Independent Regulation of DNA Recombination and Immunoglobulin (Ig) Secretion during Isotype Switching to IgG1 and IgE

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Summary

Induction of switch recombination to the γ1 and ε immunoglobulin (Ig) heavy chain loci was examined in B cells preactivated with anti-Ig (B lymphoblasts). In B lymphoblasts cultured with interleukin 4 (IL-4), IL-5 induced the accumulation of μ-γ1 rearrangements, but not ε recombination. Thus, IL-5 facilitates switch recombination directed to the γ1 heavy chain locus by IL-4, but additional signals are required to drive rearrangements to ε. Lipopolysaccharide (LPS), in the presence of IL-4, induced the accumulation of both μ-γ1 and μ-ε rearrangements, and cells treated with LPS exhibited 40-50-fold more μ-γ1 rearrangements than cells cultured with IL-5. Induction of switch recombination was not always associated with secretion of the respective Ig isotype, since concentrations of IL-4 that were sufficient to direct switch recombination to γ1 and ε in blasts treated with LPS failed to elicit secretion of IgG1 and IgE. These results demonstrate differential requirements for switch recombination to the γ1 and ε loci, as well as independent regulation of Ig gene rearrangement and secretion of each isotype.

Cytokines produced by activated T cells influence the pattern of Ig isotype expression in B lymphocytes (for a review see reference 1). For example, IL-4 stimulates the production of IgG1 and IgE in murine B lymphocytes and induces IgG4 and IgE secretion from human B cells (2-7). IL-4 stimulates transcription of the unrearranged γ1 and ε heavy chain loci resulting in the accumulation of germline γ1 and ε transcripts (8-12). Thus, IL-4 may direct switch recombination to γ1 and ε heavy chain loci by regulating switch region accessibility to a recombinase, a putative enzyme complex that mediates DNA rearrangement within the Ig heavy chain locus. Although IL-4 alone is sufficient to induce accumulation of germline transcripts in B cells, signals in addition to IL-4 are required for expression of IgG and IgE isotypes (1). In murine B cells, LPS provides a T cell-independent signal for isotype switching (2-5). B lymphocytes cultured with LPS and IL-4 exhibit μ-γ1 rearrangements on both the active and dormant alleles (13, 14), suggesting that LPS promotes switch recombination directed to the γ1 heavy chain locus by IL-4. However, in B cells activated with anti-Ig, LPS is a poor stimulus for accumulation of γ1 and ε heavy chain mRNA and secretion of IgG1 and IgE (15). Therefore, LPS does not provide all the signals necessary for robust production of these proteins.

T cells provide costimulatory signals for isotype switching by direct physical contact with B lymphocytes and by the secretion of cytokines such as IL-5 (1, 16). In conjunction with IL-4, IL-5 promotes secretion of IgG1 and IgE from B cells activated with anti-Ig (15), and IgG1 production from cells cultured with CD4+ T cell clones (17) or with membranes isolated from activated T cells (18). In contrast to LPS, IL-5 induces marked accumulation of γ1 and ε heavy chain mRNA (15, 19). Thus, IL-5 acts in concert with IL-4 to promote expression of the γ1 and ε heavy chains, but it is not clear whether IL-5 facilitates switch recombination to targeted IgH loci or acts subsequent to IgH rearrangement.

To clarify the mechanism(s) by which IL-5 promotes expression of IgG1 and IgE, we have directly examined switch recombination to the γ1 and ε heavy chain loci in anti-Ig-activated B cell blasts (B lymphoblasts). Although LPS (with moderate concentrations of IL-4) failed to stimulate secretion of IgG1 and IgE, we show that LPS induced marked accumulation of μ-γ1 and μ-ε rearrangements, indicating that switch recombination to γ1 and ε can be dissociated from secretion of IgG1 and IgE. Conversely, IL-5 stimulated marked secretion of IgG1, but was less effective than LPS as a stimulus for recombination to γ1 and failed to induce ε recombination. These results demonstrate that IL-5 regulates recombination to Ig loci targeted by IL-4 as well as expression of the rearranged γ1 and ε heavy chain loci subsequent to recombination.

Materials and Methods

Mice. Female BALB/c mice were obtained from Charles River Laboratories (Wilmington, MA) and used at 10-14 wk of age. Reagents. Affinity-purified antibodies were obtained from Jack-
son ImmunoResearch Laboratories, Inc. (West Grove, PA; goat anti-mouse IgM plus IgG, goat anti-mouse IgG, Fc fragment), Fisher Biotech (Pittsburgh, PA; biotinylated goat anti-mouse IgG1, mouse IgG1-x standard), and Pharmingen (San Diego, CA; monoclonal rat anti-mouse Thy-1.2 and rat anti-mouse-CD4, biotinylated monoclonal rat anti-mouse IgE, mouse IgE-x standard). ELISA substrate p-nitrophenyl phosphate and *Salmonella typhosa* LPS were purchased from Sigma Chemical Co. (St. Louis, MO). Alkaline phosphatase-conjugated avidin was obtained from Fisher Biotech. Recombinant murine IL-4 and rIL-5 were prepared using the Baculovirus expression system and the activity of S9 cell supernatants was determined as described (15).

**Cell Culture.** Spleen cell suspensions were enriched for B lymphocytes by incubation with anti-Thy-1.2 and anti-CD4 followed by lysis with baby rabbit serum (Pel-Freeze Biologicals, Rogers, AR). High density B cells (1.081–1.086 g/ml) were isolated on discontinuous Percoll density gradients as described (20). Low density B cell blasts (anti-lg blasts or B lymphoblasts) were prepared by culturing high density B cells with anti-Ig- Sepharose (0.5 ml of 10% Sepharose/10 ml RPMI 1640, 5% FCS, 5 μg/ml gentamicin, and 50 μM 2-ME) at 1–1.5 × 10⁶ cells/ml for 2 d. At the end of the 2-d primary culture with anti-lg, B lymphoblasts were either recultured (along with Sepharose beads) in 96-well microtiter plates at 2 × 10⁵/ml (lg secretion) or 4 × 10⁵/ml (MTT assay) with lymphokines and LPS, or secondary stimuli were added directly to bulk cultures with anti-Ig-Sepharose for experiments requiring isolation of genomic DNA.

**Proliferation and Ig Secretion Assay.** B lymphoblast growth and viability was assessed in microcultures using the MTT [(3[4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] assay of Mosmann (21). Levels of IgG1 and IgE in culture supernatants were determined by solid phase ELISA as described (15) except that biotinylated rat anti-mouse IgE was used as the secondary antibody in the IgE ELISA and NP-40 was not included in the ELISA buffer.

**Isolation of Genomic DNA.** B lymphoblasts (0.5–1 × 10⁶) cultured with LPS or lymphokines for 2–4 d were washed with PBS and lysed by resuspension in 0.6 ml of TNE buffer (1% SDS, 0.1 M EDTA, pH 8.0, 50 mM Tris, pH 8.0, 0.1 M NaCl). Cell lysates were digested with proteinase K (200 μg) for 16 h at 35°C, followed by digestion with 20 μg/ml RNase A for 1–2 h at 37°C. Genomic DNA was sequentially extracted with an equal volume of phenol, phenol/chloroform (50% vol/vol) followed by chloroform (100%), and then precipitated with an equal volume of isopropanol followed by a single wash with 80% ethanol. Genomic DNA was quantitated by UV spectrophotometry (1 OD A₂₆₀ = 50 μg/ml).

**Inverse PCR Assay.** Genomic DNA (20 μg) was digested with an initial aliquot of XbaI (Boehringer Mannheim, Indianapolis, IN) (2–3 U/μg) for 2–4 h at 37°C, after which a second aliquot was added and allowed to digest overnight at 37°C. After the overnight incubation, a third aliquot of XbaI (2–3 U/μg) was added for an additional 2–4 h. The XbaI enzyme was inactivated at 65°C for 15 min, and protein was removed from digested DNA by either phenol/chloroform (50:50% vol/vol), followed by chloroform extraction, or by centrifugation through polyvinylidene fluoride filters (Integrated Separation Systems, Natick, MA). The XbaI digests were precipitated with ethanol and quantitated by UV spectrophotometry.

XbaI-digested DNA (2 μg/ml) was circularized by incubation with 0.02 Weiss U/μl T4 DNA ligase (Boehringer Mannheim) overnight at 15°C. Aliquots from ligation reactions containing 10–150 ng of DNA were placed directly into PCR reaction tubes and denatured by heating at 94°C for 30 min and then held at 4°C while a reaction mixture containing 50 mM KCl, 10 mM Tris-HCl, pH 8.8, at 25°C, 1.5 mM MgCl₂, 1% Triton X-100, 200 μM each dNTP, 2 μM each primer, and 2.5 U Taq DNA Polymerase (Promega Corp., Madison, WI or Boehringer Mannheim) was added to a final volume of 50 μl and overlaid with mineral oil (Sigma Chemical Co.). Amplification was performed using a DNA thermal cycler (Perkin Elmer Cetus; Norwalk, CT) and the following cycling parameters: 20–40 cycles of denaturation at 94°C for 1 min, annealing at 65°C for 1 min, and extension at 72°C for 2 min. For detection of Sµ-Sε rearrangements, the following modifications were employed: (a) the annealing temperature was increased to 68°C; (b) the total reaction volume was increased to 100 μl; (c) the PCR reaction mixture was added to denatured DNA samples held at 80°C; and (d) 2 U/100 μl of Perfect Match™ Polymerase Enhancer (Stratagene, La Jolla, CA) was included in the reaction mixture.

**Primers for PCR.** DNA oligomers were purchased from Operon Technol., Inc. (Alameda, CA). The Sµ(γ1), 5′-TACA CAG AGC ATG TGG ACT GGT C-3′, and the Sµ(e), 5′-CTG AGA TAG AGA AGG GTC TCC TGG GTA GG-3′ oligomers correspond to nucleotides 3970-3946 or 4502-4478, respectively, of the antisense strand of the GenBank code MUSIGCD07 (sequences are available from EMBL/GenBank/DDJB under accession numbers J00440, J00480, and V01524). These oligomers prime synthesis of the antisense strand across the XbaI site (position 3868) immediately 5′ to the Sµ region. The Sγ1 oligomer 5′-GAG AGC AGG GTC TCC TGG GTA GG-3′, which corresponds to nucleotides 8893-8915 of the MUSIGHANB sequence (accession number M12398), primes synthesis of the sense strand across the XbaI site located 3′ of the Sγ1 region. The Sγ1 primer anneals to a sequence located 51 nucleotides upstream of the EcoRI site within the 3′ portion of the Sγ1 region (22). The SeXba4 primer, 5′-GAT GAA AGT CAT TGG GTA CCA G-3′, corresponds to nucleotides 2382-2406 of the MUSIGHAX sequence (accession numbers M57385 and X53677). The SeXba4 primer anneals to sequence immediately 5′ to the XbaI cleavage site at nucleotide 2408. The 5′-globin-XbaI oligomer, 5′-TGA CCA TTA GAT CTG CCT ACA G-3′ (nucleotides 14876-14852 of the MUSBGCDX sequence; accession numbers M36857, X14061, M27616, and M36856) primes the antisense strand 201 bases 3′ to the XbaI site at position 14675. The Sγ3-Xba4 primer 5′-ATC TCT TGT CAA GGC TCT GGA TCA G-3′ (nucleotides 17205-17229 of the MUSBGCDX sequence), primes the sense strand 388 bases 5′ to the XbaI site at nucleotide 17593.

**Southern Analysis.** Products of PCR amplification were electrophoresed through 1% agarose gels. DNA in agarose gels was denatured by incubation in 1.5 M NaCl, 0.5 M NaOH for 30 min with agitation, and neutralized by incubation in 1.5 M NaCl, 0.5 M Tris, pH 6.0, for an additional 30 min. DNA was then transferred to nylon membranes (Stratagene) (23), cross-linked to the filters using the UV Stratalinker™ 2400 (Stratagene), and hybridized to 3²P-labeled DNA probes (see below). Hybridizations were carried out at 68°C for 1–4 h using QuikHyb™ solution (Stratagene), and filters were washed according to the protocol recommended by the manufacturer. After air drying, filters were exposed to Kodak X-O MAT AR film for 1–6 h at –70°C with an enhancer screen. The migration of HaelII-digested PhiX174 molecular weight markers (New England Biolabs Inc., Beverly, MA) were used to verify the length of PCR products.

**DNA Probes.** The β-globin probe was composed of a 336-bp PCR product amplified using 3′-globin-XbaI oligomer and the 3′-globin primer 5′-GCA TTT CTT CTC TTC TGG ACA GGT C-3′ (nucleotides 17541-17517 of GenBank code MUSBGCDX), ligated
to a 463-bp product amplified using the 5' globin primer, 5'-TGA TG TCCA GCA TCT TGT ACA CCC-3' (nucleotides 14413-14436 of GenBank code MUSBGCXD), and the 5' globin-XbaI primer. The PCR-generated globin fragments described above were joined by PCR amplification to form a 799-bp $\beta$-globin probe using the 5' and 3' globin-XbaI primers. The PCR-generated globin fragments were cloned into the pCR™ II TA cloning vector (Invitrogen, San Diego, CA). Insert DNA was released from the vector by digestion with Kpnl and ApaI, or PCR amplified from vector DNA using appropriate primers. DNA fragments in low melt agarose were 3'Xlabeled using 3,000 Ci/mmole $\alpha$-$^32$P]dCTP (NEN/Dupont, Wilmington, DE) and the Prime-it II Random Primer Kit (Stratagene) according to protocols provided by the manufacturer.

Results

Differential Effects of IL-5 and LPS on B Lymphoblast Proliferation and Ig Isotype Production. The capacity of IL-5 and LPS to stimulate proliferation as well as IgG1 and IgE production by IL-4-treated blasts was compared by culturing B lymphoblasts with IL-4 (20-2,000 U/ml) and either LPS or IL-5. B lymphoblasts proliferated extensively when cultured with LPS or LPS plus IL-4 (Fig. 1) but were apparently unresponsive to IL-5 in isolation. The combination of IL-4 and IL-5 caused limited B lymphoblast proliferation (determined by the number of cells recovered at the end of 2- and 3-d cultures, data not shown) and enhanced long-term viability of cells in culture as measured by the MTT assay (Fig. 1).

Previous experiments indicated that secretion of IgG1 and IgE were differentially regulated (15). This is illustrated in Fig. 1 where culture of B lymphoblasts with optimal concentrations of IL-4 and IL-5 resulted in marked secretion of IgG1 but no detectable IgE even at supraoptimal concentrations of IL-4 (2,000 U/ml). Inclusion of LPS in cultures with IL-4 promoted both IgG1 and IgE secretion, but high concentrations of IL-4 (2,000 U/ml) were required to elicit the secretion of either isotype in the absence of IL-5. Virtually no IgG1 or IgE secretion was observed in cultures with LPS, IL-4, or IL-5 alone. These results confirm previous observations (15) demonstrating that LPS and IL-5 can each promote IL-4-dependent IgG1 production, but only LPS appears capable of inducing IgE production.

A Semi-quantitative Assay for Switch Recombination Based on Inverse PCR. To better understand the mechanisms by which LPS and IL-5 regulate Ig isotype expression, we developed a semi-quantitative assay for $\mu$-Sy1 and $\mu$-Se recombinations based on the technique reported by Chu et al. (25). $\mu$-Sy1 recombination juxtaposes two XbaI sites flanking the $\mu$ and the Sy1 regions (Fig. 2). Similarly, e recombination juxtaposes the XbaI 5' to $\mu$ with an XbaI site within the 3' portion of the Se region (26). Since this XbaI site occurs within the Se region, rearrangements downstream of this site are not detected with this technique. Products of $\mu$-Sy1 and $\mu$-Se recombination were detected by circularization of XbaI-digested genomic DNA, and subsequent PCR amplification across the religated XbaI sites flanking the respective rearrangements. Since this PCR strategy does not amplify across the site of recombination, the 835- and 661-bp products will be generated irrespective of the site of recombination within $\mu$ and the Sy1 or Se regions, but will not be generated in the absence of DNA recombination.

As an internal control for the efficiency of the XbaI digestion and ligation reactions, a segment of the $\beta$-globin gene
complex was amplified from Xbal-digested and -circularized DNA using β-globin–specific primers. All cells should provide equivalent target DNA for the β-globin primers, and a 589-bp product should be amplified independent of DNA rearrangements within the Ig heavy chain locus. In the experiment shown in Fig. 3, genomic DNA was isolated from B lymphoblasts after 3 or 4 d of culture, digested with Xbal, then circularized as described in Materials and Methods. Aliquots from each ligation were placed directly into PCR reactions containing β-globin primers. A semi-quantitative assessment of the amount of target DNA was achieved by removing 10 μl aliquots from PCR reactions in three-cycle intervals. Since PCR amplification is a geometric progression, the amount of PCR product should increase eightfold with each three-cycle interval. Specific PCR products were detected by Southern analysis using PCR generated probes (see Materials and Methods). Amplification of the globin gene segment was detected after 23 cycles in each DNA sample, and product synthesis was comparable in each reaction through 26 cycles indicating that each DNA sample contained a comparable amount of target DNA (Fig. 3).

The sensitivity of the inverse PCR assay for detecting Sμ-Sγ1 and Sμ-Se rearrangements was estimated by combining hybridoma cells with known rearrangements to either γ1 (MOPC 21, an IgGl-producing myeloma) or e (HIE cells, an IgE-producing hybridoma) with WEHI-231 cells (an IgM-expressing B lymphoma) at ratios ranging from 1:10 to 1:10,000. Genomic DNA was isolated from 10⁷ total cells as described in Materials and Methods. Amplification was performed with the appropriate primer pair over 23–28 cycles and rearrangements detected by Southern analysis of PCR products. Under these conditions rearrangements to either the γ1 or e loci could be detected at cell ratios of 1 in 10,000, and appeared linear at ratios of 1:10 to 1:1,000 (data not shown).

**IL-5 Induces Switch Recombination to the γ1 Heavy Chain Locus.** The ability of LPS or IL-5 to promote IL-4-directed recombination to γ1 was evaluated by culturing B lymphoblasts with IL-4. Genomic DNA was isolated from B lymphoblasts on day 0 or after an additional 3 or 4 d of culture with the indicated combinations of IL-4 (200 U/ml), IL-5 (50 U/ml), and LPS (20 μg/ml). The DNA was digested with Xbal, and circularized by ligation. 5-μl aliquots were removed from ligation reactions and subjected to PCR amplification using either the β-globin primers or the Sμ(γ1) and Sγ1 primers. 10-μl aliquots were removed from PCR reaction after the indicated number of cycles. PCR products were detected by Southern blot analysis.
blasts with IL-4 alone or in combination with LPS or IL-5 for 3 and 4 d. The appearance of Sμ-Sγ1 rearrangements in the DNA samples was examined by PCR amplification using the Sμ(y1) and Sγ1 primers. Although IL-4 is not by itself mitogenic for B lymphoblasts (27) it maintained the viability of blasts in culture for several days in the absence of other added cytokines. This observation allowed analysis of the effects of IL-4 alone on DNA recombination. In three independent experiments, no Sμ-Sγ1 rearrangements were detected in B lymphoblasts cultured for up to 4 d with IL-4 alone, suggesting that IL-4 cannot drive recombination to the γ1 heavy chain locus. In contrast, γ1 recombination was detected in DNA isolated from B lymphoblasts cultured with IL-5 and IL-4 for 3 and 4 d. Thus, IL-5 can promote switch recombination that is directed to the γ1 heavy chain locus by IL-4. Similar findings were recently reported by Mandler et al. (28).

Consistent with previous reports (13, 14, 25, 29), LPS also induced marked accumulation of Sμ-Sγ1 rearrangements. Switch recombination to γ1 was readily detectable in DNA isolated from cells stimulated with LPS and IL-4 (200 U/ml) after 3 d of culture, and the apparent number of Sμ-Sγ1 rearrangements increased after 4 d. In fact, Sμ-Sγ1 rearrangements were observed as early as 2 d after addition of LPS and IL-4 (Fig. 4).

**LPS Is More Effective than IL-5 as a Stimulus for Switch Recombination to γ1.** The results presented in Fig. 3 suggest that B lymphoblasts cultured with LPS contained more Sμ-Sγ1 rearrangements than blasts cultured with IL-5. This observation contrasts with data presented in Fig. 1 demonstrating that IL-5 is more effective than LPS as a stimulus for IgG1 secretion. To further explore this observation, we focused on examination of Sμ-Sγ1 rearrangements at earlier time points to reduce the possibility of selective outgrowth of switched cells by LPS. B lymphoblasts were cultured with IL-4 and optimal concentrations of either LPS or IL-5, genomic DNA was isolated after 2 and 3 d of culture, and Sμ-Sγ1 rearrangements were examined by inverse PCR. As shown in Fig. 4, Sμ-Sγ1 rearrangements were detected after only 20 cycles of PCR in DNA isolated from blasts treated with LPS and IL-4. In contrast, detection of γ1 recombination in DNA from B lymphoblasts cultured with IL-5 required an additional six cycles. Under ideal conditions, six cycles of PCR amplifies the amount of PCR product 64-fold. Therefore, blasts cultured with LPS and IL-4 appear to contain at least 40-50-fold more γ1 rearrangements than blasts cultured with IL-5. These results suggest that B lymphoblasts treated with LPS undergo switch recombination to γ1 at a higher frequency than cells treated with IL-5.

**LPS but Not IL-5 Induces Accumulation of Sμ-ε Rearrangements.** In a previous study we were unable to detect germ-line ε transcripts or secretion of IgE in B lymphoblasts stimulated with IL-4 and IL-5 (15, Fig. 1). Accumulation of germ-line ε transcripts and secretion of IgE could be induced by inclusion of LPS in cultures with IL-4 (15). These results suggested that in the absence of LPS, IL-4 did not target the ε heavy chain locus for recombination. We examined this further by analyzing the ability of IL-5 or LPS to induce Sμ-ε rearrangements in B lymphoblasts cultured with IL-4. In three independent experiments, IL-5 and high concentrations of IL-4 (2,000 U/ml) did not promote accumulation of Sμ-ε rearrangements (Fig. 5). In contrast, LPS (plus IL-4) stimulated marked accumulation of Sμ-ε rearrangements that were detectable after 3 d of culture, and the apparent number of recombination events increased after 4 d. Induction of ε recombination was dependent on IL-4 as ε alone had no effect. These results demonstrate that LPS facilitates isotype switching to IgE by promoting ε recombination.

As shown in Fig. 1, exceptionally high concentrations of IL-4 are required to elicit IgG1 and IgE secretion from B lymphoblasts cultured with LPS. Therefore, it was intriguing to find that moderate concentrations of IL-4 (200 U/ml), which did not promote IgG1 and IgE secretion, were sufficient to mediate γ1 and ε recombination with LPS. Increasing the IL-4 concentration 10-fold consistently increased the apparent

**Figure 4.** LPS is more effective than IL-5 as a stimulus for γ1 recombination. Genomic DNA was isolated from B lymphoblasts cultured as in Fig. 3 for 2 and 3 d. PCR amplification of XbaI cut-circularized DNA and detection of PCR product was carried out as in Fig. 3 except that aliquots were removed from reactions after 20, 23, and 26 cycles. Occasionally, amplification using the Sμ(y1)-Sγ1 primers yielded the anticipated 835-bp product as well as a second smaller band. The second band may result from annealing of the Sγ1 primer at a site 3' to the intended target sequence. Amplification of the β-globin sequences was equivalent for each DNA sample.
number of \(S\mu-Se\) rearrangements, but it seems unlikely that this increase in \(\epsilon\) recombination explains the marked stimulation of IgE secretion observed with high concentrations of IL-4. These results clearly demonstrate that induction of switch recombination to the \(\gamma_1\) and \(\epsilon\) loci can be dissociated from secretion of the respective isotype.

Discussion

The current model for IL-4-directed switching to IgG1 and IgE suggests that IL-4 targets the \(\gamma_1\) and \(\epsilon\) heavy chain loci for recombination by regulating accessibility to a switch recombinase. Accessibility is manifested by transcription of the unrearranged loci and by accumulation of germline transcripts (1). However, IL-4 alone is not sufficient to promote expression of IgG1 or IgE (1), or DNA rearrangement to either loci (Figs. 3 and 5). Results presented here (Figs. 3 and 4) and elsewhere (28) indicate that IL-5 induces the accumulation of \(S\mu-S\gamma_1\) rearrangements in B cells treated with anti-Ig and IL-4. In addition to inducing \(\gamma_1\) recombination, IL-5 promotes the accumulation of \(\gamma_1\) heavy chain mRNA and secretion of IgG1 (15-19). These results indicate that IL-5 provides signals that are required for completion of isotype switching to IgG1, but the mechanism by which IL-5 promotes accumulation of \(S\mu-S\gamma_1\) rearrangements has not been established. It appears unlikely that IL-5 promotes selective outgrowth of cells that have already switched to IgG. Therefore, we favor a model in which IL-5 induces the expression or activity of components of the putative switch recombinase. Direct examination of this hypothesis awaits identification and characterization of molecular components that catalyze rearrangements within the Ig heavy chain locus.

Optimal stimulation of switch recombination requires signals in addition to those provided by IL-5. Thus, IL-5 failed to induce \(\epsilon\) recombination and LPS was more effective than IL-5 as a stimulus for \(S\mu-S\gamma_1\) rearrangement. Although LPS was also a more effective stimulus for B lymphoblast proliferation than IL-5, the greater number of \(S\mu-S\gamma_1\) rearrangements in cultures with LPS is probably not due to selective outgrowth of switched cells. Recombination to \(\gamma_1\) could be detected after only 2 d of culture with LPS and IL-4 (Fig. 4), an insufficient time for substantial expansion of a small subset of cells. Instead, LPS may be a superior stimulus for recombination because of its ability to induce expression of germline transcripts from unrearranged \(\gamma_1\) and \(\epsilon\) heavy chain loci.

It is noteworthy that the appearance of \(S\mu-S\gamma_1\) or \(S\mu-Se\) rearrangements is not always accompanied by secretion of IgG1 and IgE. For example, concentrations of IL-4 that caused readily detectable \(\gamma_1\) and \(\epsilon\) recombination (Figs. 3-5), did not elicit secretion of either isotype (Fig. 1) or substantial accumulation of \(V\delta J\gamma_1\) and \(V\epsilon\) alleles or mRNA (15). This suggests that additional factors are necessary for transcription of rearranged heavy chain alleles or mRNA stabilization. It is clear that IL-5 can stimulate the production or activity of such factors since it markedly enhances the appearance of \(\gamma_1\) and \(\epsilon\) mRNA in cells that have undergone rearrangements initiated by IL-4 and LPS (15, and Figs. 3 and 4). Although IL-5 can signal recombination to \(\gamma_1\) (28, and Fig. 3), our data suggest that the primary role of IL-5 in isotype switching may be to promote expression of the \(\gamma_1\) and \(\epsilon\) heavy chain loci subsequent to recombination.

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