IDENTIFICATION OF THREE NEW Ig λ-LIKE GENES IN MAN

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An antibody molecule is constructed from four polypeptides, two identical L chains, (κ or λ), and two identical H chains (μ, δ, γ, ε, or α). In its germline state, Ig κ L chain DNA exists in three discrete groups of noncontiguous segments: V, J, and C. H chain–related DNA is slightly more complex, and includes a fourth group of discrete D segments. To form a functional L chain gene, one V region undergoes a site-specific recombination event and becomes contiguous with one J segment. H chain gene formation is a slightly more complicated variation on this theme, which includes the rearrangement of a D segment to a J segment before V-D recombination (for review see 1).

Antibody diversity can be accounted for by combinations obtained from this V-J (or V-D-J) joining and the association within a cell of such recombined L and H chain genes. Toward the expansion of antibody diversity, higher organisms have evolved hundreds of V regions capable of recombining with several separate J segments. In the case of the human κ or H chain gene systems, gene duplication has increased the number of J regions to about six, located just 5′ of the single κ or μ C region (2, 3). Because these J regions encode different amino acid sequences, and each of these J regions is capable of recombining with any appropriate V region, the number of different polypeptide chains that can be made is increased manifold.

The human λ gene system also has increased the number of available J regions by gene duplication. However, in the λ locus, the unit of duplication includes...
the C region as well as the J region. Thus, in humans, there are at least six \( \lambda \) C regions, each apparently associated with its own J region, as was reported earlier (4).

In this paper, we describe a new gene system that provides a potential third class of Ig L chain polypeptides. While most closely related to the previously reported \( \lambda \) L chain system, these genes are clearly distinct. We have characterized two genes from this new class, which have open reading frames and have J regions associated with them. A third \( \lambda \)-like gene is described that appears to be a pseudogene. These \( \lambda \)-like genes have not been described at the protein level, but may represent other L chain alleles that can increase L chain repertoire by providing additional J region coding sequences. Alternatively, these genes may be incorporated into some other gene system requiring a V-J type of activation. Such analogous systems, for example, have already been described as mediating T cell receptor diversity (5-11).

**Materials and Methods**

*DNA Preparation.* High molecular weight DNA was prepared according to Hieter, et al. (4). Intact cellular nuclei were isolated from PBS-washed leukocytes by Dounce homogenization and subsequent centrifugation through a sucrose cushion. The nuclei were then incubated overnight in proteinase K and SDS. DNA was purified from the resultant solution by phenol/chloroform extraction, precipitated in ethanol, and resuspended in deionized water.

*Restriction Enzyme Digestion and Southern Analysis.* All restriction enzyme digests were performed according to manufacturer's suggested conditions. Hybridizations were performed on genomic or cloned DNA cut by restriction endonucleases, size-fractionated on 0.8% agarose gels, and blotted onto nitrocellulose filters as described by Southern (12). Filters were then hybridized to \( ^{32}P \)-labeled probes overnight at 42°C in a 10% Dextran sulfate, 4X SSC, 40% formamide, 0.8% Denhardt's Tris-buffered solution. After hybridization, filters were washed with 2 \( \times \) SSC, 0.1% SDS three times at room temperature, and with 0.1 \( \times \) SSC, 0.1% SDS twice at 52°C before autoradiography.

*Genomic Cloning.* High molecular weight DNA was digested with Eco RI endonuclease and size-fractionated by preparative agarose gel electrophoresis. Appropriately sized fractions were ligated into \( \lambda \) phage vector Charon 4A and packaged. The resulting recombinant phage library was screened as described by Benton and Davis (13). Selected phage DNAs were mapped and areas of particular interest were subcloned into the plasmid pBR327.

*Nucleotide Sequencing Procedure.* Dideoxy sequencing was performed using reagents and instructions prepared by Bethesda Research Laboratories, Gaithersburg, MD.

*Nucleotide Alignments.* Comparisons were made on a DEC-10 computer with the DNA:NUCALN program developed at the National Institutes of Health by Lipman and Wilbur (14).

**Results**

*Identification and Cloning of Three New \( \lambda \) Genes.* Previously we have shown that at least six C region genes (each with its associated J segment) are clustered on an \(~40\) kb stretch of DNA on chromosome 22 (4). Numbered consecutively, the most 5' gene, \( \lambda_{e1} \), resides on a 14 kb Eco RI fragment; \( \lambda_{e2} \) and \( \lambda_{e3} \) on the central 8 kb Eco RI fragment (polymorphic in the human population as 8, 13, 18, and 23 kb pieces, containing from 2-5 gene copies) (15); and the last three genes (\( \lambda_{e4.5.6} \)) on a 16 kb Eco RI fragment. At the time of that analysis it became clear that these six genes represented only a part of the \( \lambda \)-like genes found in the
FIGURE 1. Demonstration of the existence of new λ-like genes. DNA from EBV-transformed B cell lines from the same patient (lane A, a κ producer; and lane B, a λ producer) was digested with the restriction enzyme Eco RI, electrophoresed, blotted, and hybridized with a 32P-labeled probe by the Southern technique (12). As shown in the diagram below, where arrows indicate Eco RI sites, the probe was derived from the C region of the second λ gene in the major functional cluster as described by Hieter et al. (4).

The existence of additional λ-like genes is confirmed by the experiment shown in Fig. 1. In lane A of this Southern blot, germline DNA from a karyotypically normal EBV-transformed κ-producing B cell line has been digested with Eco RI and hybridized to a λ C region probe. It shows the germline 14, 8, and 16 kb bands associated with λ C genes 1-6, plus a faint 18 kb band and the 5 kb processed λ pseudogene (16). In addition there is a faintly hybridizing 10 kb band that remains uncharacterized at this time. In Fig. 1, lane B, DNA from a different EBV-transformed λ-producing B cell line from the same individual, digested with Eco RI, shows the loss of the central 8 kb Eco RI fragment, and two rearranged bands, 2 and 3 kb in size. These correspond to rearrangements
that have occurred within the middle 8 kb Eco RI fragment. In these two recombinations, the more 5' 14 kb Eco RI fragment containing the \( \lambda_c \) gene is deleted from both chromosomes, but surprisingly, a 14 kb band persists. Thus, this band must represent a distinct \( \lambda \) C homologous region found on another 14 kb Eco RI fragment. Cloning experiments confirmed the presence of a 14 kb band that is distinct from \( \lambda_1 \). Interestingly, when similar experiments were performed on the 16 kb band, they revealed another \( \lambda \)-like gene that was distinct from the 16 kb Eco RI fragment containing \( \lambda_c-\lambda_a \). Therefore, Southern analysis and cloning experiments revealed two previously uncharacterized genes of man that share homology with a \( \lambda \) Ig C region.

Fig. 2 shows a comparison of the restriction enzyme maps of \( \lambda_1 \) and the new genes, called 14.1 and 16.1, plus that for a portion of a faintly hybridizing 18 kb Eco RI fragment (18.1) that we also cloned. These genes are homologous to \( \lambda \) C regions (as represented by the large black boxes), yet their restriction maps show that they are clearly different from the previously described major \( \lambda \) gene cluster. In the flanking region, however, the new genes appeared most similar to \( \lambda_1 \). Because the \( J \) region for \( \lambda_1 \) is within 2 kb 5' of the \( C \) region, we examined the equivalent area of 14.1 and 16.1 for a \( J \) region. This search was successful, as demonstrated by nucleotide sequence analysis below.

**14.1 and 16.1 are Nonallelic.** The restriction maps of 14.1 and 16.1 (Fig. 2) are very similar, and it was necessary to determine whether the two clones were
polymorphic alleles or different genes. Therefore, genomic DNA from 14 normal individuals was analyzed with a probe containing the J region from 16.1 (16.1J), which also hybridizes to 14.1, but does not hybridize with λ1 under stringent wash conditions (55°C, 0.1 × SSC). The typical result of this analysis was identical to Fig. 3, lane A. In all 14 samples, three bands were found, two of which correspond to 14.1 (4.3 kb) and 16.1 (6.4 kb), and a third band (7.7 kb), which remains unidentified at this time. Because all persons contained 14.1 and 16.1 bands, it seems unlikely that the two genes are polymorphic alleles.

When the 16.1J probe was used against the λ-producing cell line that had rearranged both λ loci in the major functional cluster, both the 14.1 and 16.1 genes persist in the germline state (Fig. 3, lane B). This result confirms that these genes are not alleles of λc, which is deleted in these cells (data not shown). Further, it supports the idea that these genes are not located between the λV regions and the major λ cluster, a region which is also deleted during V-J recombination.

**Sequence Analysis of 14.1 and 16.1.** To characterize the new genes further, we determined the nucleotide sequence of 14.1 and 16.1 in the areas that correspond to the J and C regions (Figs. 4 and 5). The C regions of 14.1 and 16.1 are encoded as single exons that are 96% homologous to each other and 86–89% homologous to the known λ genes. The number of nucleotides altered, and percentage differences between the genes is given in Table I. Compared to λc, 14.1 and 16.1 have base changes in a total of 42 positions. Of these, 31 alterations are identical in both genes, 3 occur in 14.1 alone, and 8 in 16.1 alone.
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Thus, the two genes are more closely related to each other than to the previously sequenced λ genes.

In addition, the most striking feature of the sequences of 14.1 and 16.1 is that they predict open reading frames. Their deduced amino acid sequences are 95% identical (Table I). When compared to Mcg (the myeloma protein product of λc1) (17) and Kern(−) (the protein made by λc2 and λc3) (18), 14.1 differs by 17-18 amino acids (16-17%) and 16.1 by 22-23 amino acids (21-22%). All of the replacement amino acids caused by the changes can be found at these positions in other human or mouse λ chains that have been reported (19), except methionines 161 and 197, and the alanine in position 205 of 16.1, which are unique.

A comparison of λc1-λc3 reveals a greater divergence of nucleotide sequence than amino acid sequence, as is expected in genes where selective pressure maintains specific protein sequences. In contrast, the percentage of nucleotide sequence differences between 14.1 and 16.1 is less than their percentage of amino acid sequence differences. Similarly, 14.1 and 16.1 are more closely related to λc1-λc3 in nucleotide than amino acid sequence on a percentage basis.

FIGURE 4. Comparison of the nucleotide and deduced amino acid sequences of the C regions of λc1 and 14.1 and 16.1. Dashed lines indicate identity. The site for splicing with the J region is indicated with an arrow.
### Table 1

**Interrelationships between C Regions of Genes**

| C region genes | \(\lambda_{c1}\) | \(\lambda_{c2}\) | \(\lambda_{c3}\) | 14.1 | 16.1 |
|----------------|----------------|----------------|----------------|-----|-----|
| \(\lambda_{c1}\) |                | 17 (5.3)       | 22 (6.9)       | 34 (10.7) | 39 (12.2) |
| \(\lambda_{c2}\) | 5 (4.7)        |                | 9 (2.8)        | 38 (11.9) | 43 (13.5) |
| \(\lambda_{c3}\) | 6 (5.7)        | 1 (0.9)        |                | 39 (12.2) | 44 (13.8) |
| 14.1           | 17 (16.0)      | 17 (16.0)      | 18 (17.0)      |      | 11 (5.5) |
| 16.1           | 22 (20.8)      | 22 (20.8)      | 23 (21.7)      | 5 (4.7) |

*Total bases, 318. Total amino acids, 106. Nucleotide differences are shown above the diagonal; amino acid differences below. Numbers in parentheses are percent differences.*

### J Regions of 14.1 and 16.1

The J segment of \(\lambda_1\) is located 1.4 kb 5' of the C region, and 14.1 and 16.1 each have a single J segment 1.4 kb and 1.3 kb 5' of their respective C regions (Fig. 2). In addition, the detailed structure of these areas is quite similar, and consists of the J coding region flanked on its 5' side by the recognition sequences that presumably mediate DNA rearrangements. At the nucleotide level (Fig. 5), \(\lambda_{j1}, 14.1j\), and \(16.1j\) have a nonamer recognition sequence separated by 10–12 basepairs from a heptamer recognition sequence that abuts the 5' end of the J coding sequence. The J regions of 14.1 and 16.1 are more homologous to each other than to \(\lambda_{j1}\).

For both 14.1 and 16.1, the J coding sequence appears to be one amino acid longer than \(\lambda_{j1}\). This prediction is based on the location of the heptamer recognition sequence. While the alignment of the heptanucleotide sequence appears to have been shifted by point mutations, compensating mutations have occurred that produce a good match with the heptamer consensus sequence. In fact, 14.1 and 16.1 match six and five basepairs, respectively, out of seven with the heptamer CACTGTG sequence of \(\lambda_{j1}\). Similarly, six of nine nucleotides of 14.1 and 16.1 are identical to the nonamer sequence of \(\lambda_{j1}\). This level of conservation of the heptamer and nonamer recognition sequences is comparable to that seen in the human \(\kappa\) and H chain J regions (2, 3), and the human T cell receptor \(\alpha\) chain J regions (11).

The 3' end of each presumptive J region sequence ends with a dinucleotide, -GT, which could be cleaved off with the first nucleotide of the C region to
allow a splice to form a nucleotide triplet, GGT, the codon for amino acid 108, glycine. Thus, the structure of 14.1j and 16.1j are consistent with J regions known to be functional in Ig and T cell receptor genes.

18.1 Is a Pseudogene. A clone of an 18 kb Eco RI fragment (18.1) also contained an area homologous to the \( \lambda_c \) region, as shown in Fig. 2. Nucleotide sequence analysis of this segment revealed that, although it was most closely related to 16.1 and 14.1, it had undergone many more changes than these genes when compared to \( \lambda_c \rightarrow \lambda_3 \). There are three large deletions within the putative coding region, as well as numerous point mutations, which introduce several frameshifts and stop codons (Fig. 6). We have not been able to find a sequence upstream from the 18.1 C region that encodes a J region. Thus, the structure of this gene, with many nucleotide changes and the absence of a J region, suggests that it is a nonfunctional \( \lambda \) pseudogene.

**Discussion**

We provide evidence for the existence of new \( \lambda \) \( \lambda \) chain-like genes in man. Two of them, 14.1 and 16.1, have a structure similar to the known functional locus, with a C region associated with its own J segment. However, they map on distinctly different stretches of DNA, and are nonallelic. In the C region, the nucleotide sequences of 14.1 and 16.1 are most homologous to each other, and less homologous to pseudogene 18.1 and the known functional \( \lambda \) genes. Their most important features are conserved open reading frames associated with appropriate recognition signals for recombination and splicing in the J regions.

Although there may be considerable selective pressure to preserve the coding sequences of functional genes, the flanking sequences may have fewer constraints to vary (20). As a result, interrelationships between the genes can be surmised. We observed that a probe derived from an area flanking \( \lambda_0 \) hybridizes well to 14.1, 16.1, and 18.1, but poorly to \( \lambda_2 \rightarrow \lambda_6 \). Conversely, probes obtained from 16.1j and the corresponding area of 18.1 hybridize best to \( \lambda_1 \), 14.1, 16.1, and 18.1. Probes that originate from \( \lambda_2 \) preferentially hybridize to \( \lambda_2 \rightarrow \lambda_6 \), rather than to the other genes (data not shown).

These findings, in addition to nucleotide sequence analysis, suggest a possible evolutionary model for the human \( \lambda \) gene family (Fig. 7). In this scheme, a common \( \lambda \) progenitor would be the source of the \( \lambda \) genes present in human DNA through gene duplication. The first event would lead to two divergent arms of the \( \lambda \) family, one most closely related to \( \lambda_1 \), and the other, a progenitor of \( \lambda_2 \rightarrow \lambda_6 \). We separate the latter genes from \( \lambda_1 \) at this early stage because the DNA flanking \( \lambda_2 \rightarrow \lambda_6 \) hybridizes poorly with genes on the \( \lambda_1 \) side of the family.
The individual genes $\lambda_2$-$\lambda_6$ would arise from their progenitor gene by further gene duplication; the most recent event appears to have resulted in the generation of $\lambda_2$ and $\lambda_3$, which share extensive homology in both coding and flanking regions. Insufficient data are available at this time about the relationships of $\lambda_4$-$\lambda_6$.

On the other side of the family, the progenitor gene, $\lambda_x$, appears to have given rise to $\lambda_{18.1}$ and pseudogene 18.1. $\lambda_{18.1}$ is the precursor of $\lambda_1$, 14.1, and 16.1. 18.1 appears to have separated from $\lambda_1$, 14.1, and 16.1 before their duplication, because Southern analysis and nucleotide sequence analysis indicate it is approximately equally related to all three genes. Based on nucleotide sequence analysis of the C and flanking regions, 14.1 and 16.1 are most closely related to $\lambda_1$. The suggestion that 14.1 and 16.1 arose from a more recent duplication is supported by the fact that the nucleotide sequences of the C regions of 14.1 and 16.1 differ by ~4%, but both diverge 11-12% from $\lambda_{18.1}$. In addition, most base substitutions in 14.1 also occur in 16.1, although a few mutations have occurred in each gene individually.

The chromosomal location of 14.1 and 16.1 appears to be consistent with the mechanism of gene duplication proposed in this model. While we have not formally assigned 14.1 and 16.1 to human chromosome 22, our analysis of somatic cell hybrids to map the $\lambda$C regions $\lambda_1$-$\lambda_6$ might have shown a discordance for the 14.1 and 16.1 genes had they not been on this chromosome (21). To assign these genes to a chromosome more definitively, we have cloned unique pieces of DNA from the genomic clones and are examining their chromosomal location by in situ hybridization.

Two new genes have been found that can encode $\lambda$-like C regions, and expression of these potentially functional genes is being sought. It is also possible that these two genes, 14.1 and 16.1, may function in a role different from that of the antibody molecule. Recent studies have shown the T cell receptor of man to share homology with $\lambda$ L chains (5), but neither T cell receptor $\alpha$, $\beta$, nor $\gamma$ chains are identical to these genes (5-11). No teleological rationale is obvious to explain the conservation of these $\lambda$-like sequences in man, but selective pressure to maintain their open reading frames is noteworthy. We expect that eventually they will be shown to expand the power and versatility of the humoral immune response, although we cannot rule out their use in an as yet unreported parallel system similar to the Ig $\lambda$ or T cell receptor systems.
Summary

Three new human $\lambda$ L chain–like $\text{Ig}$ genes are identified by restriction enzyme and nucleotide sequence analysis. Two genes, 14.1 and 16.1, have intact J and C regions, and are potentially functional, with open reading frames. A third gene, 18.1, is a pseudogene. The evolutionary lineage of these genes compared to the known functional locus $\lambda_{\text{c}}-\lambda_{\text{o}}$ can be surmised from Southern blot and nucleotide homologies. This study demonstrates that the human $\lambda$ gene family is more complex than previously recognized.

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