ANALYSIS OF A NOVEL $V_n$S107 HAPLOTYPE IN CLA-2 AND WSA MICE
Evidence for Gene Conversion Among Ig$V_n$ Genes in Outbred Populations

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Comparative analyses of Ig variable region genes provide a means for studying evolutionary mechanisms active in large multigene families. The Ig heavy chain V-region ($V_n$) gene families of inbred laboratory mice have been examined in detail, and several potential evolutionary mechanisms have been inferred from these studies. Among these, gene conversion has been suggested as the basis for many $V_n$ allelic differences, particularly in the $V_n$S107 family (1–8). Gene conversion within this family was initially suggested based on the cDNA sequence of a CBA/J hybridoma (6), and subsequent examination of CBA/J germline genes indicated the occurrence of gene conversion in all four members of the $V_n$S107 family (7). Additionally, gene conversion events have been postulated to explain the differences between the $V_n$S107 haplotypes of laboratory inbred strains (8).

Whether gene conversion occurs with significant frequency in outbred populations has not been addressed directly. Extrapolation from the existing data is constrained by both the small group of progenitors common to most inbred laboratory strains, and the possibility that long-term homozygosity fosters events that otherwise rarely occur. Thus, the examination of homologous gene families in feral populations may provide insight regarding this issue.

A wild population of *Mus musculus domesticus* exhibiting low immunologic responsiveness to phosphorylcholine (PC)† was described by Lieberman et al. (9), and several inbred lines have been independently derived from this population. These include the WSA (10; this article) and CLA-2/Cn strains (11). In recent studies, the nucleotide sequence of heavy chain cDNA from a PC-specific hybridoma suggested gene conversion events within the CLA-2/Cn $V_n$S107 family (11). Since the CLA-2/Cn and WSA strains were recently and independently inbred, germline sequences

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Abbreviations used in this paper: PC, phosphorylcholine.
consistent with conversion would strengthen the possibility that such events are wide-
spread and unlikely artifacts of inbreeding. Accordingly, we have examined the
VnSi07 families of the CLA-2/Cn and WSA strains. The results indicate that: (a)
The CLA-2 and WSA strains bear identical but novel VnSi07 family haplotypes;
(b) low PC responsiveness in these populations is unlikely due to an inability to ex-
press the V1 member of the VnSi07 gene family; and (c) gene conversion events have
occurred with considerable frequency in the evolution of the murine VnSi07 family,
especially among the V3, V13, and V11 members.

Materials and Methods

Mice. The CLA-2/Cn strain was derived in the laboratory of M. P. Cancro, from founders
originally obtained from Dr. M. Potter (Laboratory of Genetics, National Cancer Institute,
National Institutes of Health, Bethesda, MD; under the auspices of NIH contract NZ01-CB-
71085). These mice were selectively inbred for PC low responsiveness, and this segregates
as a single autosomal Mendelian trait (11). Adult breeding pairs of the WSA line were also
obtained from Dr. M. Potter. The progenitors of the CLA and WSA lineages were members
of an outbred population, and were initially separated based on coat color and marking traits
before inbreeding regimes (9, 10).

DNA Preparation. High molecular weight DNA was prepared using adult female animals,
freshly sacrificed by cervical dislocation. Nuclei were prepared according to the method of
Liou et al. (12), with some modifications. Briefly, livers were immediately removed, minced,
and dounce-homogenized in ice-cold PBS, containing 1 mM PMSF. All subsequent steps
were performed on ice. The homogenate was strained through two layers of cotton gauze
to remove connective tissue. Cells were spun down and washed several times in fresh PBS/PMSF.
The pellet was resuspended in 25 ml NIB (60 mM KCl, 15 mM NaCl, 5 mM MgCl2, 1
mM EGTA, 15 mM Tris-HCl, pH 7.6, 0.5 mM DTT, 1 mM PMSF), containing 0.3 M sucrose.
Cell membranes were solubilized by the addition of 0.3 ml 10% NP-40 detergent. Aliquots
of 12.5 ml were gently layered over 12.5 ml sucrose cushions (1.7 M sucrose in NIB). Nuclei
were spun down at 4°C for 15 min (13,000 rpm, HB-4 rotor, RC-5B centrifuge, Beckman
Instruments, Fullerton, CA). The supernatant was discarded and the pellet was resuspended
in a total volume of 5.0 ml NIB. Nuclei were lysed by the addition of 0.5 ml 5% SDS/100
mM EDTA and proteinase K to a final concentration of 0.5 mg/ml. Tubes were incubated
overnight at 37°C.

The DNA preparation was extracted once in an equal volume of phenol, previously
preequilibrated with TE (10 mM Tris-HCl, pH 7.5, 1 mM EDTA), extracted twice with
phenol/CHC13 (2/1, vol/vol) and once with CHC13. DNA was precipitated from the aqueous
phase by the addition of 2 vol 100% ethanol, spooled, air-dried, and resuspended to a final
concentration of ~1 mg/ml.

Individual liver DNA samples from 13 members of the original breeding stocks from which
the CLA-2/Cn and WSA strains were derived were generously provided by Drs. L. D'Hoos-
telaere and M. Potter.

Genomic Libraries. Phage genomic libraries were prepared through two different protocols.
Partial Sau3a I digests were performed and fragments ranging from 9 to 23 kb in size were
isolated by fractionation over sucrose gradients (13). Appropriate fractions were pooled, precipi-
tated, and ligated into the Bam H1-digested λ phage vector, λEMBL3 (Stratagene, La Jolla,
CA) according to supplier's instructions. Ligations were packaged with commercial extracts
(Stratagene).

An alternative method for isolation of individual VnSi07 gene segments was also used.
Genomic DNA was digested to completion with Eco R1 and electrophoresed through 0.8% agarose
slab gels. Regions containing VnSi07 gene segments, as determined by Southern blot analysis,
were excised. Gel slices were placed in dialysis tubing for electrophoresion of frag-
ments, which were precipitated and ligated into the vector λgt10 (Stratagene). Reactions
were packaged as described for EMBL3 libraries.
**Library Screening.** λEMBL3 and λgt10 libraries were plated and plaque-lifts were performed in duplicate using nylon membranes (Hybond-N; Amersham Corp., Arlington Heights, IL). Lifts were denatured by floating on 0.2 M NaOH/1.5 M NaCl. Filters were air-dried and baked for 2 h at 80°C. Lifts were washed for several hours with frequent changes of a wash solution containing 50 mM Tris-HCl, pH 8.0, 0.1% SDS, 1.0 mM EDTA, and 1.0 M NaCl at 42°C. Membranes were transferred to 50% formamide buffer for overnight hybridization.

**Probes.** A cDNA probe containing the BALB/c V1 gene segment was originally obtained from Dr. S. Crews (14); California Institute of Technology, Pasadena, CA). A Pst I fragment, containing the V1 coding region, was excised for use as a hybridization probe. A plasmid containing a rearranged V1 gene segment as well as associated 5'-sequences, was obtained from P. Brodeur (Department of Pathology, Tufts University, Boston, MA). A 1.7-kb Bam HI fragment containing V1-specific sequence was used to identify upstream sequences of that gene segment.

Probes were labeled with α-[32P]dCTP using the random-priming method of Feinberg and Vogelstein (15). A commercial kit was used (Oligolabeling Kit; Pharmacia Fine Chemicals, Piscataway, NJ).

**M13 Subclones.** A minimum of two λ phage clones from original, unamplified libraries were sequenced per gene segment. All M13 manipulations were performed as described by Messing (16). Plaque-lifts and screenings were performed as described earlier for genomic library screenings, except that filters were neutralized after denaturation. Subcloning strategies were based on the restriction maps of BALB/c Si107 Vλ phage clones described by Siu et al. (17).

The V13 gene segments of the CLA-2/Cn and WSA strains were subcloned as 2.7-kb Eco RI fragments into Eco RI-digested M13mp18. Clones were also generated by digestion with Sau3A I and shotgun cloning of resulting fragments into Bam HI-digested vector.

The V1 gene segments of both strains were subcloned as 2.5-kb Bam HI fragments into Bam HI-digested vector. Larger, 5-kb Xba I fragments were also subcloned for flanking region analysis.

The V11 gene segments were force cloned into both orientations by double-digestion with the enzymes, Eco RI and Hind III. These fragments were cloned into the vectors M13mp18 and M13mp19. The novel V11 gene segments (V11-2) of the two strains were subcloned as 4.7-kb Eco RI fragments into Eco RI-digested M13mp18.

Orientations of inserts in all M13 subclones were determined either directly, through sequence analysis or through complementation testing (16).

**Dideoxynucleotide Sequencing.** Sequencing of M13 templates was performed with a commercially sequencing kit using supplier's instructions (Sequenase; United States Biochemical Corp., Cleveland, OH).

**Oligonucleotide Primers.** All genes were sequenced in two orientations using the M13 universal primer, as well as a set of primers designed to anneal to particular regions of the coding and flanking sequences.

**Results**

**Si107 VH Allele Assignments.** To examine the contribution of gene conversion to allelism among VλSi107 haplotypes, it is first necessary to distinguish the V1, V11, V13, and V3 family members. Family member assignment was thus based on three criteria. First, genomic Southern blot analysis of Eco RI-digested CLA-2/Cn and WSA DNA reveals the presence of four VλSi107 gene segments (Fig. 1, last four lanes), three of which comigrate with VλSi107 elements in BALB/c (V1, V11, and V13), rendering preliminary family member assignment feasible. DNA from 13 members of the outbred CLA and WSA progenitors are also included in Fig. 1, and show that at least three VλSi107 haplotypes were segregating in this population. Second, coding region sequence analysis was used to distinguish the V1 element
FIGURE 1. Southern Blot analysis of Eco RI-digested BALB/c, CLA-2/Cn, and outbred progenitor DNA. 7.5 μg per lane of Eco RI-digested genomic DNA was electrophoresed, transferred to nylon filters, and hybridized to the p107V1 probe (14). Individual liver DNA samples from members of the outbred progenitor population of CLA-2/Cn and WSA are shown in lanes 1-13. BALB/c and three individual CLA-2/Cn DNA samples are in the last four lanes.

...from the other three gene segments. Finally, noncoding, flanking region sequence analyses were performed. While the coding regions of the V11, V13, and V3 gene segments exhibit a high degree of homology, the flanking, noncoding sequences have diverged considerably, allowing for unambiguous family member assignment of novel VnS107 sequences.

The CLA-2/Cn and WSA V13 Gene Segments are Pseudogenes. As in the BALB/c strain, the V13 gene segments of WSA and CLA-2/Cn lie on 2.8-kb Eco RI fragments. Alleles of V13 examined to date are believed to be functional based on nucleotide sequence analysis (14, 17, 18).

Subcloning of the CLA-2/Cn and WSA V13 gene segments revealed the presence of a novel Sau 3a restriction site within the V13 coding region. Subcloning and sequencing of the gene on the larger, 2.8-kb Eco RI fragment revealed that the novel Sau 3a site is attributable to a single nucleotide insertion within codon 37 (GTC to GATC), resulting in a shift in reading frame (Fig. 2). The new reading frame includes four translational stop codons, likely rendering the V13 gene segments of the CLA-2/Cn and WSA strains pseudogenes. A second mutation is present at codon 82c. This defect involves a single base deletion (CTG to TG) that restores the original reading frame. The defects present in these V13 gene segments are not cloning artifacts, as libraries from the two strains were generated independently. Furthermore, a minimum of three independent clones from unamplified libraries were examined per strain, all of which contained the same coding region defects. No obvious defects are present outside the V13 coding region.
Two V11 Gene Segments are Present in the CLA-2/Cn and WSA Genomes. Two V11 gene segments were identified in the CLA-2/Cn and WSA strains. One of these comigrates with and is identical in sequence to the V11 gene segment of BALB/c (Fig. 3). A second copy of V11 is also present in the CLA-2/Cn and WSA genomes on a novel 4.7-kb Eco RI fragment not present in the BALB/c strain. This extra V11 gene segment will be referred to as V11-2, to distinguish it from the V11 gene segment that comigrates with that present in BALB/c.

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Figure 2. The CLA-2/Cn and WSA V13 gene segments. The coding regions of the CLA-2/Cn and WSA V13 genes are compared with their allel in BALB/c. Insertion and deletion mutations are boxed. The insertion generates four translational stop codons. These sequence data have been submitted to the EMBL/GenBank Data Libraries.
Comparison of the coding and flanking region sequences from CLA-2/Cn V11, CLA-2/Cn V11-2, and BALB/c V11 (Fig. 4) shows that a high degree of homology exists among the three segments. It is clear from the flanking region sequences of the CLA-2/Cn V11-2 gene that it is a second copy of the V11 segment present in CLA-2/Cn and BALB/c.

The CLA-2/Cn and WSA V11-2 gene segments are functional and expressed. All promoter, leader peptide, and heptamer-nonamer rearrangement sequences are present and maintain features believed to be required for V11 expression. Most persuasively, this gene segment encodes the V11 sequence expressed in the heavy chain of the PC-binding, CLA-2/Cn-derived hybridoma, 1C4 (II). The cDNA sequence generated from 1C4 contained differences at six nucleotide positions, relative to the BALB/c V11 gene sequence. Five of these differences are present in the germline sequence of the CLA-2/Cn V11-2 gene segment (Fig. 3). The sixth difference (codon

| Gene      | Sequence                                                                 |
|-----------|--------------------------------------------------------------------------|
| BALB/c    | ACTGACATGAGGGTACTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT      |
| CLA-2/Cn  | TAAGTAAGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT      |
| BALB/c    | TAACTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| BALB/c    | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |
| CLA-2/Cn  | TAACCTAGGAGCAGAATCCTGGAGGGACAGTGGATCTGTGTGACATTTTCCTGATCAAATTCTTTACTATTGCAGGTAT |

**Figure 4.** Flanking region analysis of V11, V13, and V3 gene segments. CLA-2/Cn V11 sequences were designated alleles of V11 or V13 through comparisons of flanking regions with those determined for the BALB/c strain (17). Two different V11 and a single V13 gene segments are present in the CLA-2/Cn germline. There are no V3-like pseudogene segments present in this strain. Promoter (conserved octamer and TATA sequences) and heptamer-nonamer rearrangement sequences are indicated.
95) is not found in any of the CLA-2/Cn SI107 germline sequences and probably represents junctional diversity.

The CLA-2/Cn V1 Gene Segment. In Fig. 5 the CLA-2/Cn V1 gene segment and its associated flanking sequences is compared with its allele in the BALB/c strain. The coding region sequences of the two V1 alleles are identical. Proper promoter elements (the conserved Vn octamer and TATA elements) are present upstream of the CLA-2/Cn V1 gene segment. The heptamer-nonamer rearrangement signal is present as well. While several nucleotide differences exist in the 5'-flanking region sequences of the two strains, none of these would be expected to preclude proper expression of the V1 gene segment of CLA-2/Cn. None of the regions in which these differences occur are conserved in sequence or length among functional Vn gene segments.

CLA-2/Cn and WSA Do Not Possess an Allele of V3. Examination of the Vn SI107 gene segments of the BALB/c, C57BL/10, and CBA/J strains revealed the presence of a pseudogene segment, designated V3 (7, 14, 17, 18). This gene segment has been designated a pseudogene, as it contains several coding region defects, a nonfunc-
ional heptamer-nonamer rearrangement signal and lacks functional promoter and leader peptide sequences. None of the CLA-2/Cn or WSA gene segments contain such defects. While the V13 gene segment of CLA-2/Cn and WSA is a pseudogene, it is clearly not an allele of the V3 gene segment based on Southern blot as well as flanking region sequence analyses. We believe that we have isolated all VnS107 gene segments of the CLA-2/Cn and WSA strains, none of which are alleles of V3.

Discussion

The VnS107 genes of CLA-2/Cn and WSA mice have been examined in detail. The results suggest that: (a) Low PC responsiveness in these populations is unlikely due to an inability to express the V1 member of the VnS107 gene family; and (b) gene conversion events have occurred with considerable frequency in the evolution of the murine VnS107 family.

In accord with preliminary analyses that suggested that PC low responsiveness is unlinked to the IgH complex (11; Cancro, M. P., D. Hilbert, and H. Kersten, unpublished results), the CLA-2/Cn V1 segment contains no features that would preclude its use. All coding regions, including leader peptide sequences, are identical to BALB/c, and no defects are found in promoter or rearrangement sequences. Although a 4-bp deletion is found in a (dG-dT)n:(dC-dA)n dinucleotide repeat in the intron separating the two leader peptide exons of CLA-2/Cn strain, the length of this repeat is variable among all strains examined (7), including those known to express the V1 gene segment. Similarly, the four single base deletions found in this intron relative to BALB/c also are seen in C57BL/10 and CBA/J sequences (7, 8). Neither D nor JH elements are likely responsible for the low responder phenotype, since there is no correlation of D region use with PC specificity, and while a preference for JH1 is observed, it is not requisite for PC binding (19). Additionally, the Jα5 element, which is common to most PC-specific antibodies (19), is intact in CLA-2 and identical to BALB/c (Ferguson, S. E., and M. P. Cancro, unpublished data). Thus, either Vα genes are responsible for low responsiveness, or Ig structural gene defects do not underlie low PC responsiveness.

A large proportion (49%) of differences between alleles of VnS107 gene family members are consistent with gene conversion. In Fig. 6 all known Mus musculus domesticus VnS107 gene segments are compared. Gene segments are grouped as alleles of individual VnS107 family members. The V11-2 gene segment is grouped with other V11 genes. Differences attributed to gene conversion are boxed. Several observations suggest that these differences probably represent a minimal estimate of conversion events.

First, a difference between alleles at a particular nucleotide position was considered a conversion only if both alternative bases could be found at an identical position in at least one other member of the VnS107 gene family. For example, alleles of the V13 gene segment vary at codon position 6 (GAA or GAG). Each of these codons are present in alleles of other VnS107 family members. Because this conservative criterion of gene conversion requires identification of both donor and recipient sequences, some of the nucleotide differences shown in parentheses and assumed here to represent point mutations might also reflect conversion events. For example, at codon position 5 the V13 allele in the C57BL/10 strain contains GTG, whereas the BALB/c, CBA/J, and CLA-2/Cn alleles all contain the codon ATG. This
indeed may reflect a point mutation that occurred in the progenitor of the BALB/c, CBA/J, and CLA-2/Cn V13 alleles but not the progenitor of the C57BL/10 allele. Alternatively, all V13 gene segments may have once contained ATG, but the C57BL/10 allele may have served as a recipient for the codon GTG, using any of the other...
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members of the VnS107 family as a donor sequence. In fact, the nucleotides that flank this difference (positions 6 and 16) are likely the result of gene conversion, strengthening this possibility.

Second, a striking feature of the VnS107 family is the paucity of examples where more than two different bases are observed at a particular nucleotide position (this occurs at only 4 of the 64 positions where differences are observed: in codons 24, 40, 44, and 83). If nucleotide substitutions were random within the VnS107 gene family, the proportion of positions where identical substitutions would exist can be calculated as $(0.33)(1-H)^N$, where $H$ is the degree of homology among the sequences being compared, $N$ is the total number of sequences less the putative donor and recipient, and 0.33 is the probability of choosing one of three possible nucleotides (8, 11, 18). If calculated for the 16 known alleles this value is 21%, and the 49% of observed variations (32 out of 65) consistent with conversion is significantly higher when compared by $\chi^2$ analysis ($\chi^2 = 31.23; df = 1; p < 0.001$).

The distribution of putative conversion events is skewed to particular family members, as 31 of the 32 putative conversion events involve V11, V3, and V13 alleles as donor/recipient combinations. This is consistent with the observation that sequence homology is positively correlated with the rate of intrachromosomal gene conversion (20), since the degrees of homology between V1 and other segments range from only 85-90%, whereas this figure is 90-100% for all V11, V13, and V3 combinations. If only the V13, V11, and V3 members are considered, the expected random occurrence of matching differences is only 13%, but 53% (31 of 59) of the differences among these alleles are consistent with conversion. This deviation from the expected value based on random occurrence is highly significant ($\chi^2 = 81.08; df = 1; p < 0.001$). Thus, nonrandom events consistent with conversion have contributed to VnS107 allelic differences, and are focused among the V11, V3, and V13 alleles.

The paucity of substitutions in V1 attributable to gene conversion might reflect a selective advantage afforded by the specificity of V1-derived antibodies. However, several considerations suggest that this is secondary to homology relationships among individual VnS107 genes. Many of the observed conversions are silent (e.g., amino acid position 6), would not alter antibody specificity, and should thus serve as innocuous donor sequences for V1. Additionally, many conversions occur in framework regions, and it is clear from point mutations among V1 alleles that some framework changes can be tolerated. In fact, some of these appear less conservative than would be a potential conversion at the same position. For example, the gly → arg substitution between C57BL and BALB/c V1 is less conservative than the gly → ala substitution that would result if this codon were converted by the V3 or V13 members of the BALB/c haplotype.

The CLA-2/Cn and WSA sequences extend understanding of several features of the VnS107 gene family. First, although recent comparisons of BALB/c VnS107 members revealed that flanking regions are evolving faster than coding regions (17), this is not true for comparisons made between the alleles of individual gene segments among the CLA-2/Cn and BALB/c strains. In such comparisons (Table I), the V11-2 and V13 gene segments are evolving approximately twice as fast as are the noncoding sequences. Gene conversion may be the basis for the apparent conflict, since it can result in short-term allelic diversity, generating more than one base change through a single event. While we cannot determine the frequency of gene conversion
TABLE I

**Drift between BALB/c and CLA-2 V1S107 Coding and Flanking Regions**

| Comparison | Region          | Number of nucleotides compared | Number of base mismatches | Drift |
|------------|-----------------|--------------------------------|---------------------------|-------|
| CLA-2 V11  | Noncoding flanks| 571                            | 1                         | 0.2   |
| vs. BALB/c | Leader exons    | 57                             | 0                         | 0.0   |
| CLA-2 V11-2| Noncoding flanks| 557                            | 5                         | 0.9   |
| vs.        | Leader exons    | 57                             | 0                         | 0.0   |
| BALB/c V11 | VH segment      | 303                            | 5                         | 1.6   |
| CLA-2 V13  | Noncoding flanks| 569                            | 15                        | 2.6   |
| vs.        | Leader exons    | 57                             | 1                         | 1.6   |
| BALB/c V13 | VH segment      | 303                            | 16                        | 5.3   |
| CLA-2 V1   | Noncoding flanks| 628                            | 7                         | 1.1   |
| vs.        | Leader exons    | 57                             | 0                         | 0.0   |
| BALB/c V1  | VH segment      | 303                            | 0                         | 0.0   |

The drift for alleles of V1S107 family members in BALB/c and CLA was calculated for coding, flanking, and leader exon regions. Drift is calculated as the proportion of positions where base mismatches are observed in the designated region, and is expressed as a percentage.

With these data, it is clear that much gene conversion has occurred even during the relatively short time frame involved in divergence of alleles within a single species.

Second, these data argue against the notion that conversion events seen in the V1S107 family are inbreeding artifacts. The CLA-2/Cn and WSA gene segments are identical in sequence, and these strains were independently derived from the same feral population in which several V1S107 haplotypes were segregating (see Fig. 1). Since it is unlikely that identical mutations occurred independently in each strain subsequent to their separation and inbreeding, these gene segments probably represent those present in one haplotype of the original feral population of CNV mice.

Finally, the composition of the CLA/WSA haplotype strengthens the likelihood that a progenitor haplotype similar to CE may have given rise to many of the extant V1S107 haplotypes. As originally proposed by Perlmutter and Schroeder (18), unequal crossovers within this haplotype could account for the BALB/c and A/J haplotypes through an unequal crossover event. Similarly, the CLA/WSA haplotype could be generated through an unequal crossover in the same progenitor, resulting in two Vf members (Fig. 7).

Based upon the data presented herein, as well as other recent studies suggesting that gene conversion contributes to sequence diversity among mouse and human Vh segments (3, 4, 6-8, 21, 22), it appears that gene conversion globally affects germline Vh genes, and has contributed substantially to Igh locus evolution.

**Summary**

Gene conversion has been suggested as the basis for many Vh allelic differences, particularly in the murine V1S107 family. Whether conversion among IgVh genes
is likely to have occurred in outbred populations has not been directly addressed. The CLA-2/Cn and WSA strains, which were recently and independently derived from a feral population exhibiting low responsiveness to PC, provide the opportunity to approach this question. In previous studies, the heavy chain cDNA sequence of a PC-specific hybridoma derived from CLA-2/Cn suggested gene conversion events within the V\textsubscript{\alpha}S107 family. Accordingly, we have examined the germline V\textsubscript{\alpha}S107 genes of CLA-2/Cn and WSA. The results indicate that: (a) The CLA-2 and WSA strains bear an identical but novel V\textsubscript{\alpha}S107 family haplotype, which lacks a V3 element and contains a V1, a V13, and two V11 genes; (b) low PC responsiveness in these populations is unlikely due to an inability to express the V1 member of the V\textsubscript{\alpha}S107 gene family; and (c) when compared with the other known V\textsubscript{\alpha}S107 haplotypes, the proportion of differences consistent with gene conversion greatly exceeds that expected by random base substitution. Thus, gene conversion events appear to have occurred with considerable frequency in the evolution of the murine V\textsubscript{\alpha}S107 family, especially among the V3, V13, and V11 members.

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