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Differential Activation of Dual Promoters Alters D β 2 Germline Transcription during Thymocyte Development¹

Ruth E. McMillan and Michael L. Sikes²

Ag receptor genes are assembled through somatic rearrangements of V, D, and J gene segments. This process is directed in part by transcriptional enhancers and promoters positioned within each gene locus. Whereas enhancers coordinate reorganization of large chromatin stretches, promoters are predicted to facilitate the accessibility of proximal downstream gene segments. In TCR β locus, rearrangement initiates at two D-J cassettes, each of which exhibits transcriptional activity coincident with DJ rearrangement in CD4/CD8 double-negative pro-T cells. Consistent with a model of promoter-facilitated recombination, assembly of the DJ β 1 cassette is dependent on a D β 1 promoter (PD β 1) positioned immediately 5' of the D. Assembly of DJ β 2 proceeds independent from that of DJ β 1, albeit with less efficiency. To gain insight into the mechanisms that selectively alter D usage, we have defined transcriptional regulation at D β 2. We find that both DJ β cassettes generate germline messages in murine CD44⁺CD25⁻ double-negative 1 cells. However, transcription of unrearranged DJ β 2 initiates at multiple sites 400–550 bp downstream of the D β 2. Unexpectedly, loci from which germline promoter activity has been deleted by DJ rearrangement redirect transcription to sites immediately 5' of the new DJ β 2 joint. Our analyses suggest that 3'-PD β 2 activity is largely controlled by NF- κ B RelA, whereas 5'-PD β 2 activity directs germline transcription of DJ β 2 joints from initiator elements 76 bp upstream of the D β 2 5' recombination signal sequence. The unique organization and timing of D β 2 promoter activity are consistent with a model in which promoter placement selectively regulates the rearrangement potential of D β 2 during TCR β locus assembly. *The Journal of Immunology*, 2008, 180: 3218–3228.

The Ig and TCR genes are assembled through a series of highly regulated somatic rearrangements in developing B and T lymphocytes, respectively. Despite cell-specific segregation of Ig and TCR rearrangements, both types of Ag receptor genes are assembled via a conserved mechanism. For each locus, the Ag recognition domain is assembled from arrays of segments termed V, D, and J, by a single enzymatic complex, which targets recombination signal sequences (RSS)³ that flank individual V, D, and J segments. During rearrangement, select RSS are bound by the lymphocyte-specific components of recombinase, encoded by RAG-1 and RAG-2 (1, 2), which introduce dsDNA breaks precisely at the boundaries between the RSS and their coding sequences (3). Processed coding ends are then ligated by ubiquitous double-stranded break-repair machinery to form unique coding joints.

As they mature in the thymus, T lymphocytes transition through a series of developmental stages identified by differential expression of numerous cell surface proteins, including CD4, 8, 25, and 44. Rearrangement and expression of individual TCR genes serve

as obligate checkpoints during this maturation process. For example, before thymocytes express the CD4 and CD8 coreceptors, so-called double-negative (DN) cells must complete the assembly of a functional TCR β locus (*Tcrb*) gene (4). Cells that survive this period of β selection (5) progress to the CD4/CD8 double-positive (DP) stage of development, in which TCR α locus (*Tcra*) genes are assembled (6). Recombination of the *Tcrb* occurs in a stepwise fashion in DN cells, involving initial D-to-J joining at each of two cassettes that contain 1 D and 6 or 7 Js, followed by assembly of 1 DJ joint with 1 of ~20 distal V elements. Recombination of the upstream D β 1 to its associated J elements has been detected in the earliest stage of DN thymocyte maturation, termed DN1 (CD44⁺25⁻) (7, 8), and accumulates through DN development. In contrast, rearrangement of the second DJ cassette was not detected in granulocyte/myelocyte clones derived from the DN1 or DN2 (CD44⁺25⁺) cells of an IL-2R β transgenic mouse (7). The potentiality that D β 2 rearrangement initiates later in thymocyte development than does D β 1 assembly correlates with the long-standing observation that germline D β 2 sequences persist in the endogenous loci of TCR β transgenic mice (9) and fetal thymocytes (10–12). Whether the persistence of germline D β 2 sequences stems from a delay in the onset of D β 2 rearrangement or a relative inefficiency with which the DJ β 2 cassette gains recombinational accessibility, its consequence is to limit the availability of DJ β 2 substrates for initial V-to-DJ recombination in DN3 (CD44⁻low25⁺) cells.

The strict developmental programming of V(D)J recombination has been proposed to reflect programmed decondensation of the chromatin surrounding individual gene segments, thereby making them more accessible to recombinase. In support of this accessibility model, recombination has long been thought to require enhancer-driven chromatin decondensation and germline transcription (13, 14). Indeed, enhancer deletion impairs rearrangement to varying degrees in each of the Ag receptor loci (14). In *Tcrb*, recombination is strictly dependent upon the lone enhancer, *Tcrb*

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³ Abbreviations used in this paper: RSS, recombination signal sequence; Ct, cycle threshold; DN, double negative; DP, double positive; E β , *Tcrb* enhancer; *inr*, transcription initiator; IP, immunoprecipitation; PD β , D β promoter; SP, single positive; *Tera*, TCR α locus; *Tcrb*, TCR β locus; USF1, upstream stimulatory factor 1.

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enhancer ($E\beta$) (15, 16), which has been shown to modulate the chromatin organization of a 35-kb domain that spans both DJC cassettes (17). Recombination of the first DJ β cassette holds an additional requirement for the activation of an associated germline promoter, D β 1 promoter (PD β 1) (18, 19). Similar requirements for germline promoters in the *Tcra* (20, 21) and *IgH* loci (22) have suggested a general role for promoters in regulating the accessibility of neighboring gene segments. When PD β 1 was repositioned between D β 1 and J β 1.1, recombination of a chromatinized TCR β minilocus was severely attenuated (23). This sensitivity of DJ β 1 recombination to moving the promoter downstream of the D, together with the role of transcription from individual *Tcra* germline promoters in targeting rearrangement of downstream J α elements (24), implies that the placement and timing of germline promoter activity relative to individual Ag receptor gene segments may profoundly impact patterns of gene segment usage.

To better understand the role of germline promoters in coordinating the differential usage of Ag receptor gene segments, we sought to characterize the elements that regulate germline transcription within the DJC β 2 cassette. In this study, we describe two regions of promoter activity flanking the D β 2 gene segment. Before rearrangement, germline transcription in CD44⁺25⁻ DN1 thymocytes is initiated at a diffuse array of start sites 400–550 bp 3' of D β 2. The sites of transcription initiation overlap a poorly organized 3' promoter driven largely by p65 RelA binding at two NF- κ B sites immediately upstream of J β 2.1. The unexpected placement of PD β 2 between D β 2 and J β 2.1 precludes transcription through the D β 2 RSS, suggesting a potential model for the inefficiency of DJ β 2 assembly (9–12). We identified an additional promoter element 5' of the D β 2. This 5'-PD β 2 was only revealed when sequences between the two promoters were deleted, and mirrored the redirection of transcription to consensus initiator elements immediately upstream of D β 2 upon assembly of a DJ β 2 joint in developing thymocytes. In light of this unique functional architecture, we speculate that differential promoter usage within the DJC β 2 cassette may play a key role in coordinating the scope and timing of individual *Tcrb* rearrangements.

Materials and Methods

Cell sorting

FITC-conjugated CD8a (53-6.7) and CD25 (7D4) Abs, as well as PE-conjugated CD4 (RM4-5) and CD44 (IM7) Abs were purchased from BD Pharmingen. Thymii were isolated from 5-wk-old C57BL/6 mice, and RBC were removed by hypotonic lysis. Thymocytes were subsequently mixed with Abs to CD4 and CD8, and labeled cells were removed on magnetic beads coated with sheep Ab to rat IgG (Dyna). The remaining cells were labeled with FITC-conjugated anti-CD25 and PE-conjugated anti-CD44 Abs before sorting into CD44⁻/CD25⁻, CD44⁺/CD25⁻ (DN1), CD44⁺/CD25⁺ (DN2), and CD44⁻/CD25⁺ (DN3) cell populations on a 3 laser MoFlo cell sorter (DakoCytomation). Postsort evaluation of individual populations revealed purities of $\geq 93\%$ for each of three experiments. All mouse studies described in this work were reviewed and approved by the institutional animal care and use committee at North Carolina State University.

RT-PCR

Total RNA was isolated using TriReagent (Sigma-Aldrich), according to the manufacturer's recommendations. RNAs (0.5 μ g) were reverse transcribed using Moloney murine leukemia virus reverse transcriptase (100 U; New England Biolabs) and oligo d(T) primers. The resultant cDNAs were amplified with 1 μ mol each of primer pairs for D β 1 (forward, 5'-GGCT ACCTCACTTTGATG-3'; reverse, 5'-CCCAGGCCTCTGCATGTGATGTCTGTGTG-3'), D β 2 (forward, 5'-CAGTTCTGGAGGTAGATGGAGAATG-3'; reverse, 5'-CTGTGTGACAGGTTGGGTGAGCCCTCTG-3'), and β -actin (18) in either 10 mM Tris-Cl (pH 9), 50 mM KCl, 2 mM MgCl₂, 200 mM dNTPs, and 1 U of *Taq*, or 1 \times iQ SYBR Green Supermix (Bio-Rad). For semiquantitative PCR, reaction mixtures were amplified (94°C, 1 min; 57°C, 1 min; 72°C, 1.5 min) for 27 (actin) or 30 cycles

(*Tcrb*), and the products were assessed using gel electrophoresis and Southern blotting. For real-time PCR, reaction mixtures were cycled 50 times. Quantitation of D β cDNAs was achieved by comparison of sample cycle threshold (Ct) values with those of serially diluted C57BL/6 thymus cDNA. Variations in sample loading were controlled by normalizing D β values to values obtained for β actin.

Transcript analyses

For Northern blotting, 10 μ g of total RNA was electrophoresed on 1% agarose formaldehyde gels. Duplicate ζ -Probe (Bio-Rad) membranes were hybridized with probes 5' of D β 2 (1-kb *StuI/AluNI* fragment) or 3' of D β 2 (650-bp *AflIII/EcoNI* fragment). Relative amounts of RNA in each lane were estimated by hybridization with a probe to GAPDH.

For RNase protection, ssRNA probes were generated from a plasmid harboring a 1-kb PCR fragment spanning from 96 bp 5' of D β 2 to the J β 2.1 RNA splice donor site. The plasmid was linearized with *SmaI*, and radiolabeled antisense probes were generated using the T7 polymerase in the presence of [α -³²P]CTP. Full-length probes were purified on a 5% polyacrylamide gel, and hybridized with 50 μ g of total RNA using the RPA III kit (Ambion). Hybridization products were separated by electrophoresis on a 5% polyacrylamide gel, and visualized by autoradiography.

The 5'-RACE was performed using a BD Smart Race kit (BD Biosciences), according to the manufacturer's instructions. Touchdown PCR of individual cDNAs was performed, according to the manufacturer's instructions, using a J β 2.1-specific 3' primer (5'-TAGGACGGTGTGATCGTGTCCC-3'), and replacing polymerase activity with the Phusion polymerase blend (New England Biolabs). Amplification products were cloned into pBluescript (Stratagene), and individual clones were sequenced.

Plasmids

To construct luciferase plasmids, individual restriction fragments were isolated from p5'D2JJ-BS, which harbors a 2.5-kb PCR fragment spanning the D β 2-J β 2.1-J β 2.2 region of murine *Tcrb*. Fragments were purified by gel electrophoresis and Qiaquick gel extraction columns (Qiagen). Purified fragments were blunt ended and inserted into the *SmaI* site of the pGL3-E β or pGL2-enhancer vectors (R. McMillan and M. Sikes, unpublished data). The pGL3-E β vector carries a 570-bp *StuI/NcoI* fragment containing the mouse *Tcrb* enhancer, E β , inserted into a *BamHI* site downstream of the firefly luciferase gene. Site-specific mutations were introduced into individual reporter constructs using the Quickchange II site-directed mutagenesis kit (Stratagene), according to the manufacturer's recommendations. Sequences for the oligonucleotide primers used to introduce each mutation (shown in bold) are as follows: mG3-5, 5'-GGTCTTATAACATCCGAGCATCTT-3'; mG3-6, 5'-TTCAGCCCTTGGCATGTAA-3'; mG3-7, 5'-GAATAGATGGGCTTCCGTCC-3'; m κ B-1, 5'-AGAATGTGAGATGCCCCGGGTCT-3'; m κ B-2, 5'-AGGAAGCGCAGGAAAGAGG-3'.

EMSA

Potential transcription factor binding sites were predicted using Transcription Elements Search Software (University of Pennsylvania). Gelshift probes to each potential site were generated by annealing equimolar amounts of complementary single-stranded oligonucleotides in 10 mM Tris-Cl (pH 7.4), 1 mM EDTA, and 50 mM NaCl. Each annealed oligonucleotide (1 ng) was labeled by Klenow-mediated fill-in of 4- to 6-base 5' overhangs using [α -³²P]dCTP and [α -³²P]dATP radionucleotides. Nuclear protein extracts were prepared, as previously described (25), from untreated P5424 cells, or cells treated 4 h with either PMA (20 ng/ml) and ionomycin (1 μ M), or overnight with LPS (1 μ g/ml). Nuclear extracts were quantitated by Bradford protein assay (Bio-Rad) and stored at -80°C.

For binding reactions, each double-stranded oligonucleotide probe (1 ng) was incubated with 20 μ g of the indicated nuclear extract for 20–30 min on ice in a binding mixture (20 μ l) containing double-stranded poly(dI-dC) (2 mg), and BSA (10 mg) buffered in 20 mM HEPES (pH 7.9), 5% glycerol, 1 mM EDTA, 1% Nonidet P-40, and 5 mM DTT. For cold competition or supershift, reaction mixtures were pretreated with either unlabeled double-stranded competitor (100 ng) or Ab (1 μ g) for 30 min on ice before addition of the radiolabeled probe. Abs against Sp1 (sc-59), GATA-3 (sc-268), NF- κ B p65 (sc-372), and NF- κ B p50 (sc-8414) were purchased from Santa Cruz Biotechnology. Reactions were separated on 6% nondenaturing polyacrylamide gel, and visualized by autoradiography. Sense strand sequences for wild-type oligonucleotides are as follows: G3-5, 5'-GGTCTTATAACATCTATGCATC-3'; G3-6, 5'-3'; PD β 1 G3 (25); κ B-1, 5'-GAGAATGTGAGTAACC-3'; κ B-2, 5'-TTGAGGAAGGTGAGGAAAGAG-3'; Sp1, 5'-AACATGTGAGGAGGAGTCTAT-3'; NF-Y, 5'-AAGAGATTAACCAATCACGTA-3'. The IL-2R α κ B double-stranded oligonucleotide was purchased from Santa Cruz Biotechnology (sc-2511).

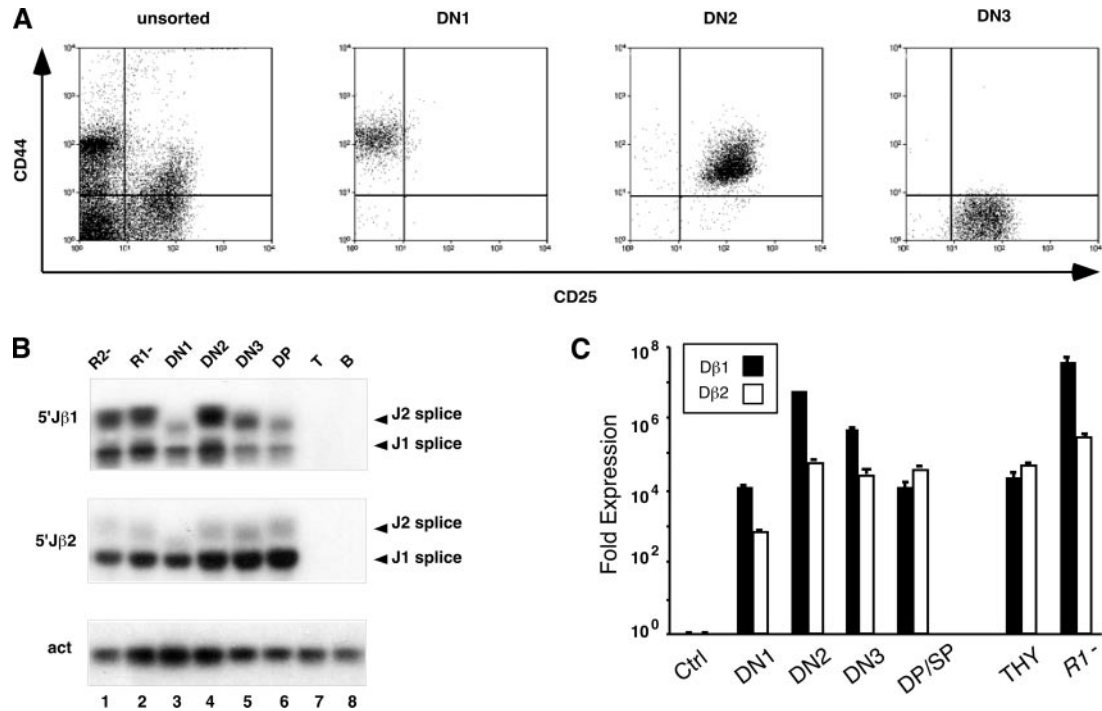


FIGURE 1. Developmental timing of DJ β germline transcription. *A*, Flow cytometry CD44 (y-axis) and CD25 (x-axis) expression in DN thymocytes following immunodepletion of CD4/CD8 DP and SP cells. Gates used to sort DN1 cells (CD44⁺CD25⁻), DN2 cells (CD44⁺CD25⁺), and DN3 cells (CD44⁻CD25⁺) are shown. Preparation of 30 thymii yielded $\sim 10^9$ total thymocytes, and sorted DN numbers of $0.5\text{--}2 \times 10^6$ DN1 (0.1% total), $0.3\text{--}1.1 \times 10^6$ DN2 (0.07% total), and $2\text{--}6 \times 10^6$ DN3 (0.4% total). Postsort evaluation revealed purities of 93–95% for DN2 and 96–98% for DN1 and DN3 for each of three independent sorts. *B*, RT-PCR analysis of germline transcripts that initiate 5' of J $\beta 1.1$ (upper panel) or 5' of J $\beta 2.1$ (middle panel), and splice to C $\beta 1$ or C $\beta 2$, respectively. R2⁻, total RAG-2^{-/-} thymocytes; R1⁻, total RAG-1^{-/-} thymocytes; T, BW5147; and B, M12 cell lines. PCR of the unrelated β -actin message (lower panel) served as a loading control. *C*, Quantitative real-time RT-PCR of D $\beta 1$ (■) and D $\beta 2$ germline transcription (□). Amplification signals were normalized for each sample ($n = 4$ replicates) to those obtained for β -actin, and expressed as fold induction above that obtained for the M12 B cell line (Ctrl). THY, unsorted C57BL/6 thymocytes.

Luciferase assays

The RAG-1^{-/-}, p53^{-/-} pro-T cell line, P5424 (26), the BW5147 mature T (27), and M12 mature B cell lines (28) have been previously described. All cell lines were cultured at 37°C/5% CO₂ in RPMI 1640 medium supplemented with 10% FCS, 2 mM L-glutamine, 0.01% penicillin/streptomycin, and 50 mM 2-ME. For luciferase assays, endotoxin-free (Promega PureYield) reporter plasmids were electroporated (260 V/950 μ F) in P5424 cells, as described. Briefly, 10^7 cells were washed and resuspended in 0.3 ml of serum-free RPMI 1640 along with 10 μ g of individual pGL3-E β luciferase reporter plasmids and 0.5 μ g of the control plasmid, pSV-RL (Promega). All transfections were performed three or more times with each of two independent plasmid preparations. In the absence of NF- κ B induction, transfected cells were cultured for 24 h, and 50 μ g of total protein from each transfectant was assayed for luciferase activity using the Dual-Luciferase Reporter system (Promega) and a Centro LB960 luminometer (Berthold Technologies). For NF- κ B induction, transfected cells were cultured 40 h, divided, and cultured an additional 8 h in normal medium or medium supplemented with PMA (50 ng/ml) and ionomycin (1 μ M) before harvesting.

Chromatin immunoprecipitation (IP) assays

Chromatin IP reactions were performed using the ChIP-IT Express Enzyme kit (Active Motif), according to the manufacturer's instructions. Briefly, P5424 cells (4.5×10^7) were cultured for 1 h in PMA (50 ng/ml) and ionomycin (1 μ M), and transcription factor-DNA complexes were cross-linked with formaldehyde and isolated chromatin was sheared by enzymatic digestion until $\geq 80\%$ of DNA fragments were < 500 bp in length. Protein-DNA complexes were immunoprecipitated overnight (4°C) from 10^9 cell equivalents sheared chromatin using protein G magnetic beads and Abs (Santa Cruz Biotechnology) specific for p65 RelA (sc-109) or upstream stimulatory factor 1 (USF1) (sc-229), or with normal rabbit IgG (110-4102; Rockland Immunochemicals). Chromatin immunoprecipitates were washed, and then chromatin was eluted from the magnetic beads

before cross-links were reversed and proteins were digested with proteinase K.

For real-time PCR, bound (5 μ l) and input (5 μ l of 1/10 dilution) samples from PMA/ionomycin-treated and untreated control cells were amplified using 1 \times iQ SYBR Green Supermix (Bio-Rad) and primers specific for the 5'-PD $\beta 2$ promoter (forward, 5'-CCCAAGGACATCTC CAAGCTCCTC-3'; reverse, 5'-GTTTCTTCCCCACAGGTGCCTACC-3') or 3'-PD $\beta 2$ promoter, respectively (forward, 5'-TTACCAGTTCTGGA GTAGATGGAG-3'; reverse, 5'-TAGGACGGTGAGTCGTGTCC-3'). Cycling parameters for 25- μ l reactions were 94°C, 30 s; 63°C, 30 s; 72°C, 30 s. Fold enrichment in the bound fractions was calculated from averages of triplicate reactions using the $\Delta\Delta$ Ct method previously described (29), and then normalized to that obtained for the nonspecific IgG-bound fraction.

Results

Developmental timing of D $\beta 2$ recombination

Tcrb recombination in developing thymocytes correlates with the appearance of sterile germline transcripts within the two DJC β gene segment cassettes. Despite transcription within both cassettes, D $\beta 2$ -J $\beta 2$ rearrangements have long been shown to accumulate more slowly than D $\beta 1$ -J β rearrangements (9–12), raising the possibility that accessibility of the D $\beta 2$ cassette may lag behind that of the D $\beta 1$ cassette during thymocyte development (7). To determine whether differences in DJ β recombination derive from delayed D $\beta 2$ expression relative to D $\beta 1$, we assessed the germline transcription status of each cassette in sorted DN subpopulations (Fig. 1). DN thymocytes account for $< 5\%$ of the normal mouse thymus. After immunodepletion of DP and CD4 or eight single-positive (SP) T cells, a pool of enriched DN cells from

30 C57BL/6 thymii was sorted into CD44⁺/CD25⁻ (DN1), CD44⁺/CD25⁺ (DN2), and CD44⁻/CD25⁺ (DN3) fractions (Fig. 1A).

Qualitative RT-PCR readily detected germline transcription at both DJC β cassettes in unsorted thymocytes from either RAG-1^{-/-} or RAG-2^{-/-} mice (Fig. 1B, lanes 1 and 2), but not in control RNA from the BW5147 T cell line (lane 7), in which both *Tcrb* alleles are rearranged, or in RNA from the M12 B cell line (lane 8). The majority of RAG-deficient thymocytes arrest as DN3 cells (30), well after the point at which D β 1-J β 1 recombination is first observed in wild-type DN1 cells (8). Consistent with such early D β 1 recombination, we found that D β 1 germline transcription was also present in wild-type DN1 cells (upper panel, lane 3), and continued throughout development (upper panel, lanes 3–6). Significantly, DN1 cells also supported transcription through the unrearranged D β 2 cassette (lower panel, lane 3), suggesting that DJC β 2, like DJC β 1, is accessible to transcription machinery during the earliest stages of T cell development. When normalized to a B cell-negative control, real-time amplification levels for each D β cassette (Fig. 1C) showed a peak in transcription through both D β cassettes at the DN2 stage of development. Consistent with the relative inefficiency of DJ β 2 recombination, germline DJ β 2 transcription remained constant in later stages of development, whereas DJ β 1 transcription markedly declined.

Initiation of D β 2 germline transcription

The D β 1 promoter, PD β 1, induces strong hypersensitivity to DNase I digestion (19, 31). No corresponding DNase-hypersensitive sites have been detected proximal to D β 2 (32). Nonetheless, the persistence of 1.0-kb germline D β 2 transcripts in mice lacking PD β 1 (31) strongly suggests the presence of a D β 2 promoter. We initially used Northern analyses to assess the origin of D β 2 germline transcription in DN thymocytes (Fig. 2B). Despite abundant transcription through C β 1 in both DN and DP thymocytes (R. McMillan and M. Sikes, unpublished observations), transcripts reading through sequence immediately 5' of D β 2 were only detected in DP thymocytes (lane 2). In contrast, a D β 2 3' probe readily detected message in RNA harvested from either RAG-2^{-/-} or RAG-2^{-/-}PD β 1^{-/-} DN thymocytes (lanes 3 and 4).

Unrearranged transcripts initiating at or upstream of D β 2 would yield an expected size of at least 1.6 kb. Indeed, trace amounts of 1.6-kb transcripts have previously been reported in CD25⁺/CD44⁺ thymocyte RNA (4). However, the enrichment of 1.0-kb transcripts in RAG-2^{-/-} RNA (DN, Fig. 2B, lane 3) and RAG-2^{-/-}PD β 1^{-/-} RNA (P1⁻, lane 4), together with previous studies in RAG-deficient animals (31) suggest that the bulk of D β 2 germline transcripts initiates 400–600 bp 3' of D β 2. We used RNase protection (Fig. 2C) and 5'-RACE (Fig. 2, D and E) to separately map transcription start site within the D β 2 cassette. An internally labeled 727-base antisense probe that extended from 94 bases 5' of D β 2 to the splice donor site of J β 2.1 (Fig. 2A) protected an array of transcription start sites in separate DN RNAs (Fig. 2C, lanes 3 and 4). Sequence analysis of cloned 5'-RACE products from P5424 RNA (Fig. 2D, lane 1) corroborated this diffuse pattern of transcript initiation in DN cells. Consistent with size expectations from the Northern blots, the great preponderance of cloned start sites mapped to a 150-nt region flanking J β 2.1 (Fig. 2E, ●). A tight cluster of start sites was mapped within the J β 2.1 coding sequence. None of the cloned sites was associated with consensus TATA, transcription initiator (*inr*), or downstream promoter elements (33).

The pattern of RNase protection in DN RNA was conserved in RNA from pooled DP and SP C57BL/6 thymocytes (Fig. 2C, lane 5). Unlike transcripts observed in RAG-deficient thymocytes (Fig.

2B) (31), early analyses of 1.0-kb D β 2 germline transcripts in wild-type mice suggested that they contained joined D β 2J β 2 segments (8). Our RNase protection assay revealed enrichment of multiple ~45- and ~90-base species (Fig. 2C, asterisks) predicted when transcripts that contain a DJ β 2 joint protect the individual D β 2 and J β 2.1 components of our internally labeled probe. Analysis of 5'-RACE products in DP/SP RNA confirmed the presence of DJ-joined segments as well as segments harboring germline and V(D)J joints. Moreover, the bulk of start sites identified in segments carrying a D β 2J β 2.1 joint was positioned 100 and 104 bp 5' of D β 2 within overlapping near-consensus initiator sequences 5'-PyPyA⁺1N^T/A₁PyPy-3' (33), whereas a single more distal start site was positioned 30 bp downstream from a perfect consensus TATA.

Promoter activity associated with D β 2

The differential use of start sites between germline and DJ-joined segments suggested that promoter elements might also be found both 5' and 3' of D β 2. To identify such promoter activities, we tested the ability of fragments upstream and downstream of D β 2 to direct expression of a luciferase reporter in the presence of the *Tcrb* enhancer, E β (Fig. 3). When normalized to a promoterless control, transient transfection of a 1.65-kb *StuI*/EcoRI fragment into RAG-1^{-/-} P5424 DN cells yielded only 2-fold promoter activity (-1104/+563 relative to the first base of D β 2), significantly less than that observed for PD β 1. However, 3' deletion of as little as 147 bp containing the majority of identified transcription start sites (-1104/+416) reduced promoter activity below that of the promoterless control. In contrast, sequential 5' deletion of all but the 147-bp start site fragment more than doubled promoter activity (+416/+563). Similar results were obtained using the SV40 enhancer or a variety of cell lines, including the 2017 pro-T and BW5147 mature T lines (R. McMillan and M. Sikes, unpublished data).

To mimic the impact of D-to-J recombination on promoter activity, plasmids bearing further truncations to +230, +98, and -61 were assessed. In fact, a fragment entirely 5' of D β 2 (-1104/-61) exhibited promoter activity ~7.5-fold above the promoterless control. When sequence between this upstream promoter activity and the downstream 147-bp promoter was deleted or replaced with a 310-bp fragment of DNA from the bacteriophage ϕ X174, activity from the two promoters appeared additive. From these functional assays, we conclude that D β 2 is flanked by two promoter elements (5'-PD β 2 and 3'-PD β 2), the upstream of which may be repressed until germline sequences are deleted by D-to-J recombination.

NF- κ B regulates promoter activity downstream of D β 2

Potential binding sites for multiple ubiquitous and lineage-restricted transcription factors were predicted within the 1.7-kb *StuI*/EcoRI D β 2 fragment (summarized in Fig. 4). Because germline promoter activity was restricted to sequences 3' of D β 2, we focused our initial analyses to this region that contained potential binding sites for Sp1, GATA-3, and NF- κ B. A duplex oligonucleotide probe spanning the predicted Sp1 binding site 105 bp 3' of D β 2 bound two distinct protein complexes in a P5424 nuclear protein extract (Fig. 5A, lane 1). Both complexes were specifically inhibited by competition with 100-fold excess unlabeled probe (lane 2), whereas mutation of the Sp1 motif severely impaired its ability to compete for protein binding (lane 3). The presence of Sp1 in each of the nucleoprotein complexes was confirmed by their inhibition upon addition of an Sp1-specific Ab (lane 4), but not upon addition of nonspecific IgG (lane 5). Longer exposure of the Ab reactions revealed a single Sp1-specific supershifted species (lane 6).

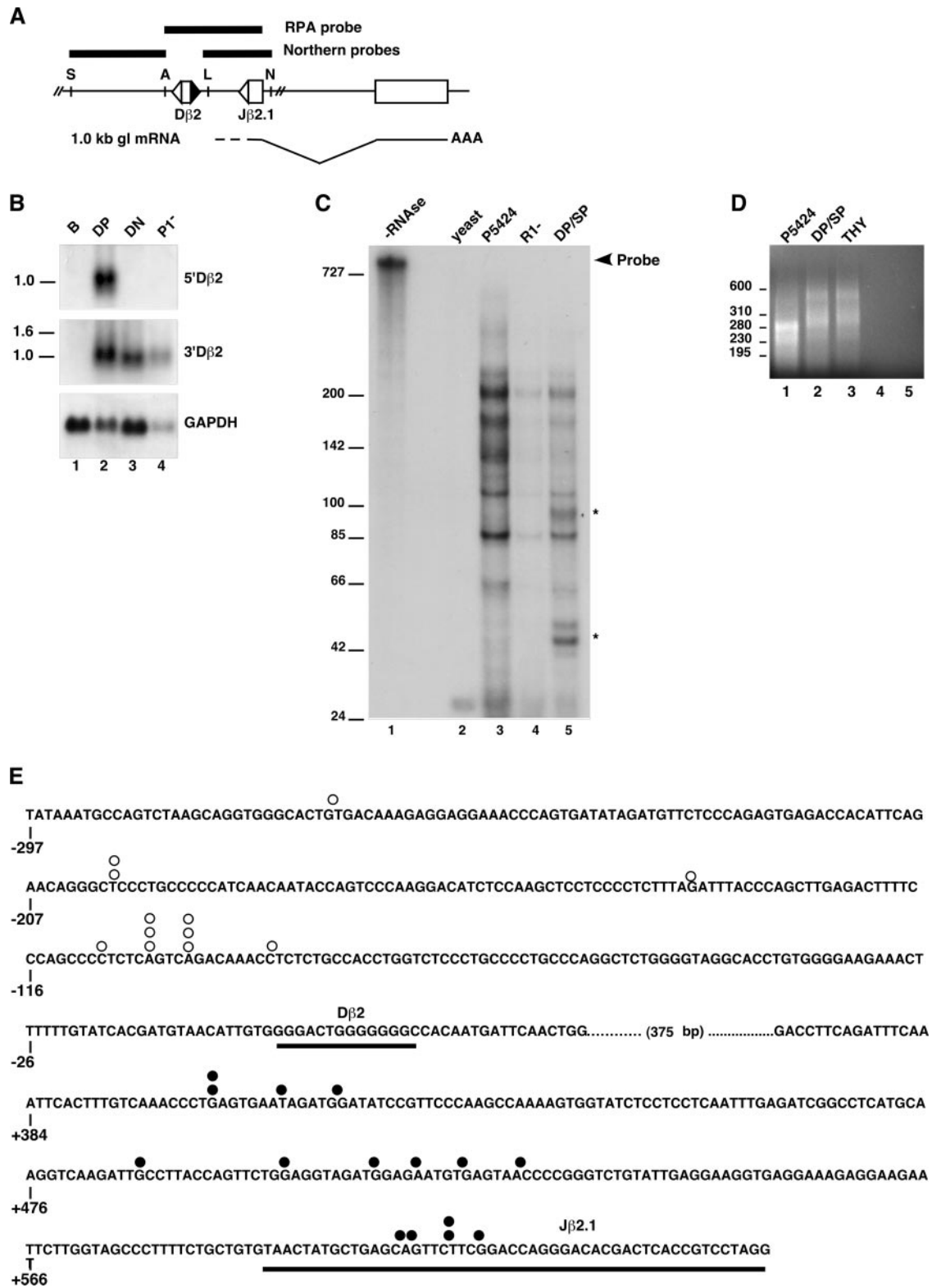
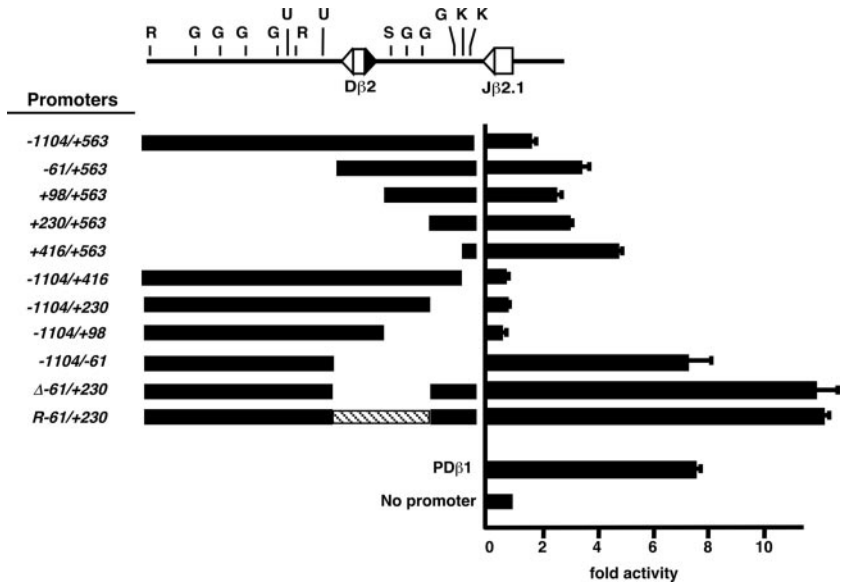


FIGURE 2. Initiation of $D\beta 2$ germline transcripts. *A*, Schematic of $D\beta 2$ region. Probes used in *B* (bars above schematic) and *C* (bar below schematic). S, *Stu*I; A, *Alu*NI; L, *Afl*III; and N, *Eco*NI. *B*, Autoradiograms of Northern hybridization with the indicated probes to RNAs from M12 (B) pooled DP and SP thymocytes (DP/SP), $RAG-2^{-/-}$ thymocytes (DN), and $RAG-2^{-/-}PD\beta 1^{-/-}$ thymocytes ($P1^{-}$). The level of GAPDH expression serves as a loading control (*lower panel*). *C*, RNase protection of a 750-base antisense probe by $D\beta 2$ germline transcripts in RNA isolated from separate $RAG-1$ -deficient DN sources (P5424 cells or $RAG-1^{-/-}$ thymocytes) or pooled DP/SP thymocytes. Asterisks mark protected species enriched in DP/SP RNA. *D*, Agarose gel electrophoresis of 5'-RACE products obtained from the indicated RNA sources, or from wild-type thymus in the presence of the 5'-RACE primer alone (*lane 4*) or the 3' primer alone (*lane 5*). *E*, Map of the 5' ends from individual RACE clones harboring germline (●) or $D\beta 2/J\beta 2.1$ joined sequences (○). $D\beta 2$ and $J\beta 2.1$ coding sequences are underlined.

FIGURE 3. Promoter activities associated with D β 2. The indicated DNA fragments were inserted upstream of the luciferase cassette in pGL3-E β . Fragment positioning is relative to the first base of the D β 2 coding region (+1). Each plasmid was co-transfected with pSV-RL into P5424 cells, and protein extracts were assayed for luciferase activity 24 h after transfection. Values from four independent transfections were normalized to *Renilla* controls. Bars represent mean normalized luciferase activity \pm SD relative to a promoterless control. Activity from the upstream PD β 1 promoter is provided for comparison. Deletion or substitution of sequences between -61 and +230 in the full-length promoter fragment (-1104/+563) is respectively indicated (Δ -61/+230) or (*R*-61/+230). The positions of putative Runx (R), GATA (G), USF1 (U), Sp1 (S), and NF- κ B (K) binding sites are schematized above the graph.



Binding of the T lineage-specific GATA transcription factor, GATA-3, has been demonstrated previously in both PD β 1 (25, 34) and E β (35, 36). Three separate GATA binding sites were predicted between 145 and 415 bp 3' of D β 2, although the third site contained mismatches in both bases flanking the core GATA. Sep-

arate oligonucleotide probes spanning the G3-5 and G3-6 sites formed multiple nucleoprotein complexes with P5424 protein extracts (Fig. 5B, lane 1), whereas the G3-7 site failed to demonstrate protein binding (data not shown). In each case, one major complex was fully competed and another weakly competed by an excess

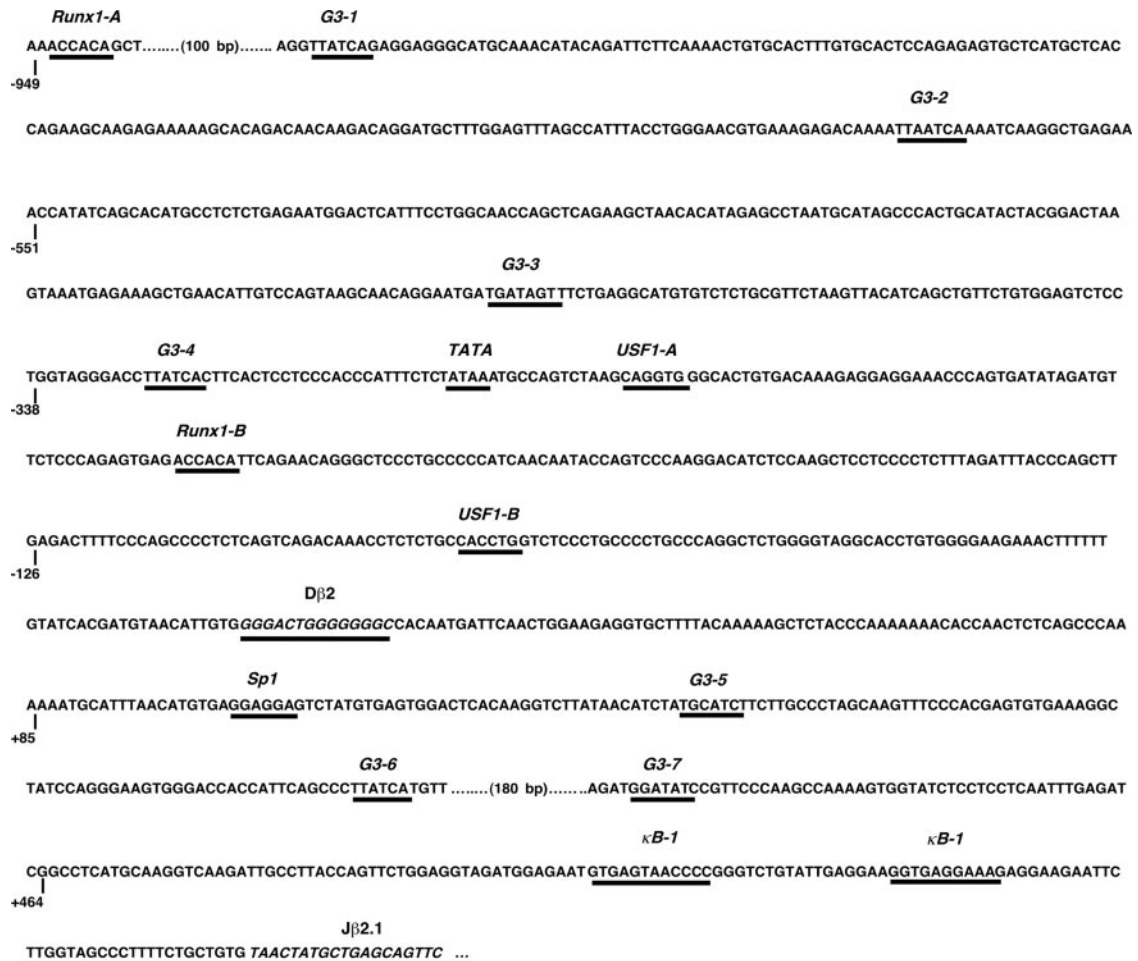


FIGURE 4. Organization of the D β 2 regulatory region. Potential transcription factor binding sites and D β 2 and J β 2.1 coding sequences are shown. Numbering is relative to the first base of the D β 2 coding sequence (+1). USF (USF1), G3 (GATA-3), κ B (NF- κ B).

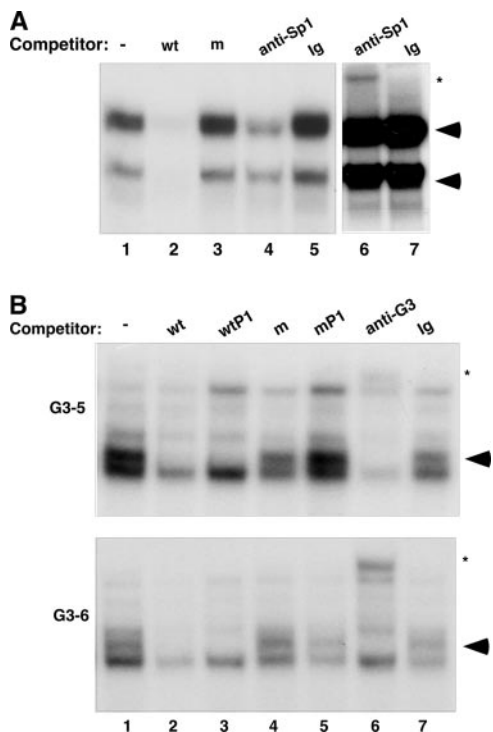


FIGURE 5. Sp1 and GATA-3 bind downstream of *Dβ2*. Nuclear extracts from the P5424 cell line were incubated with radiolabeled probes to the Sp1 (A) or G3-5 and G3-6 sites (B) alone (lane 1), in the presence of 100-fold molar excess of the indicated unlabeled competitors (A, lanes 2 and 3; B, lanes 2–5), or in the presence of the indicated Abs (A, lanes 4–7; B, lanes 6 and 7). Nucleoprotein complexes were resolved on a 5% acrylamide gel and visualized by autoradiography. Lanes 6 and 7 in A represent longer autoradiographic exposure of lanes 4 and 5. Specific protein-DNA complexes (arrowheads) and Ab-supershifted complexes (asterisks) are indicated.

of unlabeled probe (lane 2) or a GATA-3 binding site within *PDβ1* (lane 3). Neither oligonucleotide was effective at competing the major binding activities when its GATA core was mutated (lanes 4 and 5). The specificity of both primary protein complexes with G3-5 and G3-6 was confirmed by their supershift in the presence of a GATA-3 Ab, whereas other minor complexes were not specifically attenuated by the anti-GATA-3 Ab (lane 6).

Luciferase analyses suggested that the bulk of promoter activity 3' of *Dβ2* lies within a 147-bp span bounded by sites for the *EcoRV* and *EcoRI* restriction enzymes (Fig. 3). This fragment contains no identifiable TATA, *inr*, or downstream promoter elements. However, two potential binding sites for the NF- κ B family of transcription factors are positioned proximal to the *Jβ2.1* RSS (Fig. 4). Although initial gelshifts detected protein binding to probes for either of the predicted NF- κ B sites using P5424 nuclear extracts (Fig. 6A, lane 1), binding was strongly induced after P5424 cells were treated for 4 h with either PMA/ionomycin (lane 2) or overnight with LPS (data not shown). In each case, nucleoprotein complexes were specifically inhibited by incubation with 100-fold excess unlabeled probe from either site (lanes 3 and 6). Binding at the downstream κ B-2 probe was only weakly competed by unlabeled κ B-1. Competition by either unlabeled probe was abolished upon mutation of the 3' half-site within each putative κ B motif (lanes 4 and 5). The specificity of each site for p65 RelA was confirmed by inhibition of each nucleoprotein complex in the presence of p65-specific Ab (lane 8). In contrast, nuclear protein binding to double-stranded probes containing an unre-

lated NF- κ B binding site was unaffected by addition of anti-p65 Ab (lower panel, lane 8). Neither Abs to a second NF- κ B protein, p50, nor nonspecific IgGs altered the binding pattern at either κ B-1 or κ B-2 (lanes 9 and 10).

Collectively, gelshift results indicate a complex array of transcription factor binding sites arranged between *Dβ2* and *Jβ2.1*. In our initial reporter studies, a 320-bp region containing the Sp1, G3-5, and G3-6 binding sites was dispensable for 3'-*PDβ2* promoter activity (Fig. 3). The G3-7, κ B-1, and κ B-2 sites are positioned within the 147-bp fragment that exhibited peak promoter activity 3' of *Dβ2*. To assess the contributions of these three sites to promoter function, we selectively mutated each binding motif in the +416/+563 promoter fragment. Consistent with its failure to bind nuclear proteins in EMSA studies, mutation of the G3-7 site had no impact on promoter activity in transfected P5424 cells (Fig. 6B). Mutation of either NF- κ B binding site, however, dramatically impaired promoter function, with luciferase activity reduced to 38 and 49% of the wild-type fragment, respectively. When both κ B sites were destroyed, resultant promoter function was <20% of wild type.

Although gelshift analyses showed p65 RelA binding to the κ B-1 and κ B-2 sites in the nuclei of unstimulated P5424 cells (Fig. 6A), binding activity was markedly enhanced by PMA/ionomycin treatment. To determine whether the NF- κ B-dependent activity of 3'-*PDβ2* was similarly augmented, we measured the ability of transfected 3'-*PDβ2* fragments to direct luciferase reporter activity in response to PMA/ionomycin. When normalized to promoter activity in untreated cells 48 h after electroporation, addition of 50 ng/ml PMA and 1 μ M ionomycin during the final 8 h of culture led to a ~3-fold increase in 3'-*PDβ2* activity (Fig. 6C). In contrast, activity of the 5'-*PDβ2* promoter was not significantly affected by PMA/ionomycin treatment. Significantly, PMA/ionomycin treatment failed to rescue activity of the 3'-*PDβ2* promoter when the κ B-1 and κ B-2 sites were destroyed. NF- κ B has not, to date, been associated with activation of any of the defined *Tcr* promoters.

We next used chromatin IP to determine whether the κ B-1 and κ B-2 sites in 3'-*PDβ2* are bound by NF- κ B in vivo. P5424 cells were stimulated for 1 h with PMA/ionomycin before being treated with formaldehyde to cross-link protein-DNA complexes. Nuclear lysates were digested to enrich for mono- and dinucleosomes, and immunoprecipitated using Abs specific for RelA and USF1 and protein G magnetic beads. Purified DNA was amplified by real-time PCR using primers specific for the 3' and 5' *Dβ2* promoters (Fig. 6D). The enrichment of each promoter amplicon in the bound fractions was normalized to that immunoprecipitated with total serum IgG. Although modest levels of RelA binding were detected at both the 5' and 3' *Dβ2* promoters of untreated cells, PMA/ionomycin specifically induced a strong association of RelA with sequences that spanned the κ B-1 and κ B-2 sites in 3'-*PDβ2*. In contrast, binding of USF1 to its putative target sequences in the 5'-*PDβ2* was not enhanced by PMA/ionomycin treatment. The modest increase in USF1 association with 3'-*PDβ2* after PMA induction may indicate the presence of cryptic binding sites proximal to the 3' promoter amplicon, although it most likely derives from proximity of the 5' and 3' promoters and the presence of larger oligonucleosome arrays in the IP reactions. Although we cannot exclude the possibility that the G3-5, G3-6, or Sp1 sites play essential roles in directing transcription through the *DJβ2* cassette in developing thymocytes, our studies reveal that promoter activity in the germline sequences downstream of *Dβ2* is primarily dependent on the actions of NF- κ B.

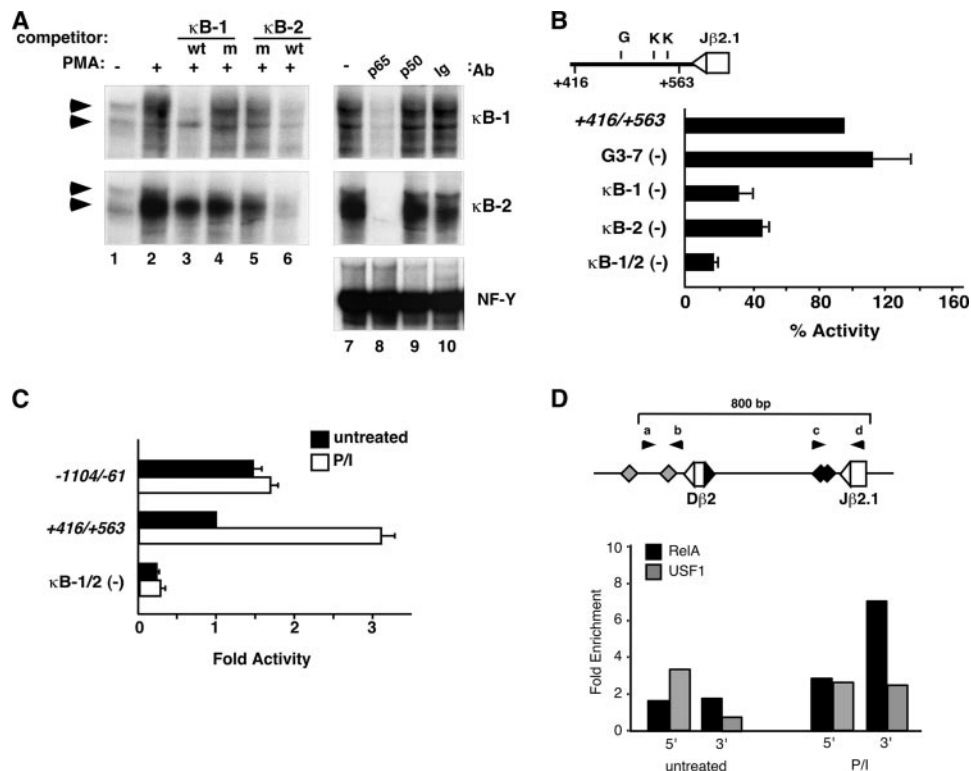


FIGURE 6. NF- κ B contributes to promoter activity 3' of $D\beta 2$. *A*, Radiolabeled probes to two potential NF- κ B binding sites 3' of $D\beta 2$ (upper and middle panels) were incubated with nuclear extracts from untreated (lane 1) or PMA/ionomycin-activated P5424 cells (lanes 2–10) alone (lanes 2 and 7), or in the presence of 100-fold molar excess of the indicated unlabeled competitors (lanes 3–6), or in the presence of the indicated Abs (lanes 8–10). As a control or Ab and nuclear extract quality, a radiolabeled probe to the NF-Y transcription factor was separately incubated with PMA/ionomycin-activated P5424 nuclear extract and each of the indicated Abs (lower panel, lanes 7–10). Nucleoprotein complexes were resolved on a 5% acrylamide gel and visualized by autoradiography. *B*, Individual transcription factor binding sites within the minimal downstream PD $\beta 2$ fragment (+416/+563) were selectively mutated to assess their contribution to downstream promoter activity. P5424 transfectants of the indicated mutant pGL3-E β (+416/+563) reporters were assessed. Results from six independent cotransfections with pSV-RL were normalized to *Renilla* levels, and expressed as percentage of that obtained for wild-type pGL3-E β (+416/+563). *C*, Each of the indicated luciferase reporter plasmids was transfected into P5424 cells. Transfected cells were allowed to recover for 40 h before 8-h treatment with PMA/ionomycin, and then protein extracts were assayed for luciferase activity a total of 48 h after transfection. Values from PMA/ionomycin-treated (\square) and untreated controls (\blacksquare) for four independent transfections were normalized to those obtained for untreated 3'-PD $\beta 2$ (+416/+563). Bars represent mean (\pm SEM) for *B* and *C*. *D*, Chromatin IP analyses of NF- κ B RelA and USF1 association with the 5' (primers *a* and *b*) and 3' $D\beta 2$ promoter sequences (primers *c* and *d*) of endogenous P5424 *Tcrb* before and after PMA/ionomycin treatment. Relative positions of the two NF- κ B (\blacklozenge) and predicted USF1 (gray diamonds) binding sites are indicated on the schematic diagram of the $D\beta 2$ region. Changes in the average Ct values of triplicate amplifications for specific Ab-bound samples relative to input controls were normalized to total serum IgG and transformed to generate fold enrichment over the isotype control. The results are representative of two separate experiments.

Organization of promoter activity upstream of $D\beta 2$

Although the GATA binding sites downstream of $D\beta 2$ appear dispensable for germline promoter activity, GATA-3 *trans* activation is essential for activation of the PD $\beta 1$ promoter (25) and E β (8). Motif searches of the -1104/-61 promoter fragment identified in Fig. 3 indicated a cluster of four GATA binding sites within a 415-bp stretch of sequence 5' of $D\beta 2$ (Fig. 4), three of which conform to the canonical GATA motif, WGATAR (37), whereas the fourth (G3-2; TGATTA) is mismatched at a single nucleotide. The promoter region upstream of $D\beta 2$ also contains multiple potential E boxes, as well as consensus Runx binding sites at -948 and -219 (Fig. 4). The Runx family of transcription factors plays essential and diverse roles in T cell development, binding to canonical RACCRC sites within the enhancers of all four TCR loci (38). Indeed, recent chromatin IP suggested that in addition to binding known sites in E β , the Runx1 protein recognizes unidentified sites flanking both $D\beta 1$ and $D\beta 2$ (39).

To identify the minimal 5'-PD $\beta 2$ promoter, we transfected a panel of pGL3-E β luciferase vectors that contained progressive deletions of the -1104/-61 promoter fragment into cultured

P5424 cells. Initial 3' deletion of 142 bp that contained the USF1-B site, both putative initiator elements, and all but one of the upstream transcription start sites identified by 5'-RACE reduced promoter activity to 37% of the -1104/-61 fragment (Fig. 7, -1104/-203). Additional 3' deletions of 372 bp (removing all but the Runx1-A and G3-1 sites) or 520 bp (leaving only the Runx-A site) progressively attenuated promoter activity to 18 and 2% of the full-length -1104/-61 fragment, respectively. Conversely, progressive 5' deletions had no negative effect on activity of the 5'-PD $\beta 2$ promoter. In fact, the -280/-61 fragment that lacked the Runx1-A site and all four potential GATA sites exhibited slightly higher promoter activity than the parental fragment. As such, our deletion assays suggest nominal roles for the four putative GATA elements in 5'-PD $\beta 2$ activity. Consistent with our findings, none of the putative GATA sites was detected in a search of the human $D\beta 2$ upstream region (data not shown). Conversely, the sequence and positioning of both putative USF1 binding sites and paired initiator elements were strongly conserved. Among the two potential Runx elements, only the upstream Runx-A site was conserved between mouse and human.

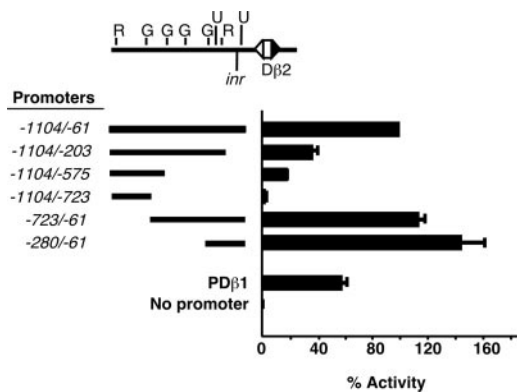


FIGURE 7. Promoter activity 5' of D β 2. The indicated DNA fragments were inserted upstream of the luciferase cassette in pGL3-E β . Fragment positioning is relative to the first base of the D β 2 coding region (+1). Each plasmid was cotransfected with pSV-RL into P5424 cells, and protein extracts were assayed for luciferase activity 24 h after transfection. Results from six independent cotransfections with pSV-RL were normalized to *Renilla* levels, and expressed as percentage of that obtained for pGL3-E β (-1104/-61). Bars represent mean (\pm SEM). The positions of putative Runx (R), GATA (G), or USF1 (U) binding sites and transcription initiator (*inr*) elements are schematized above the graph.

Discussion

A growing body of evidence has established a role for germline promoters in regulating the recombinational accessibility of discrete domains within individual Ag receptor loci. Promoters are proposed to provide a conduit through which enhancer-dependent chromatin alterations drive accessibility of promoter-proximal gene segments. Such promoter-dependent activities as recruitment of *switch/sucrose nonfermentable* (SWI/SNF) chromatin-remodeling complexes (40), initiation of germline transcription (24), and changes in CpG methylation (31) have all been implicated in efficient gene segment assembly. In this study, we have characterized the developmental profile of D β 2 transcriptional activation, and have identified separate promoter activities positioned 5' and 3' of D β 2. Significantly, whereas the downstream promoter appeared to direct transcription of germline J β 2 sequences throughout DN thymocyte development, transcription from the upstream promoter was only detected in DNA harboring D β 2J β 2 joints. We suggest that differential activation of the two D β 2 promoters may play a central role in coordinating the relative efficiencies with which the two DJ β cassettes are assembled.

The two D β 2 promoters appear structurally and functionally distinct from one another. Indeed, the 3'-PD β 2 bears little homology to any of the identified TCR germline promoters, being poorly organized and with no discernable core promoter elements. The structural simplicity of 3'-PD β 2 may account for the decentralized manner of transcriptional initiation downstream of germline D β 2. In sharp contrast, transcription directed by the 5'-PD β 2 is tightly clustered around two potential initiator elements positioned 38 bp downstream of a TTAGATT palindrome that is both a perfect match to the canonical *inr* and a single-base mismatch to the canonical TATA. Moreover, whereas activity of the upstream promoter is localized to a 340-bp sequence that contains potential binding sites for Runx1, and USF1 regulatory proteins, RelA-containing NF- κ B dimers may be sufficient to drive germline transcription from the downstream promoter.

Initial analysis of promoter activity upstream of D β 2 suggests that the four putative GATA binding sites and distal putative Runx binding site may not contribute to the activity of 5'-PD β 2 (Fig. 7). Indeed, luciferase expression in transfected cells was modestly el-

evated upon deletion of the Runx1-A and G3-1-to-4 sites. Recent studies by Oltz and colleagues (14) demonstrated localization of both Runx1 and GATA-3 to sequences immediately upstream of D β 2 5' in thymocytes of RAG-2-deficient mice, suggesting a role for Runx1 and GATA-3 in the regulation of D β 2 usage in vivo. As such, the insensitivity of our luciferase readouts to 5' truncations of the upstream promoter could reflect either the limitations inherent to large-scale deletion analyses and transient transfections. Alternatively, the gain in luciferase activity observed in 5'-PD β 2 constructs from which the distal sequences were deleted could suggest a more complex architecture in which a mixture of positive and negative effectors coordinately regulates the core 5'-PD β 2.

A role for germline promoters in controlling the recombination of downstream gene segments has been supported in the *Tcra* (20, 21, 24), *Tcrb* (19), *Tcr delta* (41), and *IgH* loci (22). Deletion of the J49 α promoter selectively impaired assembly of J segments 3' of the deletion without significantly affecting the recombination potential of upstream Js (21). Likewise, D-to-J recombination in stably transfected substrates was progressively attenuated by the repositioning of PD β 1 at increasing distances 3' of D β 1 (23). In light of the localization of germline promoter activity between D β 2 and J β 2.1, the effects of PD β 1 repositioning on DJ recombination bear a marked similarity to the relative inefficiency with which D β 2J β 2 joints are made in vivo (9–12). Alternatively, D β 2 recombination levels relative to D β 1 could reflect the presence of less efficient RSS sites within the DJC β 2 cassette, or a reduced ability of the D β 2 region to recruit enhancer-dependent chromatin alteration. Our analyses of germline transcription argue against the possibility that D β 2 accessibility is developmentally delayed until after T lineage commitment in DN2 cells (7).

What evolutionary advantage would drive the unique organization of D β 2 promoter activity? D-to-J joints in *Tcrb* are rarely out of frame. By contrast, two in every three V-to-DJ β joints will contain a frameshift that renders the assembly nonfunctional. In New Zealand White mice that lack the DJ β 2 gene segments, the frameshift potential of V-to-DJ β joining would theoretically block ~44% of early thymocytes from progressing through the $\alpha\beta$ lineage beyond β selection. However, the positioning of two complete DJC cassettes downstream of the V β elements would offer each allele an attempt to rescue nonproductive V(D)J joints involving D β 1. Such a selective advantage to $\alpha\beta$ development would only be realized if the initial DJ substrates for V β recombination were preferentially assembled from the upstream DJ cassette, because V-to-DJ β rearrangements delete the D β 1 sequences. Therefore, positioning of the germline promoter downstream of D β 2 could provide the programmed inefficiency in DJ β 2 rearrangement necessary to maximize the opportunities to assemble a functional *Tcrb* gene.

NF- κ B activation by the pre-TCR has previously been shown to be essential for the survival and progression of DN thymocytes to the DP stage (42). Likewise, the RelA and p50 subunits of NF- κ B are required for early lymphopoiesis, most likely providing protection against TNF-induced apoptosis (43). In contrast to such classical pathway-mediated activation, previous studies showed that RelA and p50 were both constitutively localized to the nucleus of wild-type (44) and RAG-2-deficient thymocytes, although at levels substantially below those in DP cells (45). Consistent with our findings in P5424, the levels of nuclear NF- κ B were greatly diminished in DN thymocytes incubated in vitro at 37°C, but remained readily inducible by PMA and ionomycin (45). A potential role for NF- κ B in 3'-PD β 2-mediated accessibility of D β 2 to recombinase would seem to necessitate such constitutive nuclear localization. However, it is intriguing to consider whether the capacity to augment NF- κ B levels in developing thymocytes might

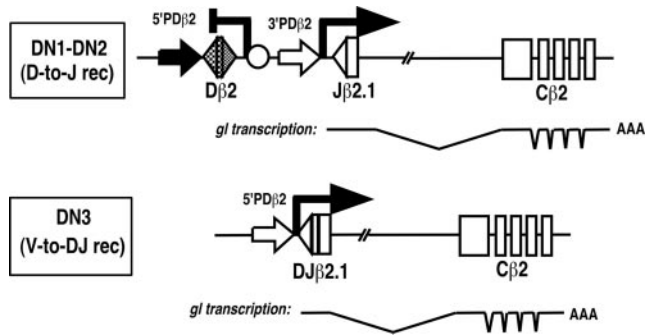


FIGURE 8. Model of transcriptional regulation at *Dβ2*. Schematic depictions of the germline (*top*) and assembled *Dβ2* and *Jβ2.1* gene segments (*bottom*). Block arrows and circle represent the 5'- and 3'-PDβ2 promoters and 5'-PDβ2 repressor element. Shading in the germline elements indicates partial (stippled) or complete transcriptional and recombinational inactivity (filled). The model proposes that repression of 5'-PDβ2 during DN1/DN2 development relegates transcription of the *DJβ2* gene segment cluster to the 3'-PDβ2 promoter, which only weakly provides recombinational accessibility to the *Dβ2* 3'-RSS. Upon deletion of the repressor and 3' promoter during D-to-J recombination, 5'-PDβ2 is strongly activated and provides *Dβ2* 5'-RSS accessibility for V-to-DJ joining in DN3-staged cells.

allow for select stimuli to impact *Dβ2* usage in the overall TCRβ repertoire.

The dependence of Vβ recombination on formation of a DJβ joint has limited study of the role promoters might play in V-to-DJ assembly. Nonetheless, the breadth of sequence that separates Vβ elements from the DJCβ cassettes, together with the dependence of *Tcra* assembly on *Jα* promoters (24), strongly suggests an analogous role for *Dβ* germline promoter activity in Vβ-to-DJβ assembly. Consequently, rearrangement of *DJβ2* joints with upstream Vβ elements would require that a second *Dβ2* promoter reside upstream of *Dβ2*, and be active only after DJ assembly. Our finding that transcription initiates predominantly downstream of *Dβ2* before recombination and upstream of *DJβ2* joints (Fig. 2) corroborates longstanding Northern analyses in wild-type (4, 46) and RAG-deficient thymocytes (31), and strongly supports such a model in which dual promoters sequentially drive inefficient D-to-J joining and then V-to-DJ joining (Fig. 8).

It remains unclear how activity of the 5'-PDβ2 promoter is limited before DJβ recombination. Deletion studies suggest that elements within a 100- to 300-bp region between the two promoters selectively repress activity of the upstream promoter without affecting the downstream promoter (Fig. 3). Indeed, a similar cryptic repressive activity immediately downstream of *Dβ1* was suggested to modestly impair PDβ1 activity (34). EMSA confirmed binding sites for the Sp1 and GATA-3 transcription factors within the region of the putative 5'-PDβ2 repressor (Fig. 5). Although Sp1 and GATA-3 have both been defined primarily as transcriptional activators, each has the capacity to drive transcriptional repression (47, 48). However, reporter constructs from which fragments containing the Sp1 and G3-5 and G3-6 sites had been deleted retained their ability to repress 5'-PDβ2 activity (Fig. 3, -1104/+98). It remains to be seen whether either element contributes to repression of 5'-PDβ2 activity in a chromatin setting. Indeed, many or all of the sites surrounding *Dβ2* could function within the endogenous chromatin as part of a single complex that preferentially targets transcription start sites 3' of *Dβ2* until they are eliminated by D-to-J recombination. Alternatively, other as yet undefined elements between *Dβ2* and *Jβ2.1* may contribute to repression of the upstream promoter, or DJ recombination could relieve topological restraints that prevent occupancy of the upstream promoter. Oltz

and colleagues (39) have recently shown that both Runx1- and TATA-binding protein are recruited to targets upstream of the unrearranged *Dβ2* sequences in RAG-deficient thymocytes, and coordinate physical interaction between the *Dβ2* and *Eβ* elements. Their findings, coupled with the general openness of chromatin within the *DJβ2* cassette of P5424 cells (49), argue against a model in which the upstream promoter is epigenetically silenced by chromatin condensation. Future studies will be necessary to unravel the complex regulatory schemes that govern differential promoter activation at *Dβ2*, and to assess the potential impact of each element on *Tcrb* assembly.

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Disclosures

The authors have no financial conflict of interest.

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