A novel Pax5-binding regulatory element in the Igκ locus

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INTRODUCTION

The B cell receptors (BCRs) are encoded in the mouse genome by the three immunoglobulin (Ig) loci, the IgH heavy chain locus, and the two light chain loci, Igκ and Igλ. In their germline conformations, the Ig loci do not give rise to functional proteins. It is only through a tightly regulated process of genome editing, termed V(D)J recombination, that the loci are reconfigured to allow transcription of an Ig gene in B cells. During the recombination process, the variable (V), diversity (D), and joining (J) segments are cleaved by the RAG complex and joined together into one continuous segment by the DNA repair machinery (1). Each rearrangement utilizes a single V, D (in the heavy chain), and J segment, and each B cell contains one productively rearranged heavy chain and light chain. In this way, B cells give rise to the multitude of antigen recognition specificities which constitutes the adaptive immune system.

The recombination of the different loci takes place in a developmentally staggered manner, with the IgH locus undergoing VDJ recombination first in the pro-B cell stage (2). Light chain rearrangement normally takes place only after a successful IgH rearrangement, which allows the cells to differentiate to the pre-B cell stage (3). In mice, the Igκ locus is the primary source for the BCR light chain and will undergo preferential rearrangement. The recombination of the different loci is kept tightly separated, despite the fact that the enzymatic machinery responsible for the processes is essentially the same and is present at both the pro- and pre-B cell stages. The light chain loci are maintained in an inaccessible chromatin state via epigenetic mechanisms prior to the pre-B cell stage, at which point they become available to the rearrangement machinery (3, 4). One such epigenetic mark is DNA methylation, a mark that is established at the Igκ locus during early embryonic development and which is hereditarily maintained during cell division (5). DNA methylation has been shown to block the activity of the rearrangement machinery in vitro (6). The Igκ locus undergoes selective demethylation at the pre-B cell stage, immediately prior to rearrangement (5, 7, 8). The rearranged Igκ allele is unmethylated from that stage onward, while alleles which do not undergo rearrangement remain methylated, even at the mature B cell stage. The low level of methylation is significant for an additional stage of Igκ editing during B cell development, namely for efficient somatic hypermutation (SHM), which will allow affinity maturation of the BCR in activated mature B cells (9). Methylated pre-rearranged Igκ sequences do not undergo proper SHM at this stage, whereas identical unmethylated sequences do (10).

The stage-specific transcription, rearrangement, and chromatin structure of the Igκ gene is mediated by regulatory sequences within and in proximity to the locus. The locus contains three characterized enhancers, including an intronic enhancer (iEκ) (11), located in the intron between the Jκ segments and the Cκ exon and two enhancers situated a few 1000 bases downstream of the Cκ exon, termed 3′Eκ (12) and Ed (13). These enhancers work in cooperation to promote stage-specific chromatin accessibility, DNA demethylation, V to J rearrangement, heightened transcription of the locus, and SHM in activated B cells, with different enhancers contributing to a varying extent to each one of these processes. iEκ and 3′Eκ have been implicated in promoting accessibility and rearrangement of the locus in pre-B cells (14–16), while 3′Eκ and Ed strongly effect the level of transcription and SHM in mature B cells (17, 18), neither of which is significantly affected by the deletion of iEκ (14, 18). All of the three enhancers contribute together to the demethylation of the locus (16, 19). Replacement of iEκ with the IgH intronic μ enhancer is enough to change the rearrangement timing of the locus to the earlier pro-B cell stage.
showing that it is indeed these sequences which direct the temporal precision of the developmental program (20).

Other than the enhancers, there are a number of additional regulatory elements surrounding the Igκ locus, increasing the complexity of the regulation. The recently discovered HS10 element, which lies downstream of Ed, appears to mostly function in plasma cells. While itself being a weak enhancer, HS10 acts as a co-enhancer to strengthen the activities of 3’Eκ and Ed (21). A matrix attachment region (MAR) lies immediately adjacent to iEκ and mediates connections between the locus and the nuclear matrix (22).

The activities of the cis regulatory elements are mediated by various transcription factors, which either activate or repress the enhancer activity. Many of these transcription factors are master regulators of the B cell lineage, which are important for maintaining B cell identity, such as E2A and PU.1. The binding sites of iEκ and 3’Eκ and substantially contribute to the enhancer activity (23–27). However, binding of Pax5, a master regulator of B cell identity, has been surprisingly missing from these enhancers in mature B cells. While binding sites have been identified in 3’Eκ (24, 25, 28), as well as in K-I and K-II (29, 30), which are regulatory regions (RR) upstream of the Jκ segments, Pax5 plays an inhibitory role in this context and is released during the pre-B cell stage when the locus is activated. This is despite the fact that Pax5 itself is necessary for the active induction of the locus (31).

In this work, we characterize a region adjacent to the MAR/iEκ elements. We had previously identified this element as a participant in the demethylation process of the Igκ locus in cell culture and thereby designated it Dm (32). Here, we find that this element binds Pax5 in B cell stages from the pre-B cell stage and onward. It is necessary for demethylation of a pre-rearranged Igκ transgene, but deletion of the element in the endogenous locus does not affect the demethylation process. We find that the element contributes to efficient SHM of the Igκ locus, indicating that the Dm element functions at more than one stage of B cell development.

MATERIALS AND METHODS

MICE

Targeted ES cells were generated for 10 generations on a BALB/c background. IgκΔDm/ΔDm mice were bred with wild-type (WT) BALB/c to produce IgκWT/ΔDm mice. Human Cκ knock-in mice (33) (gift from M. Nussenzweig) were bred with either WT BALB/c or ΔDm BALB/c to produce IgκWT/WT Cκb/m and IgκWT/ΔDm Cκb/m mice, respectively. IgκWT/ΔDm mice were bred with CAST/EiiJ (Cast) mice (Jackson Laboratory) to produce BALB/c/Cast IgκΔDmWT and BALB/c/Cast IgκΔDmWT littermates. Rag1Δ−/− mice (Jackson Laboratories) were bred onto a Cast background. IgκΔDm/ΔDm were bred onto a C57BL/6 Rag1Δ−/− (B6) background containing the 3H19 Igκ chain transgene (IgκH+). CAST/EiiJ Rag1Δ−/− mice were bred with C57BL/6 Rag1Δ−/− Igκ+ either with or without a deletion of the Dm element, giving rise to B6/Cast Rag1Δ−/− Igκ+ IgκΔDmWT and B6/Cast Rag1Δ−/− Igκ+ IgκWTWT mice, respectively. Mice were housed in specific pathogen-free conditions at the Hebrew University Medical School animal facility. Transgenic mouse lines Lk, LkΔDm, and LkΔ70 were produced, using the constructs described in the Section “Targeting Constructs,” at the Hadassah Hospital Medical School Transgenic Unit.

Two independent founder lines were produced for the Lk transgene, four for the LkΔDm and three for the LkΔ70. The copy number of the transgene for each founder line varied from low (two insertions) to high (20 insertions) with most lines having a moderate number of insertions (four to eight insertions). All animal procedures were approved by the Animal Care and Use Committee of the Hebrew University of Jerusalem.

TARGETING CONSTRUCTS

The LkΔDm construct was prepared using the following steps: the 4.3-kb KpnI–KpnI fragment, containing a unique XbaI–Csx sequence, was excised from the Lk plasmid (34) and cloned into the KpnI site of the Bluescript vector which was modified to destroy the polylinker XbaI site. The resulting pBSKpn2 plasmid was cut at unique compatible XbaI and NsiI sites, and recircularized, resulting in the deletion of 930 bp Xbal–NsiI Dm fragment from the jCκ intron. The KpnI–KpnI Dm-deleted fragment was excised and reinserted into the Lk plasmid, resulting in the LkΔDm construct.

The LkΔ70 construct was prepared using the following steps: the HindIII–blunt TaqI 2.6-kb fragment, containing the germline Jκ region, was cloned into HindIII–EcoRV sites of the Bluescript vector. Next, a blunt BstEII–BglII 2-kb fragment containing the Cκ exon was cloned into blunt EcoRI–BamHI sites of the previously described Jκ containing Bluescript vector to yield the pΔ70 construct, which had the 70 bp TaqI–BstEII deletion introduced into the HindIII/BglII 5.6-kb jCκ germline sequence. The 1-kb intact intronic Xbal–HindIII region of pBSKpn2 plasmid (previously described, containing the KpnI–KpnI fragment from the Lk plasmid) was replaced with Xbal–HindIII fragment bearing the 70 bp deletion, excised from pΔ70. The 4.2-kb KpnI–KpnI fragment with the 70 bp deletion was excised from the resulting pBSKpn2 Δ70 plasmid and cloned back into the Lk plasmid, replacing the original 4.3-kb KpnI–KpnI sequence, and yielding the LkΔ70 construct.

The ΔDm targeting vector was prepared as using the following steps: a short arm of homology (neo-SAH) plasmid was constructed by using BanI (ends filled with Klenow) and NsiI to excise the 1.25-kb MAR and Eκi containing fragment from the pBκMAR plasmid. This fragment was cloned into the sticky XbaI and blunted PstI sites of the Bluescript vector. This construct was next cut at the polylinker sites ClaI and EcoRI and used for insertion of the 1.26-kb NotI–Xbal IoxP flanked neoR gene fragment from the pMMneofox-8 plasmid (all restriction ends were made blunt by reaction with the Klenow fragment), a long arm of homology (TK-LAH) plasmid was constructed by excision of the 7.1-kb PstI–PstI germline Jκ–Cκ region containing fragment from pSPgl8 plasmid (ends were bluntly filled with reaction with T4 polymerase) and ligation into the HindIII site (blunted with reaction with Klenow fragment) of pLC19/MC1-TK. The final ΔDm targeting vector was produced by cloning of the 8.9-kb Xbal–SalI fragment from TK-LAH into the neo-SAH SalI polylinker site. Targeting strategy is illustrated in Figure S1 in Supplementary Material.

CELLS AND CULTURES

All cells in this manuscript were grown in RPMI 1640 medium (Gibco) supplemented with 10% fetal calf serum, 2 mM...
t-Glutamine, 100 µg/ml penicillin, 100 µg/ml streptomycin, and 50 µM 2-mercaptoethanol. BaF3 cell medium was additionally supplemented with IL3 secreted by WEHI-3b cells. IL-7-dependent pre-B cell cultures used for chromatin immunoprecipitation (ChiP) analysis were performed as has been previously described (35). COP8 cells were transiently transfection with a Pax5 expression plasmid (gift from M. Busslinger) using the DEAE dextran method (36).

**ISOLATION AND ANALYSIS OF LYMPHOID CELLS FROM BONE MARROW AND SPLEEN**

Bone marrow cells from femur and tibia bones were flushed out with PBS using a syringe needle. Splenocytes were disrupted and pulsed dispersed in PBS. Erythrocytes were lysed with RBC lysis solution (Biological industries) and cells were washed. When indicated, cells were isolated on magnetic MACS columns (Miltenyi Biotech) according to the manufacturer’s instructions. Cell purity following isolation was assayed as <95% by flow cytometry (LSR II, BD Bioscience).

Cells from erythrocyte disrupted spleens and bone marrows were stained with the antibodies indicated and cellular composition was analyzed by flow cytometry (LSR II, BD Bioscience). The antibodies used in this report include anti-mouse-IgG-PE (Southern Biotech), anti-human-IgK-FITC (Southern Biotech), anti-IgM-APC (eBioscience), anti-IgD-PE (Southern Biotech), anti-B220-PerCP-Cy5.5 (Biolegend), anti-CD19-(Biolegend), anti-IgG-APC (eBioscience), anti-B220 (Miltenyi Biotech), according to the manufacturer’s instructions. Cell purity following isolation was assayed as <95% by flow cytometry (LSR II, BD Bioscience).

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**ANALYSIS OF DNA METHYLATION BY SOUTHERN HYBRIDIZATION**

Cellular genomic DNA (5–15 µg) was digested with the specified enzymes, electrophoresed in native (Tris–acetate) agarose gels, denatured and transferred to nitrocellulose. DNA was then hybridized with the specific radioactive probes and analyzed by autoradiography (37). Hybridization was carried out at 65°C for 16 h. The degree of methylation was measured semiquantitatively using a PhosphorImager BAS-1800 (Fuji) and Tina2.10 g software (IsotopenMedgerate GmbH).

**NUCLEAR EXTRACT PREPARATION**

Cells (3–5 x 10⁶) were washed in PBS, resuspended in low salt buffer (10 mM HEPES pH 7.9, 10 mM KCl, 0.1 mM EDTA, 0.1 mM EGTA, 1 mM DTT, 0.5 mM PMSF, 20 µg/ml aprotinin, 10 µg/ml leupeptin) and incubated for 10 min on ice. NP-40 was then added to a final concentration of 0.66%, the mixture was vortexed briefly and centrifuged for 30 s, 16,000 g. Nuclei were resuspended in high salt buffer (20 mM HEPES pH 7.9, 0.4 mM KCl, 1 mM EDTA, 1 mM EGTA, 1 mM DTT, 1 mM PMSF, 20 µg/ml aprotinin, 10 µg/ml leupeptin) and rotated for 20 min at 4°C. Nuclear debris was removed by centrifugation at 16,000 g for 20 min at 4°C.

**ELECTROPHORETIC MOBILITY SHIFT ASSAY**

Oligonucleotide probes were end-labeled with α³²P-dCTP using Klenow fragment. Two micrograms of nuclear extract was incubated with 0.3 ng of the radioactive double strand probe in a solution containing 2 µg poly-dI-dC, 10 mM Tris–HCl pH 7.9, 10% glycerol, 100 mM KCl, and 4 mM DTT for 20 min at 25°C. In competition assay, 100-fold molar excess of an unlabeled probe was preincubated for 10 min prior to the addition of the radiolabeled probe. In supershift assays, the indicated antisera (antibodies A1 and A2 kindly provided by Meinrad Busslinger) were added to the nuclear extract 15 min prior to the addition of the probe. Samples were then electrophoresed at room temperature on a 4% polyacrylamide gel (19:1 acrylamide/bis) in 0.25x TBE buffer. Gels were dried and bands were visualized by autoradiography. Probes used for assays were Dm-70 bp 5’-CGAGTTGTAATTTATATCGCCAGCAATGGAGTGAAGTGTCCGCAACCTCTTTACAACTGGGTGACGACTCCAGAGTCGACA-3’ and the Pax5-binding site from the promoter of sea urchin H2a-2.2 5’-GGGTTGAGCGCCGGTTGGTGACGACTCCAGAGTCGACA-3’.

**DNAase I FOOTPRINTING**

TaqI–SacI fragment (130 bp) from the Dm segment, encompassing the detected Pax5-binding site, was labeled with ³²P-dCTP at TaqI end by a fill-in reaction with Klenow fragment to a specific activity greater than 10⁶ cpm/ng of DNA. Probes were incubated for 20 min at room temperature with 20 µg of nuclear extract in a 50-µl reaction mixture containing 10 mM Tris pH 7.8, 14% glycerol, 57 mM KCl, 4 mM DTT, and 0.2 µg poly(dI-dC). DNAse I (0.5–1 U; Promega) diluted in 50 mM MgCl₂, 10 mM CaCl₂ was added for 1 min. The reaction was terminated by addition of 150 µl of a stop solution containing 200 mM NaCl, 20 mM EDTA, 1% SDS, and 5 µg yeast tRNA. DNA was extracted with phenol–chloroform, ethanol precipitated, dissolved in loading buffer (deionized Formamide – 5 mM EDTA), denatured for 10 min at 85°C and separated on a 6% polyacrylamide sequencing gel containing 7 M urea. Sequencing reactions performed using the Maxam and Gilbert procedure were run parallel to each probe.

**BISULFITE SEQUENCING**

DNA was converted to bisulfite treatment using the EpiTect Bisulfite kit (Qiagen) and correctly inserted clonal amplicons were sequenced by Sanger sequencing using the Maxam and Gilbert procedure. DNA was digested with the specific radioactive probes and analyzed by autoradiography (37). Hybridization was carried out at 65°C for 16 h. The degree of methylation was measured semiquantitatively using a PhosphorImager BAS-1800 (Fuji) and Tina2.10 g software (IsotopenMedgerate GmbH).

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CHROMATIN IMMUNOPRECIPITATION

IL-7-dependent pre-B cell cultures were made from the bone marrow of IgsWT/∆Dm mice as has been previously described (35). Cells were crosslinked with formaldehyde, chromatin extracted, and immunoprecipitated with an antibody directed against Pax5 (5 μg per 30 μg DNA) (SantaCruz). Semi-quantitative PCR was carried out on input DNA compared to immunoprecipitated DNA using primers specific for the Dm element and primers spanning the Dm deletion in order to test the enrichment on the WT and ∆Dm alleles separately. PCR amplicons were visualized on an 8% polyacrylamide gel. Primers used: ∆DmChIP-F 5′-CCAAGAGATTGATCGGAGA-3′, ∆DmChIP-R 5′-CCATGACTTTTGCTGGCTG-3′, WTDMChIP-F 5′-GGCCACGGTTTTGTAAGACA-3′, WTDMChIP-R 5′-CAGGGTGAA GCCAAATAG-3′, CD19-F 5′-GATTGGAAGAGTGCCTACA-3′, CD19-R 5′-GATTTGGAAGAGTGCCTACA-3′, and biotinylated Cα-actin-F 5′-GGCCGGAGCTTGCTAATGGTTTGTAACCACATGGG-3′ and Cα-actin-R 5′-CGAAGCCGGCTTGGGAA-3′. Resulting amplicons were visualized on an 8% polyacrylamide gel. Primers used:

SYMPATRIC HYPERMUTATION ANALYSIS OF PEYER’S PATCHES B CELLS

Peyer’s patches (PP) were dissected from the small intestines of 4–6-month-old IgsWT/WT, IgsWT/∆Dm, or IgsWT/∆Dm mice. PP from three to four mice were pooled for each experiment. PP were mashed through a 70 μm nylon mesh and washed with PBS to produce single cell suspensions. Cells were washed with PBS-0.5% BSA and labeled with PNA-FTIC (Vector Labs) and uB200–PE (BD Bioscience). Germinal center B220+ cells were sorted (FACSSStar BD) to greater than 90% purity. WT and ∆Dm rearranged Igs alleles were amplified with Vκ5′-GTCCCTGCGAGTTYAGTGGCAGTGGRCAC-3′ and R3-1 5′-CAGACCCCTGTCAATGGTTGTAACCACATGGG-3′ primers using high fidelity PCR kit (Roche) with an initial denaturation of 4 min at 94°C, followed by 35 cycles of denaturation at 94°C for 15 s and annealing combined with elongation at 68°C for 3 min. A supercoiled plasmid was used as a reference. Resulting amplicons were purified and visualized on a 0.8% agarose gel. Resulting amplicons were visualized on a 1.5% agarose gel, excised and purified with the QIAquick gel extraction kit (Qiagen). Allelic distribution of BALB/c/Cast transcripts was assessed by pyrosequencing on a PyroMark Q24 instrument (Qiagen) using Cx-pyro primer 5′-ACATCAACCTTCACCATCAT-3′.

LUCIFERASE REPORTER ASSAY

M12 cells were transiently transfected using the DEAE dextran method (36) with a luciferase reporter plasmid containing the minimal β-globin promoter TATA box (pTATA), without any additional regulatory elements or with insertions of the Dm element, iEx, or four NF-κB binding sites immediately upstream of the promoter. The cells were co-transfected with pβ-GAL to normalize for transfection efficiency. Luciferase activity was measured using the Luciferase Assay System (Promega) according to the manufacturer’s instructions.

RESULTS

CHARACTERIZATION OF THE Dm ELEMENT

We have previously identified an element lying ~700 bp upstream of the iEx which facilitates demethylation of the Igκ locus in cell culture, in cooperation with iEx (32). The element, designated Dm, is not part of the previously defined core iEx (Figure 1A). The Dm element, as determined by our previous experiments, spans ~1 kb and contains numerous areas which are conserved throughout different species (Figure 1A). The element itself contains a stretch of ~200 bp with the highest density of CpG sites found within the Igκ locus. In order to see whether this element was transcriptionally active, we tested its functionality in an enhancer reporter assay. We compared its activity in a reporter plasmid to the well-characterized iEx (Figure 1B). Luciferase assays show that the Dm element acts only as a weak transcriptional enhancer which is about sevenfold weaker than the core intronic enhancer in M12 B cell lymphoma cells (Figure 1C), suggesting that the Dm element on its own does not exert its effect by direct transcriptional activation.

Pax5 BINDING AT THE Dm ELEMENT

Cis regulatory elements, such as enhancers and promoters, convey their influence on cellular phenotypes by binding trans regulatory transcription factors, which mediate transcription and changes in chromatin structure. As the Igκ locus is selectively active in B cells, starting from the pre-B cell stage, we speculated that the Dm element may bind B cell-specific transcription factors, thus mediating the changes it induces. Upon searching for potential binding sites for key B cell transcription regulators, we identified an area within the CpG-rich segment with remarkable similarity to the Pax5 consensus sequence (38) (Figure 2A). A 70–bp probe containing this sequence is shifted to a specific height when incubated with nuclear extracts from B lineage cells which have passed the pro-B cell stage, but not in other cell types tested in an electro-mobility shift assay (EMSA) (Figure 2B). These results clearly show that the binding of this protein is specific for the stages when the Igκ locus is active. Notably, this specific shift can be attained using a fibroblast extract, which normally does not produce such a shift, by forced expression of Pax5 (Figure 2C), and titrated away by competition with a probe containing the Pax5-binding site of the H2a-2.2 promoter, strongly implying that indeed the Pax5 protein is binding at this site. When the nuclear extract is incubated

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FIGURE 1 | Conservation and transcriptional activity of the Dm element.
(A) Schematic map of the Igκ locus, drawn to scale. Basewise conservation across mammalian genomes is shown graphically, adapted from UCSC genome browser “Conservation” track.
(B) Schematic map of transfected plasmid constructs.
(C) Average relative luciferase activity in M12 cells transfected with the indicated plasmids. Plasmids were co-transfected with a constitutive β-Gal-expressing plasmid and luciferase activity was normalized for transfection efficiency to β-Gal activity. Transfection and luciferase assay was carried out at least three times for each construct. Error bars represent the standard deviation of the luciferase activity.

with an antibody raised against the DNA-binding domain of Pax5 (designated A1), the shift on the EMSA gel disappears, whereas incubation with an antibody recognizing the Pax5 transactivation domain (designated A2) introduces a supershift, confirming that the 70-bp probe indeed specifically binds the Pax5 transcription factor (Figure 2D). DNase I footprinting using nuclei of the Pax5-expressing M12 B cell lymphoma cell line shows a definitive protection at the putative Pax5-binding site in comparison to S194 plasmacytoma cells which do not express Pax5 (Figure 2E).

Interestingly, this specific footprint correlates precisely with the predicted Pax5-binding site. ChIP was performed on pre-B cells from IgκWT/ΔDm mice (introduction of the ΔDm allele into mice is described in Section Characterization of Methylation, Rearrangement and B Cell Development in Dm Knockout Mice) with an antibody recognizing the Pax5 protein. While the Dm positive allele showed significant enrichment for Pax5, the deleted allele was not enriched for Pax5-binding (Figure 2F). These results indicate that Pax5 indeed binds in this region in vivo and that the binding is directly dependent on the presence of the Dm element. Altogether, the above described data shows that Pax5 specifically binds to the Dm element in vivo.

Dm FACILITATES DNA DEMETHYLATION OF Igκ TRANSGENES

We wished to further investigate the role of the Dm element in demethylation of the Igκ locus. In order to do so, we introduced a well-characterized transgene (34, 39) containing a pre-rearranged Igκ allele to mice, termed Lκ (Figure 3A). Two additional transgenic mice were produced with modified constructs, one containing a deletion of the entire Dm locus, termed LκΔDm, and the second containing a deletion of the 70 bp region containing the Pax5-binding site, termed LκΔ70 (Figure 3A). DNA from splenic B220+ cells was assayed for the methylation of these transgenes by restriction analysis, which allows for simple differentiation between the transgenic and endogenous regions. Digestion with Kpnl gave rise to a 4.3-kb fragment in the Lκ and LκΔ70 transgenes and a 3.4-kb fragment in the LκΔDm transgene, whereas the endogenous locus yields a 15-kb fragment. These fragments were further digested with methylation-sensitive restriction enzymes Acil and HhaI (HhaI was not used to assess the LκΔDm state since the HhaI site is deleted in this transgene). The digested DNA was hybridized with a probe recognizing the MAR and iEκ sequences.

To assess the level of methylation, the amount of the undigested DNA was measured using a PhosphorImager. Interestingly, while the Lκ transgene was almost completely unmethylated, with only 8% of the DNA remaining undigested (Figures 3B,C), the LκΔDm transgene was highly methylated (73%) (Figures 3B,D), indicating that indeed the Dm element facilitates the hypomethylation of the Igκ locus in B cells. Notably, deletion of only 70 bp from the Dm in the LκΔ70 transgene reduced the ability of the transgene to become unmethylated (50%) (Figures 3C,D).

Bisulfite analysis of the CpG-rich region surrounding the Pax5-binding site in the endogenous locus, Lκ and LκΔ70 transgenes showed a picture that agrees quite nicely with the above results (the LκΔDm was not assayed in this manner, since this region is deleted within the transgene). These results take into consideration the difference between the methylation levels measured by bisulfite sequencing, which probes all CpG sites in the region, and the restriction analysis which measures the methylation only at

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FIGURE 2 | Pax5-binding within the Dm element. (A) Comparison of the putative Pax5-binding site within the Dm element, known Pax5-binding site in sea urchin H2a-2.2 promoter and Pax5 consensus sequence. Bases which match the consensus sequence are marked in red, while bases which do not match are marked in gray. (B) Electro-mobility shift assay (EMSA) of TaqI–BstEII 70 bp fragment from the Dm element, containing the putative Pax5-binding site, with the indicated hematopoietic cell type nuclear extracts. Red arrowhead indicates B cell lineage-specific shift. (C) Competition EMSA. Unlabeled H2a-2.2 probe competes with radioactive TaqI–BstEII Dm probe and reduces shift in B lineage cell extracts (M12, 70Z/3), and in extracts of fibroblast cells (cop8) transfected with a Pax5 expression vector. Nonsense unlabeled probe (ns) is unable to compete with Dm probe. Red arrowhead indicates specific shift. (D) Supershift assay with TaqI–BstEII Dm probe, which was incubated with antibodies (ab) raised against the DNA-binding site of Pax5 (A1), the transactivation domain (A2), or general rabbit antisera (PL). (E) DNase I footprinting assay on end-labeled TaqI–SacI probe from the Dm element. Labeled probe was incubated with nuclear extracts from the indicated cell types. A control Maxam and Gilbert reaction (G/A) was run in parallel. Footprint specific to Pax5-expressing M12 cells is indicated with a red arrow. Location of the putative Pax5-binding site is marked in red on the nucleotide sequence. (F) Chromatin immunoprecipitation (ChIP) enrichment of Pax5 at the Dm element in Igκ wt/ΔDm cultured pre-B cells. Enrichment was measured by semi-quantitative PCR via comparison of the input DNA (Input) to the immunoprecipitated fraction (αPax5), using primers specific to the Dm element (Dm) and the deleted allele (ΔDm). One and three times the amount of PCR template were run in parallel to ensure linearity. Positive (CD19 promoter) and negative (β-actin promoter) controls for Pax5-binding were analyzed in parallel to ensure specificity of the enrichment. ChIP with a non-specific antibody (IgG) does not enrich the Dm element.

the sites which correspond to the digestion site. The endogenous locus is close to 50% methylated, as expected from a region which undergoes monoallelic demethylation (Figure 3E). The LxΔ70 transgene is 76% methylated, while the Lx transgene is completely unmethylated (Figure 3E). In order to see how these results correlate with the restriction analysis, the percent of sequences which would be protected from AciI digestion was assessed. Fifty-seven percent of the LxΔ70 sequences remain protected, supporting the restriction analysis results. These experiments clearly show that the Dm element contributes to the demethylation of the Igκ locus in vivo, results that support previously published data obtained from cell culture systems.

CHARACTERIZATION OF METHYLATION, REARRANGEMENT, AND B CELL DEVELOPMENT IN Dm KNOCKOUT MICE

Given the results in transgenic mice, we generated a knockout mouse in which the entire Dm element in the endogenous locus was replaced with a LoxP-flanked Neo gene which was then excised from the genome (Figure 4A; Figure S1 in Supplementary Material). We assessed the methylation pattern of the Igκ locus by...
the Jκ and B6/Cast Rag1 marrow cells were purified from B6/Cast Rag1
Ex vivo on the strain-specific DNA polymorphisms. We did not, however, detect significant differences in
methylation levels between the WT and the Dm-deleted alleles based
on experiments. Bold AciI line indicates multiple AciI restriction sites
within close proximity. (B,C) Southern blot assay assessing methylation
state of Lk, LkΔDm (B) and LkΔ70 (C) transgenes in B220\(^{+}\) splenic B
cells. DNA was digested with KpnI either alone or with methylation
sensitive enzymes AciI or HhaI and hybridized with the indicated probe.
Methylation levels were quantified by phosphorImager. (D) Graphical
representation of the average methylation levels at AciI sites of the
indicated transgenes, as quantified by phosphorImager. Methylation was
measured in DNA from two to four independent founder mice for each
transgene. Error bars represent standard deviation of the methylation
levels. (E) Bisulfite analysis of CpGs flanking the Pax5-binding sequence
of the Dm element. Triangles mark CpGs recognized by AciI restriction
enzyme. Black circles signify methylated, white signify unmethylated,
gray signify undetermined, and red signify CpGs deleted in transgene.
Percentage of methylated CpGs within the Dm element is indicated with arrows on schematic
representation of the genomic region. Location of primers is marked by
arrows in (A).

We explored the possibility that the Dm element may affect
other developmental processes pertaining the Igκ locus and normal
B cell development, as has been observed for cis regulatory elements in the locus such as the enhancers. There was no significant
difference seen in the levels of rearrangement of the WT versus ΔDm allele, as assessed by FACS and pyrosequencing
analyses (Figure S3 in Supplementary Material). The pyrosequencing
results also indicate that the level of Igκ transcription is not
changed by the deletion of the Dm element, supporting the above
described results showing that the Dm element is a weak tran-
scriptional enhancer. The B cell development in the bone marrow of IgκΔDmΔDm mice appeared normal, with proportions of pro-
-, pre-, immature, and mature B cells similar to those of WT mice
(Figure 4D, Figure S4 in Supplementary Material). Overall, these
results indicate that, in the endogenous locus, deletion of the Dm
element does not curtail these early stages of B cell development.

**EFFECT OF Dm ELEMENT ON SHM**

We investigated whether deletion of the Dm element affects a later
stage of Igκ maturation, specifically the process of SHM in acti-
vated B cells. Levels of SHM in IgκΔDmΔDm mice versus IgκWT/WT mice were examined, and a significant drop in amount of muta-
tions in the germinal center B220\(^{+}\) PNA\(^{high}\) B cells from Peyer’s
patches of the Dm negative mice was observed (Figures 5A,B). In
order to rule out mouse to mouse variation, which could poten-
tially give rise to such an effect, SHM in heterozygous IgκWT/ΔDm

![FIGURE 3 | Effect of Dm element on methylation of Igκ locus in transgenic mice. (A) Schematic map of the transgenes and probe used
in experiments. Bold AciI line indicates multiple AciI restriction sites
within close proximity. (B,C) Southern blot assay assessing methylation
state of Lk, LkΔDm (B) and LkΔ70 (C) transgenes in B220\(^{+}\) splenic B
cells. DNA was digested with KpnI either alone or with methylation
sensitive enzymes AciI or HhaI and hybridized with the indicated probe.
Methylation levels were quantified by phosphorImager. (D) Graphical
representation of the average methylation levels at AciI sites of the
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gray signify undetermined, and red signify CpGs deleted in transgene.
Percentage of methylated CpGs within the Dm element is indicated with arrows on schematic
representation of the genomic region. Location of primers is marked by
arrows in (A).
mice was assessed. Here too, the proportion of mutations on the ΔDm allele was lower than on the WT allele (Figure 5C). As a control, a similar number of colonies were sequenced from B220⁺PNAlow cells, with no mutations detected (data not shown). While the average number of mutations is lower in the ΔDm allele, sequences which have undergone SHM do so at an efficiency similar to the WT allele, as seen when examining the mutation rate in total sequences versus rate in mutated sequences (Figure 5), suggesting that the Dm element affects the recruitment but not the processivity of the machinery involved in SHM. These results indicate that the Dm element, which is immediately adjacent to the intronic MAR and iEκ, helps promote SHM. This is particularly notable, as deletion of the MAR/iEκ region on its own has no discernible effect on the normal SHM process (18). Previous studies have shown that deletion of any single enhancer has only a small effect on the developmentally regulated DNA demethylation, whereas the combined lack of two enhancers abolishes the demethylation process (16, 19). Another difference between the transgene and the endogenous locus is that the transgene contains a pre-rearranged Igκ. It is possible that the Dm element only affects the demethylation when the locus is in a rearranged configuration, but not in the germline conformation. In this study, we see that the deletion of the Dm sequence, which is not part of the core iEκ, greatly impedes the demethylation process in the transgene, indicating that the Dm element contributes to the activity of iEκ, possibly as a co-enhancer. As the Dm is only a weak transcriptional enhancer as a solitary element, it is the combined activity with neighboring cis acting elements which gives rise to the full activity.

DISCUSSION
In this paper, we characterized a novel cis regulatory element situated within the Jκ–Cκ intron of the Igκ locus. This sequence was originally identified as an element which lies adjacent to iEκ and contributes to its demethylating activity, as deletion of either element was sufficient to abolish demethylation in a cell culture system (32). In our present study, we find that the Dm element is necessary for hypomethylation of the Igκ locus of the Lκ transgene in vivo, but is dispensable for the demethylation of the endogenous locus. The apparent discrepancy between the phenotype in these two cases may be due to the fact that the transgene contains the sequences in the Igκ locus up to the 3′Eκ, but does not include the Ed. The three Igκ enhancers work cooperatively and, to a certain extent, redundantly to activate and demethylate the locus. Previous studies have shown that deletion of any single enhancer has only a small effect on the developmentally regulated DNA demethylation, whereas the combined lack of two enhancers abolishes the demethylation process (16, 19). Another difference between the transgene and the endogenous locus is that the transgene contains a pre-rearranged Igκ. It is possible that the Dm element only affects the demethylation when the locus is in a rearranged configuration, but not in the germline conformation. In this study, we see that the deletion of the Dm sequence, which is not part of the core iEκ, greatly impedes the demethylation process in the transgene, indicating that the Dm element contributes to the activity of iEκ, possibly as a co-enhancer. As the Dm is only a weak transcriptional enhancer as a solitary element, it is the combined activity with neighboring cis acting elements which gives rise to the full activity.

The mechanisms by which genomic loci undergo targeted demethylation have long been shrouded in mystery (40). Findings from recent years have pointed to the Tet family of enzymes as
possible catalysts of the demethylation process, via oxidation of the methyl group into a hydroxymethyl moiety (41). When acting as a demethylation intermediate, the hydroxymethylated cytosine is then either passively diluted during DNA replication (42), as it is not recognized by the methylation maintenance machinery (43, 44), or, conversely, is actively excised from the genome and replaced with an unmethylated nucleotide (45, 46). Targeting of Tet proteins to specific genomic loci is sufficient to induce local demethylation (47, 48). Tet2 has been implicated in the active demethylation of tissue-specific genes in postmitotic human monocytes (49). Additionally, Tet2 has been found to bind PU.1 (50) and EBF1 (51) in the hematopoietic system. A recent report has uncovered a different strategy to induce demethylation by which DNMT1, the maintenance DNA methyltransferase, is sequestered from specific genomic loci by binding non-coding RNA. This prevents the placement of methyl groups on DNA during replication and, in turn, brings about passive demethylation of a defined region (52). It is still unclear what mechanism is implemented by the cis regulatory elements to demethylate the Igκ locus during B cell development, especially since deletion of Tet2, the strongest Tet candidate in the immune system, causes leukemia in mice (53–55), which masks many of the tissue-specific effects that may occur as a result. As methylation is a strong barrier to the rearrangement process (8), future studies can address this issue.

We have identified a sequence within the Dm element which binds the B cell lineage specifier Pax5. This site is bound by Pax5 starting with the pre-B cell stage, up to mature B cells, but is unbound in BaF3 pro-B cells, where the Igκ locus is not yet activated and made accessible for rearrangement, nor in plasma cells where Pax5 expression is down-regulated. It should be noted, though, that the BaF3 pro-B cell line tested here does not express Pax5 (56), whereas most pro-B cells do, and as such we are unable to rule out the possibility that Pax5 is already bound at the pro-B cell stage. This is the first report, to our knowledge, of a Pax5-binding site within the Igκ locus which binds Pax5 at the time of locus activation. Previous reports have located sites in 3′Eκ (24) and in the K-I–K-II (29, 30) regulatory elements in which Pax5 plays a repressive role and where binding is lost upon Igκ locus activation. The new site we report is particularly interesting, considering that Pax5 is known to be directly necessary for Igκ locus activation and k0 germline transcription in pre-B cells (31). We find that Pax5-binding in the vicinity of the 3′κ–Cκ intron is dependent on the presence of the Dm element and that the Pax5-binding site contributes to the demethylating capabilities of the Dm element. While this clearly cannot be the only Pax5-binding site, since ΔDm pre-B cells maintain their full ability to rearrange the Igκ locus, this site highlights the potency of this B cell identity protein in one more area of B cell development.

It should be noted that the sequence of the Pax5-binding site within the Dm element is conserved among rodents, but not in the human-Igκ locus, though other aspects, such as the CpG-dense region, are. This is not the only aspect which differs between the human and murine counterparts of the Igκ locus. For example, the Sis element, a transcriptional silencer which has been shown to recruit the Igκ locus to the pericentromeric heterochromatin in mice, is not conserved in the human locus (57). It stands to reason that regulation of the human and murine Igκ loci may differ somewhat, as the strongly biased usage of the κ versus λ chain seen in mice (where 95% of mature B cells express the κ chain) is not present in humans, which have a ratio of 60:40 of κ versus λ usage (58). This could be due to differences in the RR of the human and murine Igκ loci that may contribute to this phenomenon.

While the deletion of the Dm element did not, on its own, affect the methylation status of the endogenous Igκ locus, nor the relative amount of the deleted allele which underwent rearrangement, we observed a decrease in the levels of SHM on Igκ alleles lacking the Dm element. The role of the Dm element in facilitating SHM appears to be independent of the iEκ/MAR region, since the combined deletion of the iEκ and MAR elements has no perceptible effect on SHM (18). The lower level of SHM does not appear to be the result of lower levels of Igκ transcription, since deletion of the Dm element does not lower the levels of Igκ RNA
observed in mature B cells. Deletion of the Dm element appears to cause inefficient recruitment of the mutating machinery, but once the machinery is in place, the mutation efficiency is similar to the WT locus. The element may therefore function by efficiently recruiting the mutation machinery to the locus, possibly by key regulators such as Pax5 which are bound to the Dm element. Pax5 itself has a known role in SHM by activating the transcription of the Aicda gene, encoding the AID protein, which is the deamidase responsible for SHM (59, 60). It may be that Pax5 plays more than one role in SHM induction. The role of the Dm element in SHM fits in well with its location, which is almost immediately adjacent to the Vκ–Jκ rearranged region which is the hotspot for SHM.

In conclusion, we have characterized the Dm sequence as an element that regulates the lgκ locus during different stages of B cell development. The Dm is both a team player, cooperating with the three characterized enhancers to demethylate the locus for rearrangement, as well as an element that affects the locus in its own right in allowing efficient SHM. This report adds to our understanding of the complex regulation of the lgκ locus, which undergoes many drastic changes during development and must be fine-tuned for each developmental stage.

AUTHOR CONTRIBUTIONS
Rena Levin-Klein, Andrei Kirillov, and Chaggai Rosenbluh designed the experiments, did the research, and interpreted the results. Howard Cedar and Yehudit Bergman directed the study. Rena Levin-Klein and Yehudit Bergman wrote the manuscript.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at http://www.frontiersin.org/journal/10.3389/fimmu.2014.00240/abstract

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