nature of the in vivo B-lymphocyte stimulant acting on these cells after CLP. The results of the western immunoblot analysis of the B220+ cell nuclear extracts documented an increase in expression of c-Rel (approximately 75 kD) in septic mice relative to sham (Fig 7A and B). Alternatively, it was observed that RelB (approximately 68 kD) did not appear to differ in its expression between sham and CLP mouse samples. Although not shown, no staining in the molecular weight range of either c-Rel or RelB with the control rabbit sera provided for these antibodies was seen.

The increased ΐA detected in Peyer’s patch cells during sepsis is Fas-ligand but not an endotoxin-mediated process. Because recent studies have indicated that lymphocyte activation is associated with accelerated ΐA that may be mediated via the cell surface antigen known as Fas antigen and its associate ligand, we performed experiments to assess the expression of this antigen in the Peyer’s patch cells at 12 and 24 hours after the onset of polymicrobial sepsis.

Figure 8A through F illustrates the results of typical two-color flow cytometric analyses of cell suspensions obtained from the C3H/HeN mice Peyer’s patch 24 hours after sham-CLP (sham) or CLP concomitantly stained and analyzed for Fas antigen and B220. The histograms in Figure 8A and B show the typical Fas antigen staining pattern for sham- or CLP-mouse Peyer’s patch cells over which the hamster IgG-FITC control antibody staining was superimposed. By using the isotype control the histograms were divided into two subregions (M1 =
These regions were kept consistent for all samples analyzed in a given experiment. With respect to the sham, it can be observed that the mixed Peyer’s patch cells extracted from a septic mouse at 24 hours exhibited a population of cells that are shifted to the right resulting in an increase in the percentage of Fas$^{+}$ cells (Fig 8A and B, as well as summary data for repeated experiments provided in Fig 9A and B). By using a rat IgG-PE control, B220$^{+}$ cells were again discriminated from B220$^{-}$ cells as depicted in Fig 8C and D and then the percentage of cells that were Fas$^{+}$ was again determined as depicted in the histograms in Fig 8E and F. These histograms of Fas$^{+}$ staining produced from each of the gated phenotypic populations show that the majority of the increase in Fas$^{+}$ cells was restricted to the B220$^{+}$ stained cell population (Fig 8E and F, as well as summary data for repeated experiments provided in Fig 10).
Summary data of no less than six animal’s cells samples in each group were presented in Fig 9A and B. At 12-hours post-CLP, no marked change in Fas antigen expression was evident in the mixed Peyer’s patch cell population. However, by 24 hours a significant increase in the percentage of cells that were positive for Fas antigen on septic mouse cells was evident. Also although Fas antigen mean channel fluorescence trended towards an increase in these septic mouse cells, it was not statistically significant. Additionally, the percentage of Fas antigen-positive cells was determined as a component of B220 antigen expression. The results indicated that the increase in the percentage of Fas antigen positive cells observed in the mixed septic mouse Peyer’s patch cells at 24 hours was primarily caused by a marked increase in the percentage of cells that were B220\(^+\), which were expressing Fas antigen and were not caused by changes in the B220\(^-\) population (Fig 10B). In this respect, no increase in the percentage of Fas antigen positive cells on cells expressing the T-cell phenotype, CD4 was observed (data not shown).

A series of experiments were performed to determine if the increase in A\(_o\) C3H/HeN mouse Peyer’s patch B220\(^+\) cells was due to either endotoxin (ETX; a product of gram-negative bacteria observed in CLP mice) and/or the Fas ligand.\(^{18,19}\) Figure 11A illustrates that only in the Peyer’s patch cells harvested from endotoxin-tolerant C3H/HeJ1\(^{16,17}\) septic mice was a marked decline in viable cell yield detected. Alternatively, although there was a slight decline in the viable cell yield from the Fasl-deficient C3H/HeJ-Fas\(^{\text{ld}}\) septic mice, it was not statistically significant. Figure 11B shows that only the cells obtained from CLP C3H/HeJ mouse and not the C3H/HeJ-Fas\(^{\text{ld}}\) mice exhibited increase in the percentage of A\(_o\)\(^+\) cells. This increase in the percentage of A\(_o\)\(^+\) was also associated with a significant increase in the percentage of Fas\(^+\) cells (Fig 11C). Here again, the enhancement in percentage of Fas\(^+\) cells was restricted to the B220\(^+\) cell population and was not a response to changes in the B220\(^-\) population (Fig 12).

DISCUSSION

The results presented here show that mucosal lymphocytes obtained from Peyer’s patches undergo a marked induction of Ao during sepsis. This is supported by evidence of augmented endonuclease activity in the septic animal’s Peyer’s patch cells (as detected by both flow cytometry or TBE-PAGE, as well as by the increased presence of morphologically apoptotic cells detected in the septic mouse Peyer’s patch germinal centers). The majority of these changes appear to be restricted to cells of the B-cell lineage (B220\(^+\)). It could be argued that this increase in apoptotic frequency (from approximately 3.5% in the sham to approximately 7.3% in CLP) might not be sufficient to account for the approximately 2/3 reduction in Peyer’s patch viable cell yield. However, it should be noted that the clearance of apoptotic cells is a dynamic process in which macrophages are
constantly involved in the removal of nonviable cells. Hence, the accumulation of apoptotic cells at any one time point in vivo would not necessarily have to be large. The increase in A₀ detected in these septic mice appears to be associated with an augmented endogenous IgA secretory response. We would speculate that this is in part a response to altered exposure to enteric antigens released due to increased gut permeability observed during sepsis. Inasmuch, this increase of A₀ in Peyer’s Patch B cells would appear to be an example of activation-induced lymphocyte A₀.

With respect to the kinetics of the apoptotic response exhibited here in the Peyer’s patch B220⁺ lymph node cell population after CLP, we observed that increased A₀ was not evident until 24 hours and not at 12 hours after the onset of polymicrobial sepsis. A similar observation was made with respect to viable cell yield. Interestingly, although the delay in the expression of increased A₀ in the Peyer’s patch cells differs from our previous observations made with respect to cells of the thymus, in which evidence of an increased frequency of A₀ was detectable as early as 4 hours post-CLP; these findings are comparable to our findings made with respect to bone marrow B-lymphocyte A₀, in which only late (24-hours post-CLP) after the onset of sepsis was an increase in apoptotic frequency observed. However, these results differ from splenic B cells in which no detectable increase in innate A₀ was observed up.
through 24-hours post-CLP. This supports the hypothesis that immune cells of a given lineage, be they T cells,4 macrophages,30 and/or granulocytes,31 respond differently with respect to the induction of apoptotic processes in response to polymicrobial sepsis. This is due not only to differences in their respective microenvironments, but also to their ontological/differentiation status, as well as variation in the local concentration and/or form of potential antigens and/or mediators capable of altering the apoptotic process. In this respect, the increased apoptotic response to mucosal B cells, such as Peyer's patch cells examined here, as compared to splenic B cells lack of A0, we speculate is a reflection of the difference in their exposure to enteric antigens moving across the mucosal cell barrier. It has been well documented that gut barrier function is reduced after trauma, shock, and/or sepsis that would lead to a marked influx of such enteric antigens.32,33 To the extent that the finding of augmented B-cell A0 might be a peculiarity of Peyer's patches, we have recently observed in preliminary studies that B220+ lymphocyte subset of the lamina propria also exhibit a marked increase in A0 during sepsis (24 hours; sham, 30.9 ± 2.3 v CLP; 40.0% > 2.0% A0, p < .05, n = 6/group). To the extent that this is a reflection of the teleological linkage between which is thought to exist in the development of a mucosal B-cell IgA-mediated response (antigenic stimulation of Peyer's patch B cells [commitment to IgA production] leading to differentiation, migration, through mesenteric lymph nodes and eventually to the lamina propria)18 or a reflection of a global nonspecific cellular intestinal response, must still be determined. However, we have not as yet assessed IgA production in the lamina propria B-cell population, it is unknown if this increased A0 is comparatively associated with activation of IgA in Peyer’s patch cells.

The enhanced expression of Fas antigen is in line with the onset of activation-induced B-cell A0.29 However, the mechanism underlying the induction of Fas antigen expression in this cell population during sepsis remains unknown. Fasl has been shown to act as an inducer of its own receptor’s expression19 in T lymphocytes whereby it acts to mediate lymphocyte activation-induced A0.30,31,32 In this respect, we attempted to determine the contribution of Fasl-Fas antigen interaction to the enhanced B220+ cell population A0 detected in septic mouse Peyer’s patch cells obtained from C3H/HeN-Fasl0/0 mice subjected to either sham-CLP or CLP 24 hours earlier. This mouse strain contains a point mutation that abolishes the capacity of Fasl to bind Fas receptor/antigen.19,30 By using Fasl-deficient C3H/HeN-Fasl0/0 mice it was found that CLP did not markedly enhance B-cell A0 in these animals’ cells. These findings indicate that Fasl regulates not only the upregulation of Fas antigen observed here but also the induction of A0. However, the source of Fasl that might control this mucosal B-cell process is less clear. B cells are not a typical source/producer of Fasl,33 most likely ruling out autocrine mediation of the increased expression of Fas antigen in these cells. Alternatively, paracrine release of Fasl by Peyer’s patch T cells and/or nonlymphoid cells might mediate the increased expression of B-cell Fas in septic mice. In this respect, although T cells are a documented source of Fasl,26,34,35,36 we did not detect increased Fas antigen/receptor expression on the T-helper cell population. This indirectly suggests that T-cell Fasl expression/release is not markedly augmented in the Peyer’s patch. This, however, remains to be established by direct measurement of either cell surface Fasl expression and/or its release.

Studies have also suggested that the stimulation of CD40 antigen on the surface of the B cell during activation acts to upregulate the expression of Fas antigen making the activated B cell more susceptible to Fasl-mediated apoptosis in cell death.13,14 Here again the B cell is not the primary source of CD40 ligand and the activated T cell appears to provide this stimulant.13,14 However, because the expression of CD40 antigen or CD40 ligand were not assessed it is unknown whether either of these costimulatory components are absent in the Peyer’s patch cells from septic mice.

With respect to other mediators that might be implicated in the induction of A0 in polymicrobial septic mouse lymphocytes, studies by Zhang et al38 reported that thymic A0 could be induced by the exposure of mice to endotoxin, a component of the cell wall of gram-negative bacteria (that is also a polyclonal B-cell activator) commonly associated with systemic infection associated with sepsis. However, the dosage of lipopolysaccharide required to induce marked A0 was ≥ 50 µg/mouse (ip, 18 hours; equivalent to 2.5 mg/kg/body weight). This dose of LPS typically produces blood levels of endotoxin that are significantly higher than the circulating levels of this agent detected in septic patients.39-41 It should also be noted that the model of CLP used in this study induces the release of low levels of circulating endotoxin (range 0.07 to 5 ng/mL serum) detectable as early as 1 hour post-CLP, which steadily increases over the first 24 hours thereafter to a maximum of approximately 60 ng/mL serum.15,42,43 This model of sepsis is also polymicrobial in nature as it is associated with not only gram-negative but also gram-positive bacteria. Nonetheless, because studies have suggested the role of endotoxin in the induction of thymic A0,44 we examined the potential contribution of endotoxin in increasing A0 in Peyer’s patch lymphocytes after sepsis,15,42,45,46 In this respect, the data we have presented here by using the endotoxin-resistant C3H/HeJ mouse strain16,17 show that the frequency of
A_0 is not only increased in 24-hour CLP mice when compared with sham, but is comparable to that observed in cells from septic endotoxin-sensitive C3H/HeN mouse strain. These results would imply that endotoxin is not a primary agent responsible for the induction of A_0 in the Peyer’s patch. Furthermore, this also implies that the presence of organisms not only of a gram-negative origin, but also gram-positive bacteria and/or fungal agents may be required to induce the enhanced A_0 observed here.

The present results also show that polymicrobial sepsis and not the stress associated with trauma, in the form of the midline laparotomy, produces a marked increase in the frequency of A_0 in Peyer’s patch lymphocytes.

The degree to which other inflammatory mediators released during sepsis, such as cytokines like tumor necrosis factor, transforming growth factor-β, lymphokines, prostanooids, steroids, etc, play a role in the induction of mucosal B-cell A_0 remains to be examined.

With respect to the functional significance of the enhanced A_0 observed in the Peyer’s patch B-lymphocytic population, this appears to be a reflection of an antigen-specific response to the influx of enteric antigens and not the response to nonspecific polyclonal mitogenic stimulation, such as might be mediated by endotoxin. This conclusion is based not only on our observation that A_0 is still enhanced in septic endotoxin-tolerant mouse Peyer’s patch cells but also on the ex vivo assessment of the nuclear factor footprint that is present in the cells obtained from the septic-mouse Peyer’s patch. Neumann et al recently reported that the activation of individual members of the nuclear factor-κB (NF-κB) complex (eg, RelB and c-Rel) are differentially affected (with respect to their translocation to the nucleus and/or their transcription) by various B-cell stimuli. They reported that the translocation and transcription of RelB was markedly enhanced in cells that had been stimulated through the CD40 receptor as opposed to either the receptor for endotoxin (ie, lipopolysaccharide) or the Ig receptor of the B-cell receptor complex (which is evident but weak). Alternatively, stimulation through the Ig receptor of the B-cell receptor complex induced marked translocation of c-Rel to the nucleus, but only a weak activation/translocation when stimulated through the CD40 antigen and no marked translocation of c-Rel was observed with endotoxin. Such a footprint of NF-κB nuclear translocation may provide some insight into the nature of the stimulant responsible for the induction of IgA secretion in the septic mouse Peyer’s patch cells and the associated increase in activation-induced A_0 observed here. The results of the western immunoblot analysis of nuclear protein extracts obtained from these septic animals supports the suggestion that the B-cell activation observed here is most likely Ig receptor-mediated, hence a specific response to enteric antigens, and does not appear to be due to nonspecific polyclonal B-cell activation by agents like endotoxin. However, although we speculate here that such a relationship exists, direct linkage between increased B-cell IgA release, increased c-Rel translocation, and the eventual induction of A_0 remains to be determined.

In summary, our findings indicate that mucosal B-cell A_0 appears to be markedly increased by polymicrobial sepsis and that this appears to be the result of a FasL-mediated process. Evidence of antigen-specific polyclonal activation suggests the possible induction of B-cell A_0 and eventual germinal center B-cell functional impairment.

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Increased Mucosal B-Lymphocyte Apoptosis During Polymicrobial Sepsis Is a Fas Ligand But Not an Endotoxin-Mediated Process

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