Recent evolution of mouse t haplotypes at polymorphic microsatellites associated with the t complex responder (Tcr) locus

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Summary
Microsatellites closely associated with each member of the TcplO gene family were amplified simultaneously from t haplotype and wild-type forms of mouse chromosome 17, by PCR. The t complex responder (Tcr) locus, which plays a central role in transmission ratio distortion, maps within the TcplO cluster on the t haplotype. Thus the amplified set of microsatellite loci (referred to collectively as Tcpl10ms) provides a direct marker for this central component of the meiotic drive system associated with all naturally occurring t haplotypes. A unique Tcpl10ms pattern of microsatellite alleles was obtained for a number of independent, laboratory-maintained complete and partial t haplotypes. Independent t chromosomes found in wild mice from US populations also had unique patterns, even when they were classified within the same lethal complementation group. Wild and laboratory chromosomes in the tNS group showed similarly-sized but non-identical Tcpl10ms patterns, suggesting they share a recent common ancestor. These chromosomes are likely to have derived from an ancestral chromosome within the founding population of North American house mice. The Tcpl10ms pattern was also shown to be useful in field studies for distinguishing among independent t haplotypes, when more than one is present within a single population.

1. Introduction

t haplotypes are geographically widespread variants of the proximal third of chromosome 17 that occur in natural populations of house mice (Silver, 1985). Complete t haplotypes are typified by their capacity for meiotic drive, or Transmission Ratio Distortion (TRD). Although segregation in females is normal, males heterozygous for a wild type (+) and a t haplotype (t) form of chromosome 17 will transmit the t haplotype to greater than 95% of their offspring in a clear departure from Mendelian segregation (Dunn, 1957).

The genetic basis for t-specific transmission ratio distortion is complex, and results from the interactions of a number of cis and trans-acting linked distorter loci (Tces) and a haploid-acting responder locus (Tcr) (Lyon, 1984; Lyon, 1986; Silver & Remis, 1987). The structural integrity of these loci is maintained through the suppression of recombination along the length of t haplotypes, by a series of 4 major inversions (Herrmann, Barlow & Lehrach, 1987; Hammer, Schimenti & Silver, 1989). Collectively these span almost the entire t DNA and result in a 50–100-fold reduction in normal levels of recombination. Each of the inverted regions carries one or more of the loci required for maximal expression of TRD (Hammer, Schimenti & Silver, 1989).

A candidate gene for Tcr, Tcpl10b, has been characterized in the Tcpl0 gene family. Members of this gene family are present in 2 to 4 copies on both homologues of chromosome 17 from different strains of mice (Schimenti et al. 1987; Schimenti et al. 1988; Bullard & Schimenti, 1990). Tcpl10b produces a unique, haploid-specific, alternatively spliced mRNA in elongating spermatids, consistent with its putative role as Tcr (Cebra-Thomas et al. 1991).

The high transmission bias in favour of t haplotypes is expected to lead to their rapid fixation in mouse populations; however, TRD is counterbalanced both by the complete sterility of tt homozygous males, and by the recessive lethal mutations carried by most complete t haplotypes. Thus t haplotypes are maintained as a polymorphism in natural populations (Petras, 1967; Figueroa et al. 1988; Lenington, Franks & Williams, 1988). t haplotypes express no obvious phenotype in wild mice, and only with the recent
availability of molecular probes that identify re-
striction fragment length polymorphisms (RFLPs)
between t and wild-type DNA, has it become possible
to definitively characterize the genotypes of large
numbers of wild mice.

Phylogenetic analyses of wild-type and t haplotype
forms of chromosome 17 indicate that all contem-
porary t haplotypes can be traced back to a common
ancestral chromosome that may have existed as
recently as 10000 years before present (b.p.), and not
more than 100000 years b.p. (Willison, Dudley &
Potter, 1986; Morita et al. 1992; Hammer & Silver,
1993). This recent common ancestry is further
supported by the observation of identical t-specific
protein polymorphisms and by a scarcity of RFLPs
among different t haplotypes (Silver et al. 1983; Fox
et al. 1985; Schimenti et al. 1987; Howard et al. 1990;
Horiuchi et al. 1992). Thus, although many molecular
markers have been characterized that detect
differences between wild-type and t chromosomes, in
nearly all cases, these markers do not provide a means
for distinguishing among the different t haplotypes.

One exception to this rule is provided by the Bb40
marker which detects two different t-specific RFLP
patterns based on the presence or absence of a Tcpl0
pseudogene named Tcpl0ps (Schimenti et al. 1987;
Horiuchi et al. 1992; Pilder et al. 1992). Another more
extensive set of inter-t polymorphisms is detected by
the TSE marker (Uehara et al. 1990). Unfortunately,
the Bb40 polymorphism is bi-allelic and only divides
the complete set of t haplotypes into two broad
groups. Although the TSE polymorphism uniquely
identifies individual t haplotypes, its usefulness is
limited by the requirement for very high molecular
weight DNA that must be analysed by pulsed field gel
electrophoresis, and it is rarely feasible to use this
technique for samples collected in the field, in
particular. Thus the different t haplotypes have
remained largely distinguishable only on the basis of
their traditional lethal complementation groups.

Microsatellites are regions of DNA composed of
short simple sequences repeated in tandem, such as
dinucleotide repeats (e.g. (CA)ₙ) (Tautz, 1993). They
are abundant and widespread in eukaryotic DNA
(Hamada, Petrino & Kakunaga, 1982; Tautz & Renz,
1984), and show a high degree of polymorphism in
length variation of repeat units. Because microsatellite
polymorphisms can be detected rapidly by PCR, they
provide excellent markers for genetic analysis (Weber
& May, 1989; Love et al. 1990; Dietrich et al. 1992).
They are also extremely useful in studies of variation
at the level of populations, even in species where other
genetic techniques reveal little to no detectable genetic
polymorphisms (Bruford & Wayne, 1993; Hughes &
Queller, 1993).

Dinucleotide repeats have also proven useful in
finding polymorphisms among t haplotypes. In
contrast with the lack of polymorphism between
t haplotypes for most RFLP probes, many of the
dinucleotide repeats that have been studied have
shown some polymorphism (Uehara et al. 1990;
Ebersole, Lai & Artzt, 1992; Lai & Artzt, 1992). They
not only detect differences between t haplotype and
wild type chromosomes, but have also been shown to
differentiate among some of the different t haplotypes
(Lai & Artzt, 1992).

To find polymorphic differences that distinguish
among t haplotypes, primers were designed around a
short microsatellite repeat in an intron of the Tcpl0b¹
gene. Because this gene is a candidate for the t
complex responder (Tcr), this provides an important
means for directly assessing inter-t variation at this
locus, which is of central importance to the meiotic
drive phenotype expressed by t haplotypes.

2. Materials and methods

(i) Mice and DNA

DNAs were all prepared from mice that had been
maintained in the animal colony at Princeton Uni-
versity, or were gifts to M. F. Hammer as part of an
earlier study (Hammer & Silver, 1993). Cosmid clones
containing the Tcpl0d¹, Tcpl0b¹, and Tcpl0c¹ alleles
were derived from a homozygous t⁶⁵ cell line cosmid
library provided by Dr John Schimenti (Rosen et al.
1990).

(ii) Wild caught animals

Wild mice were live-trapped in various locations
around North America. Tail biopsies were collected
from wild caught animals and preserved in ethanol
until being brought back to the laboratory. DNA was
then isolated using a standard tail DNA preparation
protocol (Hogan et al. 1994).

A number of t haplotypes from wild-caught mice were
bred to 129/SvJ animals to control the contribu-
tion to the PCR pattern from the wild-type
chromosome. The wild-caught t haplotype chromo-
somes tested here are: BM1 (a semi-lethal t haplotype
trapped in New Jersey); MV1, MV3, and MV12
(three semi- lethals from a single population trapped
elsewhere in New Jersey); PV1 (a semi-lethal t
haplotype trapped in Southern Illinois); CF109 (a
lethal t haplotype trapped in Ithaca, NY); and T57,
T59, T67, and T68 (four lethal t haplotypes collected
from a population in Tennessee). All t haplotype-
bearing mice were also confirmed independently, both
by breeding assays, and by Southern blot analysis
using the markers Bb40 and Tu119 which distinguish
RFLPs between t haplotype and wild type forms of
chromosome 17 (Herrmann et al. 1986; Schimenti et
al. 1987).

In breeding analyses, CF109, and all of the
Tennessee chromosomes failed to produce viable
homozygotes, indicating that all carried recessive
lethal alleles (data not shown). Furthermore, all
eXcept T68 failed to complement the t⁶⁵ chromosome,
and therefore carry the \( r^{os} \) lethal mutation. T68 did complement with both a number of the other chromosomes, and \( r^{os} \), indicating that it contains a different undetermined lethal allele.

(iii) **PCR analysis**

PCR primers were designed in unique sequence flanking a (CA)\(_{34}\) microsatellite located in the intron following the first amino acid coding exon of the Tcp10\( b^1 \) gene from \( r^{os} \) (Bullard & Schimenti, 1991). The complete amplification pattern obtained with this primer pair is referred to as Tcp10ms (for Tcp10 family microsatellites). Unpublished sequence for this region was kindly supplied by Dr John Schimenti. The sequence of the forward and reverse primers are,

\[
\text{TcplOms F: (5'-GCGTGCCCCTTGACAGGG-3')}
\]

and

\[
\text{TcplOms R: (5'-GCTGTACTGTAACCTTGCTTAG-3')}
\]

PCR reactions were performed both with and without radioactive end-labeling of one of the primers. The products were visualized initially on non-denaturing acrylamide gels and subsequently, for better resolution of alleles, radioactively-labelled, on denaturing sequencing gels. Approximately 50 ng of genomic DNA was amplified in a 10 \( \mu \)l PCR reaction using 0.25 units of Taq polymerase. 2.5 \( \mu \)M each of the two unlabelled primers were used, with an additional 0.75 \( \mu \)M of one of the primers end-labelled. The reactions were amplified for 25 cycles; at 95 °C for 25 s, 60 °C for 1 min, and 72 °C for 30 s. PCR products were electrophoresed on 7% denaturing polyacrylamide gels (SequaGel, National Diagnostics). Gels were dried and exposed to film at room temperature for 4–16 h. The amplified products ranged in size between ~ 250–300 bp. A sequencing reaction was used as a size marker to determine relative allele sizes.

(iv) **Sequence analysis**

Direct sequencing of the *Mus macedonicus* PCR product, to determine its microsatellite repeat length, was performed by reamplifying the initial product using one kinased primer and digesting with lambda exonuclease (Higuchi & Ochman, 1989). Primers were removed from the exonucleased reaction (Kreitman & Landweber, 1989), and the ssDNA was sequenced in both directions using the Sequenase Kit (USB). Reactions were electrophoresed on a 6% polyacrylamide gel. Gels were soaked in a 10% methanol/10% acetic acid bath, dried onto 3 mm Watman paper and autoradiographed.

3. Results

(i) **Simultaneous detection of all members of the Tcp10 gene family**

The primer pair defining the Tcp10 microsatellite was used to amplify products from the various DNA samples described in the Materials and Methods. The amplification pattern obtained with each whole genome sample (referred to as TCP10ms) was complex. This is because the primer pair simultaneously amplifies microsatellite alleles from all members of the Tcp10 gene family, both on the \( t \) haplotype and the wild type forms of chromosome 17 (Fig. 1). Individuals may vary both in the number of Tcp10 genes they contain, and in the sizes of the allelic products amplified from each of these genes.

To determine the correspondence between individual components of the complete TCP10ms pattern and particular Tcp10 genes from the \( t \) form of chromosome 17, we amplified single alleles from individual cosmid clones containing the Tcp10\( a^1 \) (CosA), the Tcp10\( b^1 \) (CosB), or the Tcp10\( c^1 \) (CosC) genes. The results obtained are shown beside the complete pattern obtained from the wild-type 129/SvJ...
Fig. 2. Autoradiograph of PCR products obtained for various complete and partial \( t \) haplotypes. The independent \( TcplO \) alleles of \( r^{es} \) are shown in the cosmid clones A, B and C, beside the 129/SvJ wild type bands, and the 129/\( r^{es} \) composite heterozygote animal is shown nearby for comparison. Most \( t \) haplotypes are shown against a 129 wild type chromosome unless otherwise indicated. The following chromosomes are partial \( t \) haplotypes: \( r^{es}, r^{e}, T^{e}, t^{e}, T^{es}, r^{es} \) (Committee for Mouse Chromosome 17, 1991). All others are complete \( t \) haplotypes. For allele sizes see Table 1.

Fig. 3. Phylogenetic tree of the \( Mus \) species related to the \( Mus \) species group. \( t \) haplotypes are present in natural populations of all four subspecies of the \( Mus \) species group, \( domesticus, musculus, castaneus \) and \( bactrianus \). Modified from fig. 2.2 (Silver, 1995).

Fig. 4. Autoradiograph of PCR products of wild type \( TcplO \) alleles obtained for four inbred strains of \( M.  m. \) domesticus and a number of related species of \( Mus \). The 2 weakly amplifying bands from \( Mus \) spretus can just be seen.

(ii) Species differentiation within the \( Mus \) genus

In addition to amplifying alleles from the wild-type \( M. m. \) domesticus and \( t \) haplotype chromosomes, the \( TcplO \) primers also amplify products from other species and sub-species within the \( Mus \) species group (Fig. 3). In both \( M. m. \) musculus, and \( M. m. \) castaneus, at least two predominant PCR products are observed (Figs 4, 5) suggesting that \( TcplO \) may have more than

Fig. 2. Autoradiograph of PCR products obtained for various complete and partial \( t \) haplotypes. The independent \( TcplO \) alleles of \( r^{es} \) are shown in the cosmid clones A, B and C, beside the 129/SvJ wild type bands, and the 129/\( r^{es} \) composite heterozygote animal is shown nearby for comparison. Most \( t \) haplotypes are shown against a 129 wild type chromosome unless otherwise indicated. The following chromosomes are partial \( t \) haplotypes: \( r^{es}, r^{e}, T^{e}, t^{e}, T^{es}, r^{es} \) (Committee for Mouse Chromosome 17, 1991). All others are complete \( t \) haplotypes. For allele sizes see Table 1.
Mouse complex responder microsatellites

Artifactual bands, however, may be due to secondary structures formed within the sample, as these can usually be eliminated by denaturing the PCR products and running them on sequencing gels (compare Figs 4 and 5). When the PCR products here were initially run on non-denaturing gels, Mus macedonicus alone did not show any echo or artifact bands, and had the smallest PCR product (see Fig. 5). Direct sequencing of this PCR product, to determine whether it’s small size was due to fewer dinucleotide repeats, revealed that the number of dinucleotide repeats was in fact reduced from (CA)\textsubscript{64} to (CA)\textsubscript{3}. This suggests that the ‘echo’ bands seen on these non-denaturing gels are internally derived and are a function of both the presence, and the number, of such dinucleotide repeats, and are probably conformational.

Fig. 5. Ethidium bromide stained, non-denaturing acrylamide gel of Tcp10 PCR products, amplified from two inbred strains of M. m. domesticus, and a number of related species of Mus. The two very faint bands amplifying from Mus spretus can just be seen. Mus macedonicus clearly has the smallest PCR product, and is also the only PCR product without associated ‘echo’, or other larger, PCR artifact bands.

One family member in these subspecies as previously shown for M. m. domesticus. The Tcp10 primers also amplify products in some of the more distantly related Mus species. A single band was amplified from Mus macedonicus, and the same sized primary band, as well as some additional fainter bands, were amplified from Mus spicilegus. These 2 species have been shown by a number of molecular phylogenies to be sister groups (Lundrigan & Tucker, 1994). At 60 °C, two very faint bands can be amplified from Mus spretus. Nothing can be amplified from Mus caroli at this temperature, but a single band can be amplified from Mus cookii, which is more distantly related to the Mus musculus sub-group than is Mus spretus. No products could be amplified from either rat or hamster (Fig. 5), nor human DNA (data not shown). Reducing the annealing temperature to 50–55 °C didn’t resolve any of these, and resulted in many non-specific bands. Both the weak bands of M. spretus, and the failure to amplify these other microsatellite alleles, may be due to base variation in the region of primer annealing (Callen et al. 1993).

Individual PCR amplified microsatellites visualized on agarose, or non-denaturing acrylamide gels, frequently show a single main product and an artifactual ladder of additional bands (Fig. 5). Some of these, particularly those near or below the size of the desired band, have been attributed to polymerase slippage (Litt & Lucy, 1989; Weber & May, 1989). This apparent strand slippage during PCR amplification is most common for dinucleotide microsatellite repeats (Beckman & Weber, 1992). Higher molecular weight (iii) Different complete \( t \) haplotypes display characteristic patterns

A survey was performed on the Tcp10 microsatellite family contained within different laboratory-maintained complete and partial \( t \) haplotypes as shown in Fig. 2. This survey demonstrated distinct patterns of Tcp10 microsatellite alleles for each independent \( t \) haplotype. Different \( t \) haplotypes are polymorphic with respect to the band sizes of any given Tcp-10 microsatellite allele, as well as to the number of visible bands (Table 1). Thus, each \( t \) haplotype has a unique set of microsatellite bands, and it is possible to distinguish between almost all of the different \( t \) haplotypes tested.

Complete \( t \) haplotypes carry three functional Tcp-10 genes, and some may carry a fourth copy as a pseudogene, while wild type chromosome 17 homologues may carry between two to three different functional Tcp10 genes (Bullard & Schimenti, 1990; Pilder et al. 1992). Thus a /+-/- animal can potentially have up to 6 or 7 different Tcp10 amplification products. Rarely are so many bands visible, however. Presumably some alleles are not visible because they overlap in band size with other alleles. An example of this can be seen in the 129/\( t \)\textsuperscript{29} compound heterozygote (Fig. 2), where the Tcp-10\textsuperscript{c} allele from \( t \)\textsuperscript{33} can be seen clearly. This allele is only slightly larger than the Tcp-10 alleles of the 129 chromosome, and is therefore obscured in the 129-\( t \)\textsuperscript{33} heterozygote, which thus appears to have only two \( t \)-associated alleles.

There is also a certain amount of size polymorphism associated with any given gene in the \( t \) haplotype complex of Tcp10. The cosmid-amplified Tcp10\textsuperscript{b} allele is derived from the \( t \)\textsuperscript{33} chromosome, and can be seen clearly in the 129-\( t \)\textsuperscript{33} individual. The partial \( t \) haplotype \( t \)\textsuperscript{1001} (C3H background) is a partial \( t \) haplotype containing only the \( t \) form of the Tcr, or Tcp10\textsuperscript{b} gene, and it is not the same size as the allele from \( t \)\textsuperscript{33}. Similarly, the \( t \)\textsuperscript{1002} chromosome, which is a naturally occurring partial \( t \) haplotype from Israel.
Table 1. Relative allele sizes and Tcp10ms band patterns of laboratory-maintained and wild-caught t haplotype chromosomes

| Genotype          | Origin* | Allele pattern and size (bp)† |
|-------------------|---------|-----------------------------|
| 129/SvJ           | Inbred  | 293, 292                    |
| C3H               | Inbred  | 288                         |
| B6                | Inbred  | 298, 278                    |
| 129/rt5‡          | NY, USA | 129/SvJ, 294 (Tcp10c), 274 (Tcp10b), 266 (Tcp10a) |
| 129/rt5/12        | Paris lab non-inbred | 129/SvJ, 286, 278, 276 |
| 129/rt5/12        | NY/PA, USA | 286, 274, 270 |
| B6/rt12           | USA     | B6, 280, 268                |
| B6/rt12           | USA     | B6, 280, 268                |
| 129/rt5/12        | USA     | 129/SvJ, 294, 274, 273     |
| 129/rt5/12        | Denmark | 129/SvJ, 290, 278          |
| 129/rt5/12        | USA     | 129/SvJ, 294, 274, 272     |
| 129/rt5/12        | Italy   | 129/SvJ, 286, 276, 270     |
| 129/rt5/12        | Italy   | 129/SvJ, 284, 278, 270     |
| 129/rt5/12        | Italy   | 129/SvJ, 288, 278, 270     |
| 129/rt5/12        | Italy   | 129/SvJ, 282, 278          |
| 129/rt5/12        | Italy   | 129/SvJ, 286, 276          |
| B6/rt5/12         | Chile   | B6, 274, 270, 258          |
| B6/rt5/12         | Germany | B6, 286, 278                |
| rt5/12/12         | Israel  | 280                         |
| 129/T58           | TN, USA | 129/SvJ, 294, 278, 272     |
| 129/T5            | TN, USA | 129/SvJ, 294, 278, 272     |
| 129/T67           | TN, USA | 129/SvJ, 294, 278, 272     |
| 129/T68           | TN, USA | 129/SvJ, 294, 278, 272     |
| 129/CF109         | NY, USA | 129/SvJ, 294, 274, 272     |
| BM1/BM1           | NJ, USA | 280, 276                    |
| PV1/PV1           | IL, USA | 280, 274, 264               |
| MV1/MV1           | NJ, USA | 286, 274                    |

* Data on origins of laboratory t haplotypes taken from Silver et al. (1987), and described in materials and methods for wild-caught chromosomes.
† May not represent all alleles, as identical or similar-sized alleles will appear as a single allele. Only chromosomes with distinct patterns in Fig. 2 are represented here.
‡ The exact size and identity of alleles is known for rt5 only.

(Silver et al. 1987), only has the Tcp-10a haplotype, and it too differs in size from the cosmid Tcp10da allele from rt5.

A few t haplotypes are indistinguishable from one another. rt5/12 and rt6/10 produce identical PCR patterns. Both of these are complete t haplotypes originally derived from Italy (Silver et al. 1987), and it is therefore possible that they are related chromosomes. The chromosomes rt5/12 and rt12 are also identical at all bands, further confirming that they are probably the same laboratory-derived t chromosome, as suggested previously by Arizt et al. (1985) and Uehara et al. (1990).

These primers additionally allow some fine scale mapping of some of the recombination breakpoints of partial t haplotypes. rt5/12 is a partial t haplotype derived from a recombination event near the Tcr locus, between the complete t haplotype, rt5, and a partial r haplotype (rt6/10/rt12) carrying the responder locus from the r haplotype (Lyon, 1984; Lyon & Zenthon, 1987). From the data presented here (Fig. 2), it can be seen that rt5/12 has the Tcp10a haplotype from rt5 but has the Tcp10b haplotype (i.e. the responder form of the allele) from r. This is consistent with other recent findings showing that the crossover event occurred between these two members of the Tcp10 gene family, which are adjacent to one another on the t haplotype chromosome (Bullard, Ticknor & Schimenti, 1992).

(iv) Polymorphism among wild-type and t haplotype chromosomes in natural populations

Wild-caught animals that are heterozygous for t haplotypes have uncharacterized, and potentially highly polymorphic, wild-type forms of chromosome 17. This variation was found to be sufficiently great that the resulting overall PCR pattern was too confusing to interpret directly in these animals. Although t haplotype-specific alleles are, on the whole, smaller in size than those from wild-type chromosomes, this distinction between t and wild-type chromosomes is less clear in wild animals. To overcome this problem, t haplotypes derived from a number of wild mice were bred opposite to the inbred 129/SvJ form of chromosome 17. With a defined
wild-type contribution, it was then possible to distinguish those PCR products that derived from wild-caught t haplotypes.

None of the PCR patterns associated with wild-caught t haplotypes were identical to any of the laboratory-maintained t haplotypes tested (Figs. 2 and 6, and Table 1). Among the wild-derived lethal haplotypes, both the CF109 and three of the four Tennessee chromosomes (T58, T59, T67) were classified as belonging to the t\textsuperscript{m5} lethal complementation group. All of these haplotypes show a general similarity in banding pattern with the t\textsuperscript{w5} chromosome, however, none are identical to it. The alleles that appear to correspond to the Tcp10\textsuperscript{a} and/or Tcp10\textsuperscript{b} alleles differ in a number of repeat units from those in the t\textsuperscript{w5} chromosome. T58, T59 and T67 have a Tcp10\textsuperscript{a} allele that is 6 bp larger, and a Tcp10\textsuperscript{b} allele that is 4 bp larger than the corresponding products in t\textsuperscript{w5} respectively. The CF109 chromosome was trapped in Ithaca, NY, from the same region from which t\textsuperscript{w5} was originally collected. It has the same sized Tcp10\textsuperscript{a} allele as the other T-chromosomes, but a Tcp10\textsuperscript{b} allele that is 4 bp smaller and corresponds in size to the same allele in t\textsuperscript{w5}. Overall, these chromosomes are all relatively similar to one another and to the t\textsuperscript{w5} lethal (Silver, 1983).

The recently trapped semi-lethal haplotypes (BM1, PV1, and MV1 through MV23), can also be seen to bear only a slight resemblance to the previously known semi-lethal haplotypes shown here (t\textsuperscript{m2}, t\textsuperscript{hob} and t\textsuperscript{w28}). Because of their phenotype of incomplete lethality, the semi-lethal alleles cannot be classified genetically into separate complementation groups, and are therefore considered as a single class.

Semi- lethals collected from different populations show quite different allele patterns. Semi- lethals derived from the same population, in contrast, have identical allele patterns (Fig. 6). Mice trapped from the MV population in New Jersey were known from breeding studies to contain a semi-lethal allele, however, it was impossible to know from breeding studies alone, whether they contained one or two such haplotypes. DNA’s from all mice trapped from the entire population were then PCR amplified, to determine whether the population contained one, or more than one, semi-lethal and/or lethal t haplotype. All mice that carried a t haplotype in this population were found to be carrying the same one. The presence of any two independent t haplotypes could not have been discerned previously in the absence of breeding studies.

4. Discussion
All t haplotypes are descendants from a recent ancestral chromosome (Willison et al. 1986; Morita et al. 1992; Hammer & Silver, 1993). As a consequence of their close relationship to each other, there is minimal inter-t variation at the molecular level. This has made it difficult to distinguish different t haplotypes, particularly in wild populations, with the use of molecular markers alone. Here we show that a single pair of PCR primers designed around a microsatellite locus in the Tcp-10\textsuperscript{b} gene can provide a means for distinguishing among nearly all well-characterized t haplotypes as well as among undefined t haplotypes from independent wild populations.
two to four copies on both the wild type and \( t \) haplotype homologues of chromosome 17. The various members of the gene family are highly conserved relative to one another, which seems to be the result of the concerted evolution of an ancient gene family (Pilder et al. 1992). Despite the high level of identity of genes in this gene family (Davies & Willison, 1991), they display considerable variation in repeat length of the microsatellite allele among the different members. All of the inbred strains tested show characteristic size differences in their alleles, and there is a similarly high level of variation in the alleles contained on the \( t \) haplotype chromosomes.

The ability to uniquely distinguish all classically-defined \( t \) haplotypes has been demonstrated by only one other means, through clusters of \( t \)-haplotype specific elements (TSES) unique to the \( Tla \) region of \( t \) haplotypes (Uehara et al. 1990). The variation observed at both the TSE andTcp10 clusters uniquely identifies individual \( t \) haplotypes to an even greater extent than the only previously available method for distinguishing among \( r \) haplotypes, which is based on complementation analysis.

Uehara et al. (1990) showed that the extensive diversity in TSES in independent \( t \) haplotypes was due to different numbers of clusters and different numbers of repeats in each cluster, which may reflect a pronounced instability of TSE clusters on an evolutionary time scale. Microsatellite alleles are similarly unstable and show relatively high mutation rates (e.g. 10\(^{-3}\)–10\(^{-8}\) (Weber & Wong, 1993; Ellegren, 1995), thus none of the diversity found in either of these highly mutable clusters is at odds with the finding that all \( t \) haplotypes share a recent common ancestor (Morita et al. 1992; Hammer & Silver, 1993).

The conservation of microsatellites between taxa has been demonstrated in a number of studies. Dinucleotide repeats in particular, are conserved by chromosomal location between closely related species (e.g. mouse and rat, and sheep and cattle) (Moore et al. 1991; Hino et al. 1993; Stallings, 1995), but not between more distantly related species (e.g. humans and rodents) (Stallings et al. 1991; Stallings, 1994). The microsatellite repeat investigated here, is conserved in a number of closely, and more distantly, related species of \( Mus \). However, it amplifies only weakly from \( Mus \) \textit{spretus}, and not at all from the rat, hamster, or human, even though humans have been demonstrated to have a homologue of the \( Tcp10 \) gene family (Islam et al. 1993). This non-amplification could result either from the absence of the microsatellite in these species, or from DNA sequence divergence in the region of primer annealing, which can produce null alleles (Callen et al. 1993; Pemberton et al. 1995).

Despite the relatively high mutation rate associated with microsatellite loci over evolutionary time scales, our results show that microsatellite loci can be sufficiently stable over shorter time periods to be used as markers to follow individual \( t \) chromosomes through multiple generations. The Tcp10ms patterns observed for both the \( r^t \) haplotype and the inbred 129 wild-type chromosome were found to have remained unchanged over, at least, the 12 years that these chromosomes have been bred in this laboratory (a number of these are shown in Fig. 2). Similarly, the identical patterns of \( r^{t2} \) and \( r^{u2} \) supports previous suggestions that these two chromosomes are descendants from the same chromosome. These two chromosomes have been maintained in separate stocks since 1957 (Silver, 1983) during which time their Tcp10ms patterns have not diverged (Fig. 2).

In contrast, while newly identified wild-derived chromosomes from the \( r^{o3} \) complementation group (CF109, T57, T59 and T67) share similar TCP10ms patterns relative to one another and to the laboratory defined \( r^{o3} \) haplotype, they are all distinguishable on the basis of small allele size differences of 2–6 bp at the Tcp10a and Tcp10b loci. Both empirical and theoretical research on the mutational generation of new microsatellite alleles (Tautz & Renz, 1984; Schlötterer & Tautz, 1992; Valdes, Slatkin & Freimer, 1993; Stephan & Cho, 1994), suggest that new alleles form predominantly through processes such as slipped strand mispairing, so that the size of a new allele depends on the size of the allele that mutated. Thus the small allele size differences seen here among alleles of the \( r^{o3} \) complementation group may well reflect a recent common ancestry of these alleles.

It seems likely that house mice probably invaded North America for the first time with their human counterparts from Western Europe sometime during the early colonial period ~1620–1650. While a large number of different \( t \) haplotype complementation groups are found in Europe (Klein, Sipos & Figueroa, 1984) only a relatively small number of \( t \) complementation groups are found in North America, and a single lethal, \( r^{o3} \), predominates (Bennett, 1975). Thus all current North American \( t \) haplotypes are probably descendants from a few original founders that existed no more than 375 years ago. Among the independent members of the \( r^{o3} \) group investigated here, all of the observed changes in Tcp10ms patterns are small, and are therefore likely to have occurred during this time period.

In summary, the Tcp10ms pattern of PCR products provides a powerful tool for the analysis of \( t \) haplotypes in both laboratory and field studies. In the laboratory, the Tcp10ms pattern can be used as a means to test and control for the purity of different \( t \) bearing stocks that are not easily distinguished except through complementation testing. In the field, the Tcp10ms pattern can be used for population genetic studies where identifying particular alleles is not of concern (Bruford & Wayne, 1993; Slatkin, 1995). Moreover, the pattern can be used to demonstrate the presence of a single, as opposed to multiple, \( t \) haplotype founders within a given population.
Mouse t complex responder microsatellites

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