Characterization of a Cis-Acting Regulatory Element which Silences Expression of the Class II-A β Gene in Epithelium

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Summary

Class II major histocompatibility complex (MHC) genes encode for α/β chain pairs that are constitutively expressed principally on mature B cells and dendritic cells in mice. These gene products are easily induced on macrophages with cytokines, and may also aberrantly appear on the surface of epithelium during immune injury. The appearance of class II determinants in parenchymal tissue potentially renders these somatic cells capable of antigen presentation to circulating CD4+ T lymphocytes, and their absence may be protective for normal tissues expressing self-antigens. The low surface class II expression observed on parenchymal cells generally correlates with low levels of mRNA, suggesting that transcription rate is a major element in class II regulation. To understand the transcriptional mechanism maintaining low basal surface expression of class II in somatic cells, we transiently transfected mini-gene reporter constructs to study the regulation of the murine Aβ promoter in a cultured renal epithelial cell line. We describe here a negative cis-acting regulatory region located between −552 and −489 bp upstream of the Aβ cap site that silences the transcriptional activity of the Aβ promoter in epithelial cells in an orientation-dependent manner, and is also able to silence a heterologous promoter. This region is not active in class II-expressing B cells (BAL-17) in culture, but is functional in two other murine class II-negative cell lines, fibroblasts and thymoma T cells. Using competition electrophoretic mobility shift assays, we have localized the core protein binding site within this region to an 8-10-bp response element, designated AβNRE, at −543 to −534 bp. A nuclear extract from BAL-17 cells does not bind to this element. Mutation of this site abrogates the transcriptional silencing activity of the region. We conclude that the transcription of class II-Aβ in parenchymal cells, and some lymphocytes, can be actively repressed by an upstream silencing element.

The gene products of the class II region of the MHC are required for the recognition of antigen by CD4+ T lymphocytes (1). Class II molecules are normally expressed constitutively in mature B lymphocytes (2) and dendritic cells (3), and are easily induced in antigen-presenting macrophages with IFN-γ or TNF-α (4). Other somatic cells, such as organ epithelium, normally express little or no class II on their cell surface, but can increase their expression during episodes of local inflammation. This upregulation has been noted in such settings as type I diabetes mellitus (5), GVHD (6), and various forms of tubulointerstitial nephropathy (7, 8). Again, IFN-γ has been implicated as the primary mediator (for reviews see references 9 and 10).

Under circumstances of increased class II expression, some nonlymphoid cells (11-13) including renal tubular cells (14-17), are capable of presenting both self and foreign antigen to CD4+ T lymphocytes or hybridomas. The surface expression of class II as a gene product correlates well with levels of mRNA transcript, and a variety of studies suggest that levels of mRNA encoding AαAβ/EαEβ are regulated by transcriptional controls rather than by modulation of stability of transcript levels (9). Transcript levels are generally abundant in mature B cells, and very low in somatic parenchymal cells. Thus, an insight into how class II determinants are normally repressed in somatic cells, particularly at the transcriptional level, may be important in understanding the antigen-presenting function of parenchymal cells in contributing to the maintenance of a local immune response against self (8, 18, 19).

The cis-acting response elements and associated trans-acting
nuclear proteins involved in the positive enhancement of class II transcription in murine and human lymphoid cells have been well characterized, with most of the key elements clustering in a proximal promoter region in the first 200 bp upstream of the transcription start site (9). Two lines of evidence suggest that the absence of class II expression on other cell types is at least partly a function of active silencing by negative-acting nuclear factors, as opposed to simply a lack of transcriptional enhancers. First, fusion of class II-positive B cells with either class II-negative plasma cells (20) or fibrosarcoma cells (21) extinguishes both basal and inducible mRNA expression of class II genes in the hybrid. Second, many of the trans-acting proteins that bind to the conserved X and Y box promoter elements upstream of all class II genes can be detected in both class II-positive and negative cell lines (22, 23), although it has been suggested that these do not occupy their binding sites in class II-negative cells in vivo (22). Here we describe a novel cis-acting element upstream of the murine αβ gene which acts to silence transcription of that gene in low class II-expressing renal tubular epithelium, fibroblasts, and T cells, but not in class II-positive B cells.

Materials and Methods

Cell Lines. MCT epithelium is a low class II-expressing (AαAβ\textsuperscript{+}/EαEβ\textsuperscript{+}) murine proximal renal tubular cell line (14) that has been maintained in culture in DMEM supplemented with 10% FCS, 100 U/ml penicillin, and 100 µg/ml streptomycin. BAL-17 is an H-2\textsuperscript{b} B lymphoma cell line that expresses high levels of class II (kindly provided by Dr. John Monroe, University of Pennsylvania; 24), and was maintained in RPMI 1640 medium with 10% FCS, penicillin, and streptomycin, and 5 × 10\textsuperscript{-5} M 2-ME. NIH-3T3 is a class II-negative murine fibroblast cell line maintained in media described for MCT cells, and 2B4 is a class II-negative murine T cell thymoma line (kindly provided by Dr. Helen Quill, University of Pennsylvania) maintained in media described for the BAL-17 cell line.

Plasmids. The plasmids 8.54, 8.64, 12.19, and B9 contain, respectively, −322, −489, −669, and −2,600 bp of genomic DNA upstream of the Aβ cap site (taken from a combination of H-2 haplotypes b, d, and k) placed 5’ of the chloramphenicol acetyltransferase (CAT) gene (25). AoCAT contains the CAT gene with no promoter elements, and SV\textsubscript{40}CAT contains the CAT gene driven by an SV\textsubscript{40} enhancer and promoter. SprCAT contains the SV\textsubscript{40} enhancerless promoter. The plasmids Δ669/566.8 and Δ669/489.8 were constructed by cloning a 103-bp fragment (−669 to −566) and a 179-bp fragment (−669 to −489), respectively, from 12.19, into the unique Sall site at the 5’ end of 8.54 in the forward (F) and reverse (R) orientations. The plasmid Δ542/492.8 was constructed by PCR amplification of an 88-bp fragment from −605 to −517 of plasmid 12.19 using the flanking oligomers 5’-gtcaggtgttgttgttggagtattgCAGG-3’ and 5’-gtcagctctactggtttttgttgataagc-3’ (lower case letters represent added Sall restriction site). PCR amplification reactions were performed using the GeneAmp kit\textsuperscript{(R)} (Perkin Elmer Cetus, Norwalk, CT) with 40 pM of primer, 200 µM each of dNTP, amplification buffer (10 mM Tris-HCl, pH 8.3, 50 mM KCl, 1.5 mM MgCl\textsubscript{2}, and 0.001% gelatin) and 1.25 units Taq polymerase (27). 40 cycles of amplification were completed by annealing at 55°C for 90 s, performing the extension step for 120 s at 72°C, and the denaturation step for 60 s at 92°C. The amplification products were subcloned into the pCR\textsubscript{II} vector (Invitrogen, La Jolla, CA), and then transferred into the Sall site of 8.54, again in both orientations. This same fragment was also cloned into the Sall site upstream of the promoter of SprCAT to create Δ552/489.Spr. Plasmid Δ552/489M.8 was constructed in a similar manner to Δ552/489.8 except that the 5’ primer was replaced with the mutated primer gtcagAGCTCAATGACCTTCATAGGAGGCCCA (mutation underlined). The insertional orientation and sequence of all constructed plasmids were verified by dyeoxy chain termination (26, 27).

Detection of Aβ mRNA by Northern Hybridization. Cultured cells were washed in RNA-free PBS, and poly(A)\textsuperscript{+} mRNA was extracted using the FastTrack oligo(dT)-cellulose kit (Invitrogen). 5 µg poly(A)\textsuperscript{+} mRNA was electrophoresed through a 1.2% agarose gel with 2.2 M formaldehyde. The RNA was blotted by capillary action onto a GeneScreen membrane (New England Nuclear, Boston, MA), prehybridized, and hybridized with a 0.79-kb fragment from cDNA clone p7l3a encoding murine Aβ (26). Selected filters were stripped and rehybridized with a cDNA probe encoding the housekeeping gene glyceraldehyde-3-phosphate dehydrogenase (GAPDH) to account for small loading and transfer variations.

Transfections and CAT Assay. Transient transfections of MCT, 2B4, and 3T3 cells were carried out using calcium phosphate precipitation (28); replicate experiments were performed as indicated in the figure legends. 5 × 10\textsuperscript{5} MCT cells were plated in 60-mm dishes and transfected 24 h later with 10 µg of plasmid DNA (or equimolar equivalents), incubated in 5% CO\textsubscript{2} at 37°C for 24 h, after which time the precipitate was removed and the cells were incubated for a further 24 h in complete medium. Cells were lysed by repeated freeze–thaw cycles. Transfections of BAL-17 cells were carried out using DEAE–dextran (29). 10\textsuperscript{7} BAL-17 cells were transfected with 20 µg plasmid DNA in 500 µg/ml DEAE-dextran for 30 min followed by a 3-min shock in 10% DMSO. Cells were harvested 64 h after transfection. CAT assays were performed with all cell lysates heat inactivated at 65°C for 10 min, and incubated for 2 h in the case of MCT cells, or 16 h in the case of BAL-17 cells (28). Loading of cell lysates for incubation was normalized for protein content.

Nuclear Extract Preparation and Gel Shift Assay. MCT nuclear extracts were prepared according to the method of Dignam et al. (30) with the following modifications: three additional protease inhibitors, leupeptin, aprotinin, and pepstatin A (Boehringer Mannheim Biochemicals, Indianapolis, IN) were added to all buffers. Nuclei were extracted in 20 mM Hapes, 15% glycerol, 1.5 mM MgCl\textsubscript{2}, 5 mM dithiothreitol (DTT), 0.4 M KCl, 1 mM EDTA, and 0.5% NP-40. Buffer D contained 15% glycerol and 0.5 mM ZnSO\textsubscript{4}, but was otherwise unchanged. BAL-17 nuclei were extracted in the presence of 0.43 M NaCl rather than KCl, and in the absence of NP-40. The probe S89A was created by amplification of an 88-bp fragment from −605 to −517 of plasmid 12.19 using the flanking oligomers 5’-gtcaggtgttgttgttggagtattgCAGG-3’ and 5’-gtcagctctactggtttttgttgataagc-3’ (lower case letters represent added Sall restriction site). PCR amplification reactions were performed using the GeneAmp kit\textsuperscript{(R)} (Perkin Elmer Cetus, Norwalk, CT) with 40 pM of primer, 200 µM each of dNTP, amplification buffer (10 mM Tris-HCl, pH 8.3, 50 mM KCl, 1.5 mM MgCl\textsubscript{2}, and 0.001% gelatin) and 1.25 units Taq polymerase (27). 40 cycles of amplification were completed by annealing at 55°C for 90 s, performing the extension step for 120 s at 72°C, and the denaturation step for 60 s at 92°C. Amplification products were subcloned into the pCR\textsubscript{II} vector (Invitrogen, La Jolla, CA), and then transferred into the Sall site of 8.54, again in both orientations. This same fragment was also cloned into the Sall site upstream of the promoter of SprCAT to create Δ552/489.Spr. Plasmid Δ552/489M.8 was constructed in a similar manner to Δ552/489.8 except that the 5’ primer was replaced with the mutated primer gtcagAGCTCAATGACCTTCATAGGAGGCCCA (mutation underlined). The insertional orientation and sequence of all constructed plasmids were verified by dyeoxy chain termination (26, 27).
Results

MCT Epithelial Cells Express Low but Detectable Levels of Aβ mRNA. We sought to determine the transcriptional mechanism whereby parenchymal epithelium maintains immunologic neutrality by means of low MHC class II expression, and made use of a renal tubular cell line to do so. MCT epithelial cells normally express little to no Ac~/A~ on their cell surface, but this expression can be upregulated somewhat with IFN-γ (8). Unlike mature B cells which constitutively express class II determinants, MCT cells would be predicted to have very low levels of mRNA encoding class II. Fig. 1 presents a 10-d exposure of a Northern blot of the Aβ mRNA message level in MCT cells compared with the class II–expressing BAL-17 B cell line. Basal class II mRNA message levels in MCT cells are clearly much lower than in B cells. We have also previously isolated full-length cDNA clones encoding Aβ from an MCT library, sequence analysis of which indicated that there are no mutations in the coding region of Aβ which would explain its low surface abundance by aborted translation (26).

A DNA Segment Upstream of the Aβ Gene Silences the Homologous and a Heterologous Promoter in MCT Cells. In Fig. 2, a representative transfection of MCT cells with Aβ gene deletion constructs demonstrates strong CAT activity with the shorter promoter segments which drops off in activity between −669 and −489 bp, indicating the possible presence of a negative cis-acting element in this region. The position independence of this region was established by creating plasmid Δ669/489.8, in which region −669 to −489 was ligated directly to the homologous proximal promoter for Aβ contained in a 322-bp segment upstream of the gene. As can be observed in Fig. 3 A, this region was still able to silence the strong positive CAT activity of the downstream region in MCT cells, but only when placed in the forward orientation (F); similar results were observed with 3T3 fibroblasts (Fig. 3 B) and 2B4 thymoma T cells (Fig. 3 C). Transfection of BAL-17 cells with these constructs (Fig. 4), however, revealed no silencing of proximal CAT activity by Δ669/489.8, suggesting the absence of a relevant trans-acting factor in these class II–expressing B cells.

To functionally map the silencing region more closely, further constructs were created in which portions of this region were linked to the proximal homologous promoter (31), as diagrammed in Fig. 5. Transfection of these constructs into MCT cells further localized the silencing element to a region between −552 and −489. Again, the orientation dependence of this silencing effect is evident. This same region was also able to silence the heterologous SV40 enhancerless promoter, as shown in Fig. 6. This region in Aβ was of identical sequence to that found in the genomic DNA used to assemble the original Aβ CAT constructs used above (25), with the exception of a single base pair substitution at position −547.

Mapping the Core Binding Site of the Putative Silencing Factor. Gel shift analysis was performed using an 88-bp probe from −605 to −517, and revealed one major shifted band of interest (Fig. 7, lane 1). Competition analysis using 100-fold molar excess of the unlabeled double-stranded oligomers diagrammed in Fig. 7 revealed that the S12 oligonucleotide (lane 3), but not S11 or S14 (lanes 2 and 4), was able to com-

Figure 1. Northern blot for Aβ transcripts in BAL-17 B cells and MCT renal epithelium. 5 μg poly(A)+ RNA extracted from BAL-17 cells (lane 1) or MCT cells (lane 2) were fractionated in a 1.2% agarose gel, blotted, and probed with a 790-bp Aβ cDNA probe and exposed for 10 d. The blot was stripped and reprobed with a cDNA probe for the housekeeping gene GAPDH.

Figure 2. Transient transfection of deletion constructs into MCT cells. (Right) Representative CAT experiment derived from transfection of the diagrammed constructs. All numbering of constructs represents the number of base pairs upstream of the transcription start site for the Aβ gene (+1). Radioactive spots were cut out of the TLC plate and counted, and expressed as percent acetylation relative to SV2CAT, which is arbitrarily set at 100%. Values represent the average of three independent transfections.
Figure 3. Effect of –669 to –489 region on homologous promoter in MCT cells (A), 3T3 fibroblasts (B), and 2B4 thymoma T cells (C). The –669 to –489 region was linked directly to the proximal promoter region (contained in a 322-bp segment upstream of the cap site) in the forward (F) and reverse (R) orientation. Percent acetylation (average of three independent transfections) is expressed relative to that of 8.54, which was set at 100%.

Figure 4. Effect of –669 to –489 region on homologous promoter in BAL-17 cells. Results expressed as described in legend to Fig. 3.

pete with the labeled probe, narrowing the binding site of the relevant nuclear protein to the region between –556 and –532 bp. By competing with mutants of the S12 oligonucleotide (lanes 5–9 [32]), it was observed that only M.4 and M.5 (Fig. 7, lanes 7 and 8) fail to abolish the shifted band, indicating that the base pairs mutated in these two oligomers are essential for the binding of the putative silencer protein. The core binding site was thus localized to an 8–10-bp segment, (CC)ATTATTAA, from –543 to –534 bp, which was designated AβNRE. The other faint bands observed on gel shift were either not consistently present or were not abolished by competition with cold full-length probe (data not shown), and were thus not felt to represent specific protein binding. As demonstrated in Fig. 8, this binding complex was absent in an extract derived from BAL-17 cells.

To establish the functional importance of this binding site, a mutation ([CC]ATTAGGCC) was introduced into a CAT construct in the last four nucleotides of AβNRE. Thus, plasmid Δ552/489M.8 links the –556 to –489 region, containing a mutation of bp –537 to –534, in a forward orientation to the homologous promoter region. As shown in Fig. 9, mutation of the binding site largely abrogates the silencing activity of this region.

Discussion

The immunologic implications of aberrant expression of class II MHC gene products on nonlymphoid cells, as seen

Figure 5. Localization of silencer region to –552 to –489 bp. Diagrammed constructs were transfected into MCT cells with results expressed as described in legend to Fig. 3.
in a wide array of settings and likely mediated primarily by IFN-\(\gamma\), have been the subject of considerable interest in recent years. Although several groups have been able to demonstrate a T cell proliferative response to intact (15, 33) or processed (11) antigen presented by class II-expressing parenchymal cells, implying that such cells contribute, under certain pathologic conditions, to the local immune response, others have suggested that such "nontraditional" antigen presentation may in fact lead to T cell anergy, and thus contribute to peripheral self-tolerance (34). In either event, studying the normal mechanisms maintaining low epithelial class II expression (and thus immunologic silence) will contribute to an understanding of issues pertaining to peripheral tolerance and organ-specific autoimmunity.

It has become increasingly appreciated that transcription of eukaryotic genes may be modulated by negative-acting elements in addition to the transcriptional enhancers that have long been recognized. Since the first description of silencing elements in yeast (35), numerous studies have demonstrated factors that exert a negative effect on the transcription of a wide range of genes (31, 32, 36, 37) including MHC class II (38). That such a silencing factor might be operative in suppressing the transcription of class II genes in nonexpressing cells is suggested by fusion studies that have demonstrated extinction of class II expression in hybrids of class II-negative and -positive cells (20, 21). In one of these (21), multiple clones of four independent fusions of L929 fibrosarcoma cells with splenic B cells were studied for both basal and inducible class II message and gene product. Although the hybrids retained expression of class I genes derived from both fusion partners and retained the E\(\beta\) gene at the genomic level, none of the clones had any detectable class II expression. Similar findings were noted using transient fusion techniques, so that chromosomal loss appeared not to have been the mechanism of class II extinction. A more recent study found reactivation of low levels of class II transcript in plasmacytoma cells and human T cells on transient fusion with B lymphoblastoid cells and to a much lesser extent with splenocytes (39). Whereas these opposing results are difficult to reconcile, the study by Stuart et al. (21) is highly suggestive of the presence of a negative trans-acting factor operating in at least some cell types. In addition, two studies have previously demonstrated functional evidence of a class II silencer element in the region we report. Boss and Strominger (40), using stable transfection of DQ\(\beta\) constructs into fibroblasts, found a three-fold increase in transcription on deletion of upstream DNA from \(-2,500\) to \(-590\) bp, and a further 3.3-fold increase on deletion to \(-159\) bp. Thanos et al. (41), using transient transfection of E\(\alpha\) constructs into a xeroderma cell line, found an increase in basal expression on deletion from \(-873\) to \(-353\) bp.
of A3NRE. (*) Site of mutation. Results are expressed as described in legend to Fig. 3.

Figure 8. Gel shift of BAL-17 nuclear extract. S89A probe was incubated with MCT (lane 1) or BAL-17 (lane 2) nuclear extract. (Arrow) Position of major shifted band.

Figure 9. Effect of mutation of AβNRE. Δ552/489M.8 links silencing region -552 to -489 to proximal promoter region with a mutation introduced in bp -537 to -534 corresponding to the last four base pairs of AβNRE. (†) Site of mutation. Results are expressed as described in legend to Fig. 3.

bp. To date, the region in the range of -1,200 to -400 bp upstream of the transcription start site for class II genes has not been examined in further detail.

We have mapped a novel silencing region to a segment between -552 and -489 bp upstream of the murine Aβ cap site. This region, which acts in a position-independent, but orientation-dependent manner, is active in a low class II-expressing renal tubular epithelial cell line (MCT), but not in a class II-positive B lymphoma cell line (BAL-17). Gel shift competition analysis using a series of mutated oligonucleotides localized the core protein binding site, termed AβNRE, to an 8-10-bp AT-rich element at position -543. Mutation of this binding site abolished the ability of the region to silence the transcriptional activity of the downstream positive-acting elements of the Aβ gene when linked directly upstream of them. Our studies of this region in two other murine class II-negative cell lines (one a fibroblast and the other a T cell) suggest that this upstream silencer may represent a relatively ubiquitous mechanism of class II transcriptional repression.

A search of the GenBank™ database revealed no homology between AβNRE and other known transcriptional elements. However, a novel member of the bZIP family of DNA-binding proteins, E4BP4, has recently been isolated from a human placental cDNA library (42), and its optimum binding site, (G/A)TTATGTAA(C/T), bears a marked degree of homology of AβNRE, (CC)ATTATTAA(C). Cotransfection of an E4BP4 expression plasmid with a reporter bearing its binding site upstream of a heterologous promoter revealed it to have transcriptional repressing activity. The function and distribution of this binding protein is currently unknown. Although AβNRE is not found in the first 1,296 bp 5' of the translation start site in the Aα gene, the first 8 bp of the element, CCCATTATT, are found 480 bp upstream of the transcriptional start site of Aβ's analogous human gene, DQβ. In fact, an examination of the proximal promoter region reveals greater degree of identity of conserved elements between Aβ and DQβ than between Aβ and Aα, with at least one element (a nuclear factor xB binding site) which is present in both Aα and DQα but not in the corresponding β chain promoters (10). Whether Aα is coordinately silenced by a poorly conserved version of AβNRE or by an entirely different mechanism remains to be determined.

Transcriptional repression of eukaryotic promoters is mediated by DNA-binding proteins which act either by competing for DNA-binding sites, by quenching or sequestering of activator proteins, or by direct repression of the basal transcription complex (43, 44). The mechanism of action of the putative silencing protein that binds to AβNRE can only be speculated upon, but any model must take into account two key findings. First, Liou et al. (45) have found that class II-expressing cells possess three DNase I-hypersensitive sites in their Aα genes, at -1,400 bp, in the first intron, and in the proximal promoter region. The latter of these, D2, was absent in class II-negative cell lines, and on fusion of T and B cells (in which class II expression was extinguished), the authors demonstrated the absence of the D2 site derived from the Aα gene of either fusion partner in the hybrid. Second, it has been shown that the class II promoter elements are not occupied in vivo in class II-negative cell lines, although their binding factors can be detected in vitro gel shift assays (22). Thus, the AβNRE binding protein might be expected to act in a manner analogous to the α2 repressor of the yeast S. cerevisiae, which is able to place a nucleosome near a promoter site (46). This would have the effect of rendering the promoter both inaccessible to activating factors and insensitive to DNase I. However, silencers that function in this manner tend to demonstrate both position and orientation independence (43), and so the marked orientation dependence of AβNRE is somewhat puzzling. A lesser degree of orientation dependence was noted when a 274-bp segment from the first intron of the α1 collagen gene was placed upstream of the promoter of that gene (36), so our finding is not without precedent. IFN-γ (or factors induced by it), which exerts its class II upregulating effect through the proximal promoter region (47), could disrupt this nucleosome structure, as has been shown in the case of the glucocorticoid–glucocorticoid receptor complex and the murine mammary tumor
virus promoter (48). Alternatively, both constitutive class II-expressing and IFN-γ induced cells might produce a factor that is able to sequester the silencing protein and prevent its effects on promoter accessibility. A fuller understanding of the mechanism of action of the nuclear factor recognizing AßNRE must await isolation of this protein.

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