Presence of Germline and Full-Length IgA RNA Transcripts Among Peritoneal B-1 Cells

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Next to conventional B cells (or B-2 cells), peritoneal B-1 cells have been shown to contribute significantly to the production of IgA-secreting plasma cells in the gut. Evidence for this was mainly based on studies comprising manipulated animals, including lethally X-irradiated and transgenic mice. To examine the ability of peritoneal B-1 cells from untreated mice to switch actively to IgA in vivo, we performed RT-PCR analysis on FACS-sorted peritoneal B-cell subsets from untreated BALB/c mice in order to examine the presence of germline Ca mRNA and mature Ca mRNA transcripts. Germline Ca and mature Ca transcripts were readily detectable in peritoneal B-1 cells (defined as IgM^bright/IgD^bright), but not, or very little, in peritoneal B-2 cells (defined as IgM^null/IgD^null). Moreover, by subdividing the B-1-cell population in CD5+ B-1a cells and CD5- B-1b cells, it was shown that in vivo expression of germline Ca and mature Ca transcripts was largely restricted to the B-1b-cell lineage. These results indicate that peritoneal B-1 cells indeed are capable to switch to IgA under normal physiological conditions and hereby further support the view that B-1 cells contribute significantly to the mucosal IgA response, albeit this function appears to be restricted to the B-1b-cell subset.

Keywords: B-1 cells, germline transcripts, IgA

INTRODUCTION

The intestinal lamina propria is characterized by a mucosal preponderance of IgA-secreting plasma cells (Van der Heijden et al., 1987). In the mouse, these cells have been shown to originate from two different lineages of B cells. An important source for IgA-secreting plasma cells is formed by Peyer’s patch B cells (Craig and Cebra, 1971) of which nearly all are bone-marrow-derived conventional type B cells or B-2 cells (IgM^null, IgD^bright). After antigen stimulation, committed B cells leave the Peyer’s patches, migrate via mesenteric lymph nodes and thoracic duct into the circulation to the spleen, and finally home back to the
gut lamina propria. During the migration, the cells expand, differentiate, and mature into IgA-secreting plasma cells (Tseng, 1984).

Another source of progenitor cells that contribute to the pool of lamina propria IgA plasma cells is formed by B-1 cells (Kroese et al., 1989, 1995), which reside abundantly in the murine peritoneal cavity (Herzenberg et al., 1986). The B-1 cell population (defined as IgM<sub>bright</sub>, IgD<sub> dull</sub>) can be divided, based on surface CD5 expression, into a CD5<sup>+</sup> B-1a cell and a CD5<sup>-</sup> B-1b “sister” cell population (Kantor and Herzenberg, 1993). The B-1 cell lineages differ from the conventional B-cell populations with respect to cell-surface-marker expression, localization, development, and antibody repertoire (Stall et al., 1996). Conventional B cells may undergo somatic hypermutation in the germinal centers and are responsible for high-affinity antibody responses to various antigens, whereas B-1 cells, which show a very low frequency of hypermutation of their V<sub>H</sub> genes (Kantor, 1996), primarily produce low-affinity IgM immunoglobulins, most of which are cross-reactive with bacteria-related and self-antigens (reviewed in Kroese et al., 1996).

Evidence that B-1 cells are involved in IgA production was demonstrated in different chimeric mouse models and B6-Sp6 μ,κ transgenic mice (reviewed in Kroese et al., 1995). Additionally, it was shown that several B-1-cell lines showed a high frequency of switching to IgA (Whitmore et al., 1991). However, all these studies comprised manipulated in vivo mouse models or in vitro experiments that left many questions regarding the B-1-cell lineage in normal untreated mice unanswered.

IgA class switching of IgM<sup>+</sup> B lymphocytes is preceded by the synthesis of germline C<sub>α</sub> mRNA transcripts (Lebman et al., 1990). Germline IgA transcription starts 5′ of an Iα exon, proceeds through the switch region, and terminates downstream of the C<sub>α</sub> exons. This leads to splicing of the Iα exon to the C<sub>α</sub> exon, which forms the germline IgA transcript (Lebman et al., 1990). Downregulation of germline IgA transcripts has been shown to inhibit IgA isotype switching that implies a role for IgA germline transcripts prior to the expression of full-length IgA transcripts (Lin and Stavnezer, 1992).

To study the ability of peritoneal B-1 cells (both B-1a and B-1b) to switch to IgA under normal physiological conditions, we sorted peritoneal B-cell subpopulations from untreated BALB/c mice and examined those populations for the presence of germline IgA transcripts and full-length IgA mRNA transcripts.

**RESULTS AND DISCUSSION**

**Peritoneal Washout Cells Express Germline Cα Transcripts**

The presence of germline Cα mRNA transcripts and mature IgA heavy-chain mRNA transcripts was determined by RT-PCR in unsorted peritoneal cells from 3-month-old untreated BALB/c mice. cDNA was synthesized from mRNA extractions of peritoneal washout cells as well as from sorted Peyer’s patch cells, which already have been shown to express Cα and mature IgA transcripts (Weinstein et al., 1991). Sorted splenic T cells served as negative controls. Additionally, all cell suspensions were tested for mature C<sub>μ</sub> transcripts. The C<sub>α</sub> germline transcripts were revealed by the use of the Iα-leader and Cα1-Cα2 primers, whereas full-length IgA and IgM transcripts were identified by usage of an universal V<sub>H</sub> primer in combination with C<sub>α</sub> or C<sub>μ</sub> primers, respectively. In each set of experiments, cDNAs of various different cell suspensions were normalized by means of serial dilutions with β-actin mRNA (650 bp). The specificity of the primers was tested on RNA derived from mouse IgA and IgM secreting hybridoma cell lines 2F7 (Bos et al., 1996) and NEO4211 (Bos and Meeuwsen, 1989), respectively. The germline primer did not result in a PCR product in either cell lines, whereas the universal V<sub>H</sub> primer in combination with the Cα primer and C<sub>μ</sub> primer did result in the correct PCR product for the corresponding hybridoma (Table I). Both unsorted peritoneal cells and Peyer’s patches cells express
Germline IgA Transcript Expression Is Restricted To Peritoneal B-1 Cells

To specify the phenotype of peritoneal B cells that switch to IgA, peritoneal B-cell subpopulations were sorted based on differences in IgM and IgD expression. The IgM^bright and IgD^null populations are formed by the B-1 cells, whereas the B-2 cells are found within the IgM^null and IgD^bright populations. Figure 1 shows a typical example of the staining used and the sorting gates set. The purity of the two sorted peritoneal B-cell subsets was approximately 95%. RNA was extracted and cDNA was synthesized from the sorted B-cell fractions and analyzed for mature IgM, mature IgA, and germline Cα transcripts.

Cα germline transcript and mature IgA expression occurred mainly in the peritoneal B-1-cell subpopulation (Figure 2 and Table I). In some B-2-cell sorts, a 20-50-fold lower expression of Cα germline transcript and/or mature IgA transcripts compared to the B-1-cell sorts could be detected. This could be due to contaminating B-1 cells, since in these sorts higher percentages of B-1-cell contamination was observed.

Our parameter for determining switching to IgA is the expression of germline Cα and full-length IgA mRNA transcripts. Germline transcripts are translated preceding the actual isotype switching. A strong correlation between germline transcript expression and isotype switching has been shown (Lebman et al., 1990). However, the function of germline transcripts is less clear, since mice lacking the Iα exon are still able to switch to IgA (Harriman et al., 1996).
they differentiate and mature to IgA plasma cells after migration out of the peritoneal cavity. For Peyer’s patch B cells, it is known that they leave those sites after initiation of isotype switching and migrate through the circulation toward the lamina propria while they further differentiate to IgA plasma cells (Tseng, 1984). Where and how B-1 cells migrate from the peritoneal cavity and differentiate to IgA plasma cells in the lamina propria remain to be established.

Also the factors that are involved in the initiation of isotype switching of B-1 cells in the peritoneal cavity are still unknown. Different interleukins, such as IL-4, IL-5, and TGF-β, already have been shown to play a role in the regulation of IgA switching and secretion (Harriman et al., 1988; Lebman and Coffman, 1988). Additionally, CD40 ligation by direct B-cell-T-cell interaction seems to be important for isotype switching (Jumper et al., 1994). Both peritoneal T cells as well as other cell types, such as mesothelial cells, might produce the correct cytokines and provide the stimuli for B-1 cells to start isotype switching to IgA. Whether peritoneal B-1 cells also can switch to other isotypes such as IgG and IgE remains also to be established.

IgA Expression Within the Peritoneal Cavity Is Confined to the B-1b-Cell Population

The B-1-cell population consists both of CD5+ B-1a cells and CD5− B-1b cells. To examine the ability of both peritoneal B-1a- and B-1b-cell subpopulations to switch to IgA, the subsets were sorted (Figure 1), RNA was isolated and cDNA was synthesized. Both B-1a and B-1b cells express mature IgM transcripts, however, the expression of germline Cα and mature IgA transcripts appeared to be largely confined to the peritoneal B-1b cells (Table 1 and Figure 2). The PCR products of the B-1b-cell population has also been cloned and sequenced, which confirmed the identity of the product as germline Cα transcripts compared to the EMBL sequence databank (data not shown). In one out of four experiments also, a very low expression of germline Cα transcripts was found in the B-1a subset, which was most probably due to contaminating B-1b cells in the B-1a-cell fraction.

Two alternative explanations can explain this preferential expression of IgA transcripts among B-1b cells. First, it might be argued that B-1a cells downregulate CD5 expression after switching to IgA. This might explain reconstitution experiments with
purified B-1a cells, which resulted in IgA plasma cells in the lamina propria (Beagley et al., 1995) and our own preliminary experiments with sorted peritoneal B1 cells showing that both B-1a and B-1b cells may contribute to the pool of intestinal IgA plasma cells (Kroese et al., unpublished observations). Alternatively, B-1a and B-1b cells might belong to closely related, but separate B cell lineages, whereby the switching to IgA is mainly restricted to the B-1b-cell lineage.

In conclusion, our experiments show that peritoneal B1 cells actively switch toward IgA in vivo, thereby confirming our previous data in manipulated animals that B-1 cells contribute to the IgA production in the mouse. The function of B-1-cell-derived IgA might be important in the establishment and maintenance of the normal gut flora, since we have shown that B-1-cell-derived monoclonal IgA antibodies primarily react with normal gut bacteria (Bos et al., 1996).

MATERIAL AND METHODS

Animals

Both male and female BALB/c mice at 3 to 4 months were studied. Mice were bred in the animal facility at the Stanford Department of Genetics.

Antibodies

Rat and mouse monoclonal antibodies used in this study were as follows: Rat anti-mouse IgD (11-26), rat anti-mouse IgM (331), rat anti-mouse Ly-1 (CD5; 53-78), mouse anti-mouse Igh-6a (IgM of “a” allotype, DS-1), and mouse anti-mouse Igh-5a (IgD of “a” allotype, AMS 9.1). Purification and conjugation of antibodies to biotin, fluorescein, phycoerythrin, and allophycocyanin (APC) are described elsewhere (Hardy et al., 1984).

Cell Preparation

Peritoneal washout cells were obtained from the mouse peritoneal cavity by injection of chilled deficient RPMI 1640 medium (Irvine Scientific, Santa Ana, CA) supplemented with 10 mM Hepes, 3% newborn calf serum and 0.1% NaN3. Single-cell suspensions from spleen and Peyer’s patches were prepared by mincing tissue fragments in the same medium between the frosted ends of microscope slides. All cell suspensions were treated with red-blood-cell lysis buffer to eliminate erythrocytes.

Staining and Cell Sorting

The B-cell subpopulations were sorted on an extensively modified fluorescence-activated cell sorter (FACS II; Becton-Dickinson, Mountain View, CA), as described (Hardy et al., 1984). For sorting subpopulations, single-cell suspensions were stained in tubes on ice with optimal concentrations of conjugated antibodies. Biotinylated antibodies were detected with avidin-Texas Red. Dead cells were stained with propidium iodide and were excluded from sorting. After sorting, 30,000 viable sorted cells were reanalyzed to test the purity of the sorted cells.

RT-PCR

Total RNA, isolated from different cell suspensions by use of TRIzol (Life Technologies) according to the manufacturer’s instructions, was used as a template for cDNA synthesis in a 30-μl reaction mix containing 1.6 μg oligo-dT 12-18 (Pharmacia), 10 mM dNTP mix (Pharmacia), MilliQ-DEPC, First Strand Buffer (Life Technologies), 0.1 mM DTT (Life Technologies), 30 U RNA guard (Pharmacia), and 200 U Superscript™ II (Life Technologies). The reaction proceeded at 45°C for 30 min, followed by heating to 94°C for 5 min. For the RT-PCR, an aliquot of cDNA
was added to a cocktail composed of 5 μl (10×) RT-PCR buffer (Life Technologies), 0.5 μl 20 mM dNTP mixture, 1.5 μl MgCl₂ (Life Technologies), 2.5 μl 1% W-1 (Life Technologies), 0.6 μl 5’ primer (25-35 pmol/μl), 0.6 μl 3’ primer (25-35 pmol/μl), and 0.5 μl Taq DNA polymerase (5 U/ml) (Life Technologies). Primers sequences, 5’ to 3’, used in this study were as follows:

3’ β-actin: TCTTCATGGT GCTAGGAGCCA
5’ β-actin: CCTAAGGCCAACCGTGAAAAG
Iα-leader: CCAGTCCTAAGCTTFCTACCATAG
IgL Co2: GAGGAGTAGGACCAGAGCAATTC
(Cebra et al., 1996)
IgL BamHI: CTCGGATCCTCACATT CATCGTGCC (Bos et al., 1996)
Universal Vh: ACGAATTCAAGTSMARCTGCAGSAGTCWGG (M A or C; R A or G; S C or G; W A or T) (Orlandi et al., 1989)
Cμ 1.3: CCCCCTGGATGACTTCAGTGTTG

The RT-PCR was performed on a thermal cycler (Pharmacia LKB—Gene ATAG Controller) with an initial denaturation cycle at 94°C (2 min), an annealing step at 60°C (1 min), and an extension step at 72°C (1 min), respectively. Then 35 steps of amplification were performed (each step of the cycle: 1 min) with a denaturing temperature of 94°C, an annealing temperature of 60°C, and an extension temperature of 72°C. The PCR products (10 μl per reaction) were analyzed on an 1.2% ethidiumbromide-agarose gel and visualized in UV light.

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