Alternative use of 5' exons in the specification of Ly-5 isoforms distinguishing hematopoietic cell lineages

(hematopoietic differentiation/Ly-5 molecular isoforms/alternative splicing/S1 nuclease mapping)

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Contributed by Edward A. Boyse, April 20, 1987

ABSTRACT Previous inferences that Ly-5 glycoprotein isoforms of murine hematopoietic cells are generated by alternative splicing of primary transcripts of a single Ly-5 gene are supported by the present study. A cDNA library was prepared from B-cells by extension from primer representing a known T-cell cDNA sequence. Three different Ly-5 clones from this library included sequences missing in T-cell cDNA clones. From the constitution of cDNA clones and of the Ly-5 gene, and from S1 nuclease mapping, it is concluded that at least two exons, provisionally numbered Ex-6(B) and Ex-7(B), in the 5'-proximal region are mainly represented in mRNA of the B-cell lines examined but not of the T-cell lines examined. Also, exons 1 and 2 appear to be used alternatively in different species of B-cell mRNA and probably also in different species of T-cell mRNA.

Interest in the Ly-5 system of the mouse (1), independently described as the T200 system (2) before it was known that Ly-5 and T200 refer to the same system, stems largely from representation of the products as multiple glycoprotein isoforms of the cell-surface defining stages and lineages of hematopoietic development (3-5). Two of these isoforms, a 200-kDa isoform from T cells and a 220-kDa isoform from B cells, are conveniently named T200 and B220 (6). Evidence from protein composition, the constitution of cDNA clones and details of mRNA (7, 8) have suggested that the larger B220 isoform is encoded by a longer mRNA containing extra sequences shortly following a leader peptide sequence and that T200 and B220 RNAs are generated by alternative splicing of the primary transcript of a single Ly-5/T200 gene. That conclusion is supported by evidence presented here that B220 cDNA clones include extra sequences at the predicted position, representing at least two exons absent from T200 mRNA in consequence of alternative splicing.

MATERIALS AND METHODS

Preparation of RNA. Total RNA was prepared from the cell lines named using guanidinium isothiocyanate followed by centrifugation through a cushion of CsCl (9). Poly(A)* RNA was then isolated by oligo(dT)-cellulose chromatography (10).

Construction of a Primer-Extension B-Cell cDNA Library. Antisense oligonucleotide primer comprising 18 nucleotides (nt) was synthesized according to a sequence 151-168 nt downstream from the putative interolation site of extra sequences used by B cells (8). Primer was hybridized with 10 μg of poly(A)* RNA prepared from the B-cell leukemia L.29. The first strand was synthesized by reverse transcriptase at 42°C (11) and second-strand synthesis was by RNase H and DNA polymerase I according to Gubler and Hoffman (12). Double-stranded DNA fragments, blunt-ended by RNase A and DNA polymerase I, were ligated into the bacterial alkaline phosphatase-treated Smal I site of pUC13 vector (Pharmacia).

S1 Nuclease Protection Mapping. Probes for S1 nuclease mapping were prepared from clones pLy-5-B15 and -B51 (see below). End-labeled probe DNA (2 × 104 cpn) was hybridized with total RNA (30 μg) at various temperatures and digested with S1 nuclease (37°C for 30 min) (13). Protected fragments (PFs) were analyzed on 6% polyacrylamide/7 M urea sequencing gel.

RNA Transfer Blotting. Electrophoresis and blotting of RNAs and hybridization with nick-translated 32P-labeled probes were done as described (13).

Construction and Screening of a λ Genomic Library. This library was made from cells of the T-cell leukemia variant EARAD1-β2M in EMBL4 vector as described (8).

Sequencing of DNA. DNA was sequenced according to Sanger et al. (14) on fragments subcloned in M13mp18 and M13mp19.

RESULTS AND DISCUSSION

B-Cell Ly-5 cDNA Has Sequences Not Present in T-Cell Ly-5 cDNA. By screening the B-cell primer-extension library with a T-cell Ly-5 cDNA probe, three varieties of B-cell cDNA were obtained, represented by clones pLy-5-B11, -B12, and -B15. Fig. 1 illustrates that all three clones have sequences that are lacking in the T-cell cDNA clones and are located as inferred from S1 nuclease mapping (8). All three B-cell clones share a common B-cell distinctive sequence upstream of position 196 of the T-cell cDNA clone T4 (the present notation T4 (pLy-5-T4) replaces the former notation pLy-5-R4 (8)). Upstream of this common B-cell sequence (141 nt; Fig. 1, box E) the three B-cell clones differ (Fig. 1)—B11 is completed by a sequence (194 nt) identical to T4 (boxes C and B). B15 has a longer B-cell sequence (141 and 147 nt; boxes E and D), followed by a shorter sequence shared with T4 (135 nt; box C), and is completed by a 91-nt sequence (box A) not shared by any other T- or B-cell cDNA clone. B12 is completed by a sequence that contains stop codons in the same reading frame, suggesting an intron (confirmed by genomic sequencing; see below).

Reading frames for the B-cell distinctive sequences of B11 and longer B15 clones (equivalent to 47 and 96 amino acids, respectively) are in phase with T4. The predicted 96-amino acid region contains two potential N-glycosylation sites and is rich in serine and threonine residues (potential sites for O-linked glycosylation). This region contains no cysteines for disulfide linkage and is relatively hydrophilic, allowing that this region is exposed and may be an epitope for antigen known to be unique to the B200 Ly-5 isoform (6).

Abbreviations: nt, nucleotide(s); UT, untranslated; PF, protected fragment.
B-Cell-Distinctive Ly-5 mRNA Sequences Are Encoded by Exons Lacking in T-Cell mRNA. A genomic library in EMBL4, screened with a T-cell Ly-5 cDNA probe, yielded 13 overlapping clones spanning ≈120 kilobases (kb) and inferred to comprise a single Ly-5 gene from which diverse Ly-5 isoforms are generated (8). The B-cell-distinctive cDNA sequences noted above also imply an origin of Ly-5 isoforms by alternative splicing of primary Ly-5 transcripts.

Fig. 2 shows exon–intron organization of the 5′ end of the Ly-5 gene, including the first 8 exons that comprise the 5′ region of T4.

The first exon represented in T4 cDNA is Ex-2 (Fig. 1, box B) and belongs to the 5′ untranslated (UT) region. The next is Ex-3, composing the rest of the UT region and the leader sequence (15). Ex-4 contains the first seven N-terminal amino acids common to all Ly-5 isoforms (15). These exons form the region denoted by box C (Fig. 1).Fig. 2, 8ex, is approximately 10 kb downstream from Ex-4.

To verify the prediction from cDNA sequences that B-cell-distinctive sequences should lie between Ex-4 and Ex-8, a B-cell cDNA fragment containing these sequences was used to probe DNA from the λ genomic clones in Southern blotting. Two exons, Ex-6(B) and Ex-7(B) (Fig. 1, boxes D and E), were identified by sequencing genomic fragments thus selected and are seen to lie between Ex-4 and Ex-8. [A further upstream B-cell-distinctive, called Ex-5(B) in our provisional notation (129 nt), is predicted by the complete sequence of a cDNA clone isolated from the pre-B-cell line]
Inclusion versus exclusion of Ex-6(B), it appears that there is at least one additional site for alternative splicing in the 5' UT region. Thus, clone B15 begins with Ex-1, whereas clones B11 and also T4 begin with Ex-2. S1 nuclease mapping was used to find out whether some Ly-5 mRNA species of T cells also may contain Ex-1 in place of Ex-2. For this purpose, a probe with its 5' end in Ex-3 and incorporating all of Ex-1 (probe I, Fig. 3) was prepared from B15.

Fig. 3 shows that a protected fragment (PF1, 161 nt) was observed with mRNA of T cells (leukemias ISL-57 and ASL1) as well as B cells (leukemias I-29 and pre-B 70Z/3), suggesting that some T-cell mRNA may use Ex-1 in place of Ex-2. A second PF (PF2, 70 nt) is probably protected by T4 mRNA and B11-like mRNA, which incorporate only the Ex-3 region of the probe.

Thus, tentatively, T cells or sublineages of T cells, like B cells, may use either Ex-1 or Ex-2, and in both cases transcription may begin at alternative sites. Preliminary data from primer extension suggest alternative initiation sites for both T- and B-cell transcripts and that the presence of Ex-1 and of Ex-2 are mutually exclusive. Alternative use of Ex-1 and Ex-2 would seem to generate diversity of sequence in the 5' UT region without altering the reading frame.

In Both T- and B-Cell Lineages, Ly-5 mRNAs Are Diverse, Using Alternative 5' Exons. The diversity of B-cell transcripts inferred from the composition of clones B11 and B15 was studied further by S1 nuclease mapping. Probe II (767 nt, from the 5' end to the EcoRV site in Ex-7(B)) was prepared from clone B51 (see legend of Fig. 4), end-labeled, hybridized with total B-cell RNA (I29, 70Z/3), and digested with S1 nuclease.

Fig. 4 shows four PFs. The longest (PF1, 531 nt) is evidently protected by mRNA of B15 type [Ex-1, -3, -4, -6(B), and -7(B)]. PF2 (414 nt) is probably protected by mRNA containing Ex-3, -4, -6(B), and -7(B), without Ex-1; because Ex-1 is missing in these RNAs (RNA type 1 predicted), we surmise that they use Ex-2 instead of Ex-1, which T4 and B11 use. PF3 (278 nt) appears to be protected by RNAs (RNA type 2 predicted) that contain Ex-6(B) and -7(B) and presumably relates to a third B-cell exon, provisionally called Ex-5(B), noted by Thomas et al. (16) (see above), which is not represented in our cDNA clones. PF4 (131 nt) is probably protected by mRNA representing B11, containing only Ex-7(B) among B-cell specific sequences, and/or B12, which contains the upstream intron sequence.
Since the 5' end of probe II (Fig. 4) is located within Ex-7(B), T-cell mRNA should not protect this probe and should yield no PFs. Accordingly, total RNA of ISL-57 T cells yielded neither PF1 nor PF2. However, small amounts of PF3 and PF4 were observed (Fig. 4). This finding, that a B-cell-specific probe can be marginally protected by T-cell mRNA, called for further research of T-cell transcripts. Therefore, we prepared two genomic probes—probe 6(B) containing Ex-6(B) and probe 7(B) containing Ex-7(B)—for RNA transfer blotting (Fig. 5). Both probes hybridized strongly with B-cell RNA as expected. However, probe 6(B), and to a lesser extent probe 7(B), hybridized with RNA of T cells (ISL-57 and ASL1), suggesting that some species of T-cell RNA may include Ex-6(B) and/or Ex-7(B), although most T-cell Ly-5 RNA does not, as concluded previously (8). After complete removal of probes 6(B) and 7(B), the same bands were hybridized with probe A, a 2.3-kb BamHI fragment of cDNA clone pLy-5-T4, which is common to T and B cells. Comparing the hybridization patterns of probes 6(B) and A in Fig. 5, it is clear that probe A gives a broader band, probably signifying more than one RNA species, in contrast to probe 6(B), which gives only the upper part of that band. The same may apply to Ex-7(B) in lesser degree according to the lesser intensity of the hybridization bands evident in Fig. 5. Thus, on present evidence PF3 and PF4 generated from T cells (as shown in Fig. 4) may be construed to represent minor species of T-cell RNA that include Ex-6(B) and/or Ex-7(B).

Possible Relationship of Genomic Sequence to Alternative Modes of Processing of Primary Transcripts. In this context, we searched regions flanking intron–exon boundaries of the 33 Ly-5 exons identified, which include the sequences of all present cDNA clones, to find sequences that might distinguish constitutive exons (used for all Ly-5 isoforms) from conditional exons (used for some Ly-5 isoforms). The first eight exons shown in Fig. 6 are flanked by consensus splicing junction sequences and follow the AG/GT rule (18). All six exons of the coding region terminate within a codon, splitting the first and second nucleotides. We note that a stretch of similar sequences [GG(T/A)GTGAT] (doubly underlined) occurs in the upstream intron of Ex-6(B) (−31 to −38) and Ex-7(B) (−27 to −34) but not of the other 31 exons. The last nucleotide of the upstream intron and first six 5' nucleotides

Thus B-cell mRNAs are diverse in their 5' coding regions. Whether each transcript is functional remains to be seen; there appears to be a single Ly-5 isoform (220 kDa) in 70Z/3 pre-B cells but I.29 B leukemia cells appear to yield more than one Ly-5 isoform, all of somewhat smaller molecular mass (according to Laemmli-type concentration-gradient NaDodSO4/PAGE; unpublished observation), which might imply more than one functional transcript in the I.29 cell population.

Fig. 5. RNA transfer blotting with probes 6(B) and 7(B) (Left) prepared from genomic clones and containing Ex-6(B) and Ex-7(B), respectively, followed by probe A (Right) prepared from cDNA clone pLy-5-T4 as a 2.3-kb BamHI fragment and representing a sequence common to both T- and B-cell cDNA. Ly-5 RNA of I.29 cells appears slightly smaller than RNA of 70Z/3 cells, in keeping with the observation (unpublished) that I.29 cells express slightly smaller and diverse Ly-5 isoforms.

Fig. 6. Exon–intron boundary sequences of the first eight Ly-5 exons. Aligned boundary sequences show close similarities to consensus sequences for both donor and acceptor splicing sites (18). Doubly underlined and heavily underlined sequences are unique to Ex-6(B) and Ex-7(B).
(heavily underlined) of the splicing acceptor sites of Ex-6(B) (G/GTGTGT) and Ex-7(B) (G/GTGTGC), but not of the acceptor sites of the 31 other exons, resemble the consensus sequence of the splicing donor site, G/GT(G/A)AGT (18). Possibly these junction sequences peculiar to Ex-6(B) and Ex-7(B) may be recognized as splicing donor sites in the T-cell T200 program but not in the B-cell B220 programs, the succeeding exon being spliced out only in T cells.

We thank Mr. Thomson C. Pancoast for excellent technical assistance. E.A.B. is American Cancer Society Research Professor of Cell Surface Immunogenetics. Y.S. receives a fellowship from the Cancer Research Institute. This work was supported in part by Grants CA-39827 and AI-21840 from the National Institutes of Health.
