INTRAGENIC RECOMBINATION IN AN Eβ GENE
FOR A MURINE Ia ANTIGEN*

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The plasma membrane-located murine Ia antigens consist of two polymorphic
chains: the larger chain, designated α, is ~35,000 mol wt, whereas the smaller chain,
designated β, is ~28,000 mol wt (1). Whereas the I-A antigens are encoded by genes
(Aa and Ab, located within the I-A subregion, the I-E antigens are encoded by one
gene (Eα) located in the I-E subregion and a second (Eβ or Aβ) in the I-A subregion
(2).

Previous studies from our laboratories have used F1 hybrids to explore the serological
(3, 4) and biochemical (5) basis for the complementation between the Eα and Eβ
genes. These studies, which established that complementation could occur between
the genes involved in the trans configuration, provided evidence that the structural
genes for the α and β chains were the complementing genetic elements. During the
course of the biochemical studies (5), we compared the Eβ chain derived from a
(D2.GD × A.TFR5)F1 (Aa, Eb × Aa, Eb)2 to the Eβ chain from B10.D2. Although
the Eβ chain from the F1 animals (presumably encoded by the D2.GD parental
chromosome in the F1) was quite similar to the Eβ chain from B10.D2, they were not
identical. This result was not expected because D2.GD (H-2k) is a recombinant
strain previously typed as Kd Ad[Bl Ab Eβ Cβ Sβ Dβ, and therefore it should possess
the same Eβ gene as B10.D2 (H-2k). This difference between the Eβ chain encoded by
the D2.GD genome and other bona fide Eβ chains was also noted by Jones (6), who
used two-dimensional (2D) electrophoresis to document the difference.

Two obvious and mutually exclusive hypotheses may be put forward to explain the
aberrant nature of the Eβ chain from D2.GD. The first of these is that the recombi-
national event in D2.GD occurred not between the I-A and the I-B subregions, as

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Jones and her co-workers originally used the designation Aa to denote the I-A subregion location of this
gene. We will use the Eα notation because the gene product, the Eα chain is part of the I-E antigen.
§ Strain D2.GD, like all I-Eb type strains, does not contain a functional Eα gene, whereas the I-A' region
of A.TFR3 does not contain a functional Eα gene. Thus, (D2.GD × A.TFR3)F1 animals express only a
single I-E antigen produced by trans complementation of the Eα and Eβ genes.

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previously indicated, but rather within the \( E_{\beta} \) gene within the \( I-A \) subregion. Alternatively, the previous assignment of the recombinational event in D2.GD was correct, and mutations have occurred within the \( E_{\beta} \) gene of D2.GD, allowing it to drift slightly from a true \( E_{\beta} \) gene. Data are presented in this paper that support the first of these two alternatives.

Materials and Methods

**Mice.** (D2.GD × A.TFR5)\( F_1 \) mice were from the colony (CSD) at the Mayo Clinic and Medical School. Mice of the B10.D2 and B10.A(SR) strains were obtained from The Jackson Laboratory (Bar Harbor, ME).

**Isolation of I-E Antigens.** The \( \alpha \) and \( \beta \) chains of I-E antigens were isolated as previously described (5). Briefly, leukocytes from normal spleens were metabolically radiolabeled with \(^3\)H-labeled or \(^14\)C-labeled amino acids by incubation for 21 h, and, after labeling, Nonidet P-40 (NP-40) lysates of the cells were prepared. The I-E antigens were isolated by lentil lectin chromatography and indirect immunoprecipitation, using an (A.BY × B10.A(18R))\( F_1 \) anti-B10.A(3R) alloantisemirum and *Staphylococcus aureus* Cowan I strain. Alternatively, the immunoglobulin-depleted NP-40 lysate was applied directly to an affinity chromatography column, prepared by coupling purified anti-Ia.7 monoclonal antibody (14-4-4S, produced by hybridoma cells generously provided by Dr. David H. Sachs, National Institutes of Health, Bethesda, MD) to Sepharose, followed by elution of the bound I-E antigen from the column with glycine-HCl (pH 2.1). The \( E_{\alpha} \) and \( E_{\beta} \) chains from the isolated I-E antigens were separated without reduction by preparative polyacrylamide gel electrophoresis in sodium dodecylsulfate-containing buffers (SDS-PAGE).

**Tryptic Digestion and Analysis of Peptides.** Purified \( E_{\alpha} \) and \( E_{\beta} \) chains obtained from SDS-PAGE were reduced and alkylated and digested with trypsin, as previously described (5). Analysis of tryptic peptides was by medium pressure ion exchange chromatography on a DC-6A cation exchange resin (Dionex Corp., Sunnyvale, CA) eluted with a pH 3.13 to pH 5.05 pyridine/acetic acid gradient (5).

**Amino-Terminal Amino Acid Sequence Analysis.** The \( E_{\alpha} \) and \( E_{\beta} \) chains to be used for amino acid sequence studies were extracted from the appropriate SDS-PAGE slices with 0.01% SDS and then lyophilized. The lyophilized material was redissolved in 1 ml water, dialyzed twice against 1 liter of 0.001% SDS, and re-lyophilized. The sample was dissolved in 2 N formic acid and applied to a Beckman 890C sequencer (Beckman Instruments Inc., Fullerton, CA) for amino acid sequence analysis (7).

**Results**

**Tryptic Peptide Comparison of the \( E_{\beta} \) Chain with the \( E_{\beta} \) Chain from (D2.GD × A.TFR5)\( F_1 \) Hybrids.** Previous studies of the I-E antigen obtained from (D2.GD × A.TFR5)\( F_1 \) hybrids have shown that the \( E_{\alpha} \) chain contributed by the A.TFR5 parent is identical with the \( E_{\alpha} \) chain from B10.BR(5).

However, the \( E_{\beta} \) chain contributed by the D2.GD parent differed somewhat from the \( E_{\beta} \) chain from B10.D2(5). Fig. 1 shows the comparison of \(^{14}\)C]arginine and lysine-labeled tryptic peptides from B10.D2 with the \(^3\)H]arginine and lysine-labeled tryptic peptides derived from the \( E_{\beta} \) chain from the D2.GD parent in the \( F_1 \) hybrid. The use of arginine and lysine labels allows visualization of all tryptic peptides in the molecules, except the carboxy-terminal peptide. Although the two profiles are quite similar, sharing fourteen peptides between them, there are three peptides unique to the \( E_{\beta} \) chain (marked by the open arrows) and one unique to the \( E_{\beta} \) chain (marked by the filled arrow).

**Tryptic Peptide Comparison of the \( E_{\beta} \) Chain with the \( E_{\beta} \) Chain from (D2.GD × A.TFR5)\( F_1 \)**

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Hybrids. We next compared the tryptic peptide profile from the $E_a$ chain with that from the $E_b$ chain from B10.A(5R). The rationale for this experiment was as follows: if the $E_a$ chain from the $g^2$ haplotype arises from an $H-2^g/H-2^b$ intragenic recombinational event, then those tryptic peptides that are unique to $E_a^g$ as compared with $E_b^g$ should correspond to some of the peptides produced by tryptic digestion of the $E_b$ chain. Fig. 2 shows the comparison of the arginine and lysine-labeled tryptic peptides from the $E_a^g$ and $E_b^g$ chains. As may be seen from the figure, the $E_a$ chains from these two haplotypes share a number of peptides (peptide homology of ~55%) but, significantly, those three peptides by which $E_a^g$ differs from $E_b^g$ are shared between $E_a^g$ and $E_b^g$. These peptides are indicated by arrows in Fig. 2.

Sequence Analysis of the Amino-Terminal Portion of $E_b$ Chains. Radiochemical sequence analysis was carried out on $E_b$ chains metabolically radiolabeled with $[\text{3H}]$phenylalanine. The results of the sequencer runs are presented in Fig. 3 and are summarized in Table I. These data demonstrate that the amino-terminal $[\text{3H}]$phenylalanine profiles are identical for the $E_b^g$ and $E_b^b$ chains, and these differ from the profile for $E_b^d$. This suggests that the amino-terminal portion of the $E_b^g$ chain is derived from the $d$ haplotypes as opposed to the $b$ haplotype. The $E_a^g$ sequence (Fig. 3, panel D) from B10.A(5R) is included because the $E_a$ chains that were sequenced contained $E_b^d$ chains at the level of ~10% contamination. (In panels A–C, compare the size of the peaks at step 8 with the heights of the peaks at step 12.) All assigned phenylalanine positions are from peaks that fit a single repetitive yield line (~92%). These data support the postulate that each sample sequenced was a single species that, except for the limited...
contamination by the Eα chain, was homogeneous Eβ chain. The phenylalanine residues identified within positions 1-32 agree with data previously published for Eα, Eβ, and Eγ chains (8-11), whereas those identified at positions 41 and 48 (see Table I) have not been reported previously.

**Discussion**

Because the D2.GD (12) and B10.D2 (13) strains were each derived from crosses involving DBA/2 (H-2^d^), any genetic elements with the d haplotype in these strains should be identical because they were derived from the same donor strain. For the A^d^ and A^b^ chains this appears to be true, as Jones (6) was able to demonstrate identical two dimensional gel patterns for these chains from all three strains. The analysis of the Eβ chains is complicated somewhat by the fact that D2.GD will not express normal amounts of the Eβ^d^ chain, and an F1 hybrid between D2.GD and A.TFR5 (A"^d"E^b^ × A^d^E^b^) must be used to obtain usable amounts of the Eβ^d^ chain. Again, Jones' two dimensional gel analysis (6) suggested that there was a difference in electrophoretic mobility among the Eβ chains from the three strains and that the Eβ chain derived from the D2.GD genome in the F1 hybrid was clearly different from the Eβ^d^ chains from B10.D2 and DBA/2. Furthermore, previous observations from one of our laboratories (14) have identified a serological difference between these Eβ chains: bona fide Eβ^d^ chains (such as that from B10.D2) express a specificity, Ia.50, absent from the Eβ^d^ chain.

In our previous analysis (5) of the Eα and Eβ chains from (D2.GD × A.TFR5)F1 animals, we made two observations: (a) the Eα^b^ chain, putatively contributed by the
A TFR5 parental chromosome, was identical to authentic Eα chains; and (b) the Eβ chain, putatively contributed by the D2.GD parental chromosome, differed slightly from authentic Eβ chains. As the results presented in Figs. 1 and 2 amply demonstrate, the three peptides unique to the Eβ chain, when compared with the Eα chain, do co-elute with tryptic peptides from the Eβ chain.

There are two likely explanations for the "aberrant" Eβ chain in D2.GD. The first is that a series of mutations have occurred within the Eβ gene. This would make the Eβ vs. Eα system analogous to the system of mutations observed in the H-2Kb antigen (15). As few as two amino acid substitutions, both of the X → lysine or X → arginine type, could explain the differences observed between the Eβ and Eα profiles. The second explanation is that a recombinational event occurred within the gene for the Eβ chain. Hence, the haplotype for D2.GD should be: H-2Kd, Aαb, Bb, Jb, Eβ, Cb, Sb, Db. We feel that the results presented in Fig. 2 strongly support this latter model because it is highly unlikely that two-point mutations in the Eβ gene would give rise to tryptic peptides that co-elute with peptides derived from the Eβ gene.

Current understanding of segmented genes for major histocompatibility complex molecules, e.g., the eight gene segments in a pseudogene for the H-2-linked Qa-2 antigen (16), does not permit one to say whether the crossover event in D2.GD occurred within a coding sequence (exon) or within an intervening sequence (intron) between two exons. Either of these possibilities (both of which would be a classical "intragenic" recombination) would be consistent with our results, and distinguishing between them will require full knowledge of the genomic DNA in this region of the 17th chromosome of D2.GD.

We explored the genetic origin of the Eβ chain further by comparing its amino-terminal sequence (phenylalanines only) to that of the Eα and Eβ chains. The choice of labeled phenylalanine was based on previously published (9, 10) data demonstrating that the positions of phenylalanines in Eα and Eβ chains were different. The results presented in Fig. 3 and Table I suggest that the amino-terminal portion of the Eβ chain, at least up to position 48, is derived from the d haplotype. This observation, when coupled with the information available on the genetic origin of the other H-2 regions of D2.GD, suggests that the orientation of the coding strand for the Eβ chain (at least for D2.GD) is centromeric → telomeric for the 5' → 3' reading direction.

Recently, Rose and Cullen (17) have published a similar study using the recombinant haplotype H-2d11 to suggest that the Aα gene most likely lies to the centromeric side of the Ab and Eβ genes. These results combined with those presented here suggest that the gene order for the I-A subregion of the H-2 complex should be (H-2K), Aα, Aβ, Eβ . . . (H-2D).

Summary

The recombinant strain D2.GD was originally typed as I-A^d by serological methods. Indeed, the Aα and Aβ chains of the I-A antigens appear to exhibit normal behavior by the criteria of serology and two dimensional gel analysis. However, the Eβ chain encoded by the I-A subregion of this strain, one of the two components of the plasma membrane located I-E antigens produced in (D2.GD × A.TFR5)F1 animals, has been demonstrated to be the product of an intragenic recombinational event between Eβ genes from the d and b haplotypes.

Sequence analysis suggests that the amino-terminal portion of the Eβ chain is
derived from the d haplotype and, therefore, that the coding strand for this gene is oriented centromeric → telomeric (5' to 3' direction). Finally, these data combined with the data of Rose and Cullen (17) allow the ordering of the genes within the I-A subregion as (H-2K), A\alpha, \beta, E\beta . . . (H-2D).

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