Homologous recombination in the tandem calmodulin genes of *Trypanosoma brucei* yields multiple products: compensation for deleterious deletions by gene amplification

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Homologous recombination between a calmodulin–neomycin-resistance fusion gene and the *Trypanosoma brucei* chromosome takes place not only in the large 5’- and 3’-flanking segments of the calmodulin locus but also in any of the four tandem genomic calmodulin genes. This results in a recombined locus consisting of the chimeric neo' gene and four, three, two, one, or zero functional calmodulin genes. Cells bearing this latter event have half of their normal number of intact calmodulin genes and an accompanying phenotype of slow growth. Over months of propagation, these lines acquire additional calmodulin genes, frequently by amplifying a calmodulin gene at the untargeted locus, and concomitantly revert to normal growth rate. This response could be related to the property of the trypanosome of maintaining most housekeeping genes in tandem chromosomal arrays. Recombination appears to be initiated by a crossover event between the linearized end of the transfecting plasmid and a homologous region in the host genome; the second crossover generally occurs internally and in that region requires no more than 87 bp of homology.

**Key Words:** Homologous recombination; calmodulin genes; *Trypanosoma brucei*; gene amplification

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Our laboratory and others have reported that stable transformation of *Trypanosoma brucei* and related organisms appears to occur by integration of the exogenous DNA into the homologous chromosomal locus (Cruz and Beverly 1990; Lee and Van der Ploeg 1990; Ten Ashroek et al. 1990; Eid and Sollner-Webb 1991). The other laboratories studying *T. brucei* used plasmids singly cut within a tubulin gene region; five resultant stably transformed cell lines were analyzed, and these had up to 16 copies of the entire plasmid integrated into the endogenous tubulin locus [Lee and Van der Ploeg 1990; Ten Ashroek et al. 1990]. Our system [Eid and Sollner-Webb 1991] utilized an excised calmodulin–neomycin-resistance fusion gene (CNeoC), and seven resultant stable cell lines all showed targeting of a single copy of the transfecting DNA to the calmodulin locus. In this paper we demonstrate that this targeting event involves homologous recombination and is accurate to the nucleotide. Characterization of the 5’- and 3’-recombination products from 33 transfected cell lines showed that the CNeoC sequences targeted to the genomic calmodulin locus in all cases. By using doubly digested plasmids, recombination events were found to occur both in large terminal regions of calmodulin locus homology and in any of the four tandem genomic calmodulin gene repeats. This results in deletion of one, two, three, or all four of the calmodulin genes in the targeted locus. Plasmids linearized within a region of homology also yielded transformants in which the entire plasmid had inserted into the calmodulin locus without accompanying loss of chromosomal sequences. Interestingly, resultant cell lines that contain only the four intact calmodulin genes of the unrecombined chromosomal locus exhibited impaired growth. Over time, these cell lines reverted to normal growth rate and, in concert, acquired an additional calmodulin gene copy, frequently by duplicating one of their remaining calmodulin genes.

**Results**

**Homologous integration of CNeoC involves recombination with various segments of the tandem calmodulin locus**

We have reported stable transformation of the procyclic form of *T. brucei* using the chimeric CNeoC DNA [Fig. 1A] cleaved with *XbaI* and *NotI* to remove vector sequences [Eid and Sollner-Webb 1991]. This construct contains the neomycin resistance (neo')-coding region inserted, in-frame, within a complete *T. brucei* calmod-
would occur within the 6 kb of 3' flanking region. However, the calmodulin-coding sequence is repeated four times at both chromosomal calmodulin loci in the recipient *T. brucei* strain (a, b1, b2, and c genes), and most of this sequence is located just downstream of the neo' sequences in CNeoC [Fig. 1A], so the downstream recombination event could also involve these homologous gene segments. Thus, we wanted to analyze the products of the targeted integration of CNeoC DNA into the trypanosome genome to determine in which regions of the calmodulin locus the downstream recombination occurred, to confirm that homologous recombination was responsible for the gene targeting, and to ascertain whether gene insertions in addition to gene replacements occurred when the transfecting DNA was singly cleaved within a region of homology.

Trypanosomes were transfected by using CNeoC DNA cleaved with *XbaI* within the 5' flanking calmodulin region, with *NotI* at the end of the 3' flanking calmodulin region, or with both *XbaI* and *NotI* (see Fig. 1A). Thirty-three resultant cell lines, derived from two to three separate electroporation experiments for each differently cleaved DNA, were selected by growth in the presence of G418 and were cloned by dilution. Their genomic DNA was restricted and analyzed by using probes for the neomycin-resistance (*neo'*, calmodulin [cal], and pBR322 vector [pBR] regions, sequentially (diagramed at the bottom of Fig. 1). In all cases, the CNeoC sequences targeted to the genomic calmodulin cluster. However, the structure of the resultant cellular DNA was not identical but fell into six distinct classes, events A–F (diagramed in Fig. 1B; the cell lines constituting each event are listed in Table 1). Data for representative members of each are presented in Figures 2 and 5 (below) as indicated: *ClaI*-cut DNA, *neo'* probe [Fig. 2A], *ClaI*-cut DNA, *cal* probe [Fig. 2B], *EcoRI*-cut DNA, *neo* probe [Fig. 2C], *ClaI* and *SalI*-cut DNA, *cal* probe [Fig. 5], additional analyses (not shown).

Initial evidence that all transformants were not identical came from Southern analysis of genomic DNA digested with *ClaI*, which cleaves at the 3' border of the *neo'* sequence in the chimeric CNeoC construct [Fig. 1A]. When probed for *neo'*, sequences, most cell lines yielded a 12-kb fragment (events A–D; Fig. 2A; Table 1), indicating that the upstream recombination occurred within the 7.2 kb of 5' homology between CNeoC plasmid and genomic DNA (Fig. 1A; for details, see Eid and Sollner-Webb 1991). However, the *ClaI*-cleaved DNA from a few lines yielded *neo'*-containing fragments of two other sizes [Fig. 2A], either 14.5 kb [event E] or ~18 kb plus 12 kb [event F], as described below, whereas control untransformed cells showed no hybridizing material [Fig. 2A, lane Co]. Thus, transformation with CNeoC DNA yields at least three different recombination products.

### Cell lines of events A–D

When the filter of Figure 2A was stripped and reprobed by using the calmodulin-coding region (diagramed in Fig. 1A) the organization of the endogenous *T. brucei* calmodulin locus [Tschudi et al. 1985; Eid and Sollner-Webb 1991] is depicted in the top two lines. The solid box represents the calmodulin-coding regions. (The calmodulin gene repeat is 0.85 kb, and the 5'- and 3'-flanking *EcoRI* fragments are 2.4 and 12 kb, respectively.) (B) Events A–E, diagramed below, are the products of double crossovers. In event A, one crossover occurred in the 7.2 kb of 5' homology to the *neo'* sequence, and the other in the 6 kb of homology to the *neo'* sequence. In events B–D, the first crossover occurred in the 7.2 kb of 5'-flanking homology and the second in the 420 bp of homology with calmodulin genes b2, b1, and c, respectively. In event E, the first crossover occurred in the 6 kb of 3'-flanking homology and the second in the 87-bp homologous segment flanking the translation initiation site of the c gene. Event F represents homologous insertion of two entire copies of plasmid CNeoC in the 5' region of the calmodulin locus. Probes *neo'*, *cal*, and pBR are illustrated below. (C) *ClaI*, *EcoRI*, *SalI*, *XbaI*, *NotI*, *X* mutated site. Parentheses denote mutated sites.

### Figure 1. The products of recombination of plasmid CNeoC and the calmodulin locus. (A) The organization of the endogenous *T. brucei* calmodulin locus [Tschudi et al. 1985; Eid and Sollner-Webb 1991] is depicted in the top two lines. The solid box represents the calmodulin-coding regions. (The calmodulin gene repeat is 0.85 kb, and the 5'- and 3'-flanking *EcoRI* fragments are 2.4 and 12 kb, respectively.) (B) Events A–E, diagramed below, are the products of double crossovers. In event A, one crossover occurred in the 7.2 kb of 5' homology to the *neo'* sequence, and the other in the 6 kb of homology to the *neo'* sequence. In events B–D, the first crossover occurred in the 7.2 kb of 5'-flanking homology and the second in the 420 bp of homology with calmodulin genes b2, b1, and c, respectively. In event E, the first crossover occurred in the 6 kb of 3'-flanking homology and the second in the 87-bp homologous segment flanking the translation initiation site of the c gene. Event F represents homologous insertion of two entire copies of plasmid CNeoC in the 5' region of the calmodulin locus. Probes *neo'*, *cal*, and pBR are illustrated below. (C) *ClaI*, *EcoRI*, *SalI*, *XbaI*, *NotI*, *X* mutated site. Parentheses denote mutated sites.
Table 1. Cells lines corresponding to the various recombination events obtained with differently cleaved CNeoC DNA

| Event | Replaced calmodulin genes | Cell lines derived from transfected CNeoC DNA cleaved with |
|-------|---------------------------|----------------------------------------------------------|
|       |                           | XbaI + NotI | XbaI | NotI |
| A     | a b₁ b₂ c                 | 5, 6, 7, 9, 21 | R, Z |
| B     | a b₁ b₂                   |             |      |
| C     | a b₁                      | a, b, d, e, f | 8, 10 |      |
| D     | a                         | 17, 18, 20   |      |
| E     | c                         | m           | l, i, h, t, E |
| F     | insertion                 |             | 2, 11 |      |

All the cell lines of event A changed upon continued propagation. It is unlikely that event A was greatly under-represented in our original collection of clones as a result of growth rate selection, because the characterized secondary mutations occurred slowly, over a period of several months. Lines 21 and R acquired an additional calmodulin gene copy at the untargeted calmodulin locus.

1B), six different patterns were observed (Fig. 2B). All of the cell lines, like the control untransfected cells (Fig. 2B, lane Co), gave a 20-kb calmodulin-containing Clai fragment. Thus, this fragment represents the uncombined calmodulin locus. All of the transfected cells yielded additional calmodulin-containing fragments. Most lines showed one such fragment, which contains the DNA sequences extending downstream from the Clai site of the integrated CNeoC DNA (Fig. 1B). This fragment falls into four different length classes (Fig. 2B, events A–D). The smallest is 8 kb, the size expected when the downstream recombination event took place within the 6 kb of 3' homologous flanking DNA (event A). Accordingly, this Clai fragment comigrates with the analogous fragment excised from the untransfected cellular DNA by EcoRI and Clai digestion (see Fig. 1A, B; data not shown). Thus, in the cell lines of event A the four tandem genomic calmodulin-coding regions have been replaced by the CNeoC fusion gene.

The analogous fragments from the cell lines of events A, B, C, and D differ in size by units of ~0.85 kb (Fig. 2B). Because the tandem chromosomal calmodulin genes a, b₁, b₂, and c form a 0.85-kb repeat (Tschudi et al. 1985; see Fig. 1A), the successively larger fragments evidently represent cell lines retaining one, two, or three of the chromosomal calmodulin genes downstream of the CNeoC region (Fig. 1B, events B–D). These transformants presumably arose from a recombination event between the 420 bp of the common calmodulin-coding region present downstream of the neo' sequences in CNeoC and the chromosomal calmodulin b₂, b₁, or a genes (whose sequences are virtually identical; Tschudi et al. 1985). The recombinant calmodulin locus of events B, C, and D therefore retains, 3’ to the CNeoC gene, the intact genomic calmodulin gene c, genes b₂, and c₁ or genes b₁, b₂, and c, respectively (Fig. 1B). Because the calmodulin probe hybridizes to each of the calmodulin gene copies, the diagnostic bands should become increasingly intense going from events A to B to C to D, as is observed (Fig. 2B).

The organization of the recombined calmodulin locus of events A–D is supported further by the analogous mapping Clai- plus Sall-digested DNA (see Fig. 5, below), which yields smaller fragments that can be sized more accurately. Cell lines of event A yielded a 2.5-kb calmodulin-containing fragment that comigrates with the equivalent fragment from CNeoC plasmid DNA.

Figure 2. Analysis of the DNA organization of representative transfected cell lines. DNA from transformant cell lines of events A–F was cleaved with Clai and hybridized first with the neo' probe (A) and then with the cal probe (B) after complete stripping of the neo' probe. Lane Co represents untransfected cellular DNA. (C) Genomic DNA from lines representing the various events was cleaved with EcoRI and hybridized with the neo' probe. Lane M shows size markers of HindIII-cleaved λ DNA. Lane P contains plasmid CNeoC, cleaved with EcoRI. Sizes are shown in kilobases.
[data not shown], and the corresponding fragments from cell lines of events B, C, and D, indeed, show the 0.85-kb progressive increase in size. Thus, the cell lines of events A–D are as diagramed in Figure 1B.

Cell lines of events E and F

Lines representing recombination event E (lines 1 and t) showed calmodulin sequences on the same 8-ko $Clal$ fragment and the same 2.5-kb $Clal$–$SalI$ fragment as those of event A (Figs. 2B and 5A), indicating that their downstream recombination occurred within or downstream of the genomic calmodulin gene $c$ (illustrated in Fig. 1B). However, their upstream recombination was different from that discussed above. Lines of event E have a 14.5-kb $neo^o$-containing $Clal$ fragment, 2.5 kb larger than the analogous fragment from the other cell lines (Fig. 2A). This difference was confirmed by sizing $SalI$- and $Clal$-digested DNA (which yielded a 5.5-kb fragment from cell lines of event E and a 3-kb fragment from the other lines, not shown). The extra 2.5 kb represents three 0.85-kb calmodulin gene repeats, as shown in Figure 1B, event E. This organization would result if the upstream recombination was within the 87 bp surrounding the 5’ end of the CNeoC gene that is also present in the calmodulin $c$ gene (see Fig. 1B). As predicted, the cell lines of event E give a 1.85-kb $neo^o$-containing $EcoRI$ fragment (the 1-kb $neo^o$ region plus a 0.85-kb calmodulin repeat) rather than the 3.4-kb $EcoRI$ fragment (Fig. 2C) and 2.5- and 5.5-kb calmodulin-containing $Clal$ plus $SalI$ fragments (Fig. 2A). The $Clal$ fragments (see Fig. 5, below), this organization is confirmed in Figure 3 (see below).

Lines of event F have two copies of $neo^o$ sequences, yielding 12- and 18.2-kb $Clal$ fragments (Fig. 2A). The 12-kb fragment comigrated with that of events A–D (Fig. 2A, as did the analogous 3-kb $SalI$ plus $Clal$ fragment; data not shown), indicating that the upstream recombination also took place within the 7.2-kb region of homology upstream of the calmodulin-coding sequences.

The ~18-kb $neo^o$-containing fragment of event F comigrated with linearized full-length CNeoC plasmid DNA (data not shown), indicating that more than one copy of the entire CNeoC plasmid integrated into the cellular genome. Reprobing this $Clal$ digest for calmodulin sequences showed the ~18-kb plasmid-size fragment and a ~27-kb fragment, as well as the 20-kb fragment from the other calmodulin chromosomal locus (Fig. 2B, event F); the former two fragments were also detected upon reprobing for pBR322 vector sequences (data not shown). These data led to the structure for event F (Fig. 1B) in which two copies of the entire CNeoC plasmid homologously inserted upstream of the calmodulin locus. As expected, this event resulted only when the transfecting CNeoC DNA was cleaved once within a region of homology, using $XbaI$ (Fig. 1A).

Reprobing the filter of Figure 2, A and B, for pBR322 sequences showed that only cells of event F contained them. This provides further evidence that event F resulted from insertion of the whole transfecting plasmid but that the other cell lines arose from a double cross-over surrounding the $neo^o$ region, and therefore lack pBR322 sequences (Fig. 1A, B).

Sequence of a junction derived by homologous recombination

To confirm that the gene targeting of transfecting CNeoC DNA occurred by homologous recombination to the nucleotide, the junction region of cell line 1 (event E) was sequenced. We chose to analyze this junction because it occurs within an 87-bp region (see above), while all other recombination events can be localized only to within >420 bp regions (Fig. 1B). This recombination junction was polymerase chain reaction (PCR)-amplified by using a 5’ primer of calmodulin-coding region and a 3’ primer for $neo^o$ sequence (see diagram in Fig. 3). Dideoxy sequencing of the cloned junction showed that it begins in the $b_2/c$ intergenic calmodulin sequence, continues

Figure 3. Integration of the CNeoC gene by homologous recombination. The diagrams at the top depict the transfecting CNeoC gene (left), the $b_2$ and $c$ wild-type calmodulin gene region (center), and the region surrounding the integrated $neo^o$ gene of event E, cell line 1 (right). [Below] The sequence of the indicated region surrounding the 5’ end of the CNeoC gene (left), Eid and Sollner-Webb 1991), the wild-type calmodulin $c$ gene (center, Beck et al. 1982), and the $neo^o$ gene of cell line 1, event E (right). The dideoxy sequence analysis of this region of cell line 1 is shown at the bottom. The 87-bp region contained in all three of the sequences is indicated by underlining, it includes the first 57 bp upstream of the translation initiation codon (Tschudi et al. 1985) and the first 30 bp of the common coding region (Eid and Sollner-Webb 1991). Double underlining represents the ATG calmodulin translation initiation codon. The sequence of cell line 1, event E, is identical to that of the genomic calmodulin locus upstream of the common region and identical to that of the CNeoC gene downstream of the common region.
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into the 87-bp homologous sequence, and extends into
the neo sequence [Fig. 3], thereby verifying that cell line
1 results from a homologous recombination event [Figs.
1B and 3].

The number of functional calmodulin genes affects
the growth rate of T. brucei

While propagating the cell lines, those representing
event A (replacement of all four calmodulin genes of
the targeted locus) were consistently less dense than the
other cell lines and only reached a lower density
(\(1 \times 10^7\) vs. \(2 \times 10^7\) cells/ml). To measure growth
rate, log-phase cultures representing all of the recombi-
nation events, as well as control untransfected cells,
were diluted to equal concentration and grown in me-
dium lacking G418. From cell counts, growth rates were
calculated and compared with that of the untransfected
cells [Fig. 4]. Figure 4 represents at least three different
experiments of at least two different cell lines for each
event (except event B with only one existent cell line).
The lines of event A consistently grew at only 40–50%
the rate of untransfected cells. In contrast, those of
event B, with only one extra calmodulin gene, generally
grew like the untransfected cells. Thus, cell growth is
impaired in lines bearing four intact calmodulin genes
but is virtually wild type in lines bearing five or more
calmodulin genes.

The slow-growing cell lines phenotypically revert,
often by duplicating a calmodulin gene
in the untargeted chromosome

The transformed cell lines have been passaged continu-
ously for over 1 year, and their growth rates and DNA
organizations have been monitored. The growth rates of
the cell lines shown to be event A, 40–50% of the un-
transfected cells when initially analyzed [Fig. 4], began to
increase over subsequent months, until they eventually
became indistinguishable from the untransfected cells
[Fig. 4, column A* and data not shown]. The growth rates
of cell lines containing five or more intact calmodulin
genes remained constant.

Over the months, a corresponding change in the diges-
tion patterns of the DNA from the cell lines of event A
was also observed. DNA cleaved at early times with ClaI
and SalI, and analyzed with the calmodulin probe,
showed the same 7-kb fragment representing the un-
targeted calmodulin locus that is obtained from all of the
cell lines [Fig. 5C]. However, this 7-kb fragment began to
disappear from the DNA of cell line 21, when prepared 2
months later [Fig. 5B] and was virtually absent from
DNA prepared 3.5 months later [Fig. 5A]. Concomi-
tantly, a 7.8–7.9-kb fragment appeared [Fig. 5A, B; frag-
ments sized from semi-log plots of the data of Fig. 5].
Cell line R (another event A) also showed a transition
from the 7-kb to the ~7.85-kb fragment [Fig. 5A]. Thus,
the calmodulin locus of the untargeted chromosome mu-
tated, becoming larger by the size of one (0.85 kb) cal-
modulin gene repeat. Further probing of this new ~7.85-
kb fragment revealed that it contained neither neo' nor
pBR322 sequences [data not shown]. Proof that the ~0.85
kb increase in the size of this fragment was the result of
an increase from four to five copies of the calmodulin
gene repeat came from comparing EcoRI-cleaved DNA of
cell lines R and 21 with that of untransfected cells (re-
peating the experiment of Fig. 2C, but probing for the
entire region contained in the 7-kb genomic SalI frag-
ment). As expected, four fragments were observed (the
0.85-kb calmodulin repeat, the 12-kb 3'-flanking region,
and the 2.4- and 3.4-kb 5'-flanking regions from the two
chromosomes; Fig. 1; data not shown). If the 0.85-kb
insertion had resulted from any rearrangement other
than a duplicated calmodulin gene repeat, a new frag-
ment of a different size would have been observed.

Notably, the acquisition of the extra calmodulin gene
by the cell lines forming event A* occurred coincident
with their reversion to wild-type growth rate. Evidently,
their growth rate was increased by the duplication of one
of the four calmodulin gene copies on the chromosome
that had not received the neo' gene. The loss of the pa-
rental 7-kb fragment in event A* also confirms that the
calmodulin locus was diploid in the original T. brucei
cells.

In conjunction with their regaining wild-type growth
rate, the other cell lines of event A also changed their

![Figure 4. Correlation of growth rate with the number of intact calmodulin genes in the transformant cell lines. The growth rate of various transformants of events A–F, relative to that of control untransformed T. brucei grown in parallel, is plotted. The growth rates were consistent throughout the logarithmic growth period. Data for lines of event A were obtained prior to any evidence of altered genotype [i.e., on or prior to August 9, 1990 for line 21; see Fig. 5C]. A* represents the derivatives of event A, lines R and 21, after acquisition of an additional calmodulin gene copy on the nontargeted chromosome and reversion to normal growth rate (measured on or after November 23, 1990; see Fig. 5A)]. The other cell lines of event A also reverted to normal growth rate but through different genetic alterations.](image)
Figure 5. Analysis of DNA from transfected cell lines prepared after different time periods of propagation. Genomic DNA from the indicated cell lines was cleaved with *SalI* and *ClaI* and was detected with the *ca1* probe. The dates of DNA preparation are shown (month/date/year). The transfections took place in January, 1990; but because of the low cell density inherent in the cloning protocol, rapid growth did not commence for 4-5 months. (Lane Co) DNA of untransfected control *T. brucei*.

Figure 6. Analysis of RNA from transfected cell lines. Total cellular RNA was electrophoretically resolved and detected by using the *neo* probe. (Lane Co) RNA from control, untransfected *T. brucei*.

**Discussion**

**Homologous recombination events in *T. brucei***

We have studied transformation of *T. brucei* using a calmodulin–neomycin-resistance fusion gene (CNeoC) and found that all of the resultant neomycin-resistant cell lines arose from targeting of the exogenous DNA to the homologous calmodulin locus of the host chromosome [Fig. 1B; Table 1]. Restriction mapping indicated that homologous recombination was responsible for the gene targeting in all of the cell lines [Fig. 2], and sequencing the junction of one common transformant demonstrated that the recombination, indeed, was homologous to the nucleotide (Fig. 3).

Notably, the transformants show several different recombination patterns [Fig. 1B]. This occurs because the calmodulin gene, like most genes in *T. brucei* [Michels et al. 1986], is present in multiple tandem copies [Tschudi et al. 1985]; the recipient cell line used in our studies has four copies at both chromosomal loci [Eid and Sollner-Webb 1991]. Thus, the CNeoC DNA can combine with the genome in more than one way, either by using only the >7 kb and 6 kb of 5' and 3' calmodulin-flanking sequence present in the CNeoC plasmid or by also using the calmodulin gene sequence surrounding the *neo* sequence, which can recombine with any of the chromosomal calmodulin genes. Thus, double crossovers can result in the loss of one, two, three, or all four of the intact calmodulin genes from the recipient chromosomal locus [Fig. 1B]. In addition, CNeoC insertion can result from integration of the entire plasmid, without a corresponding loss of chromosomal sequences.

Transformants were obtained by using CNeoC DNA that was singly cleaved to generate large 5'- or 3'-flanking regions of homology or doubly cleaved to excise the CNeoC region. In all cases, the vast majority of the transformants were substitution type, presumably re-
resulting from double crossovers. In all 31 of these, at least one of the crossovers took place in the large homologous region abutting the linearized end of the transfecting DNA. Unexpectedly, however, the second crossover generally was not in the large homologous region on the other side of the neo gene (event A) but in the smaller homologous regions of calmodulin gene sequence (events B–E). All of the cell lines of events B, C, and D (with the terminal crossover in the 5'-flanking homology region) were obtained by using CNeoC DNA cleaved in the 5'-flanking region (with XbaI), whereas none were obtained by using DNA cleaved only in the 3'-flanking region; all cell lines of event E (with the terminal crossover in the 3'-flanking homology region) were obtained by using CNeoC DNA cleaved in the 3'-flanking region (event F; Fig. 1B), as shown by electrophoretic analyses performed 6 months or more after the transfection date (Fig. 2A,B, data not shown). However, analysis of DNA extracted a shorter time after the transfection indicated that both cell lines originated as single-insertion events which, over time, changed to double insertions (data not shown). This might be the result of subsequent unequal crossing-over, facilitated by the 13 kb of duplicated calmodulin sequence flanking the neo gene in the single insertions, and selection of these cells during continuous growth in G418.

When the transfecting DNA was singly cleaved within the 5'-region of homology (XbaI cleaved, Fig. 1A), plasmid insertions were also obtained (event F). As expected, such insertions were not similarly generated from CNeoC DNA cleaved at the end of a region of homology (NotI) or doubly cleaved (XbaI and NotI). Notably, however, plasmid insertions only account for 2 of the 12 transformants analyzed from XbaI-cleaved DNA (Table 1). This is unlike the situation in yeast and mammalian cells, where plasmids linearized within a region of homology yield primarily insertion-type transformants (Orr-Weaver et al. 1981; Thomas and Capecchi 1987).

Both cell lines representing the insertion event contain two complete copies of the transfecting plasmid (event F, Fig. 1B), as shown by electrophoretic analyses. This is in contrast to the yeast and mammalian systems, where one target is typically present in a single copy per haploid chromosome complement, in trypanosomatids genes generally occur in multiple tandem copies (Thomashow et al. 1983; Clayton 1985; Tschudi et al. 1985; Michels et al. 1986; Dragon et al. 1987). It has been speculated that these duplicated copies may be necessary to obtain sufficient quantities of essential RNAs from the gene expression machinery of trypanosomes. The process of gene expression in trypanosomes is still largely characterized, but evidently differs from most other eukaryotes in a number of ways, any of which could affect the efficiency of gene expression in these primitive organisms. For example, trypanosome RNAs appear to be initially transcribed in a monocistronic fashion from still unknown but distant promoters and then cleaved to monocistronic RNAs in a process involving the transsplicing of a common leader sequence as well as a polyadenylation directed by noncanonical sequences (Thomashow et al. 1983; Clayton 1985; Gonzalez et al. 1985; Tschudi et al. 1985; Michels et al. 1986; Murphy et al. 1986; Sutton and Boothroyd 1986; Dragon et al. 1987).

The calmodulin gene is a typical tandemly repeated T. brucei gene. It encodes the calcium-binding protein calmodulin, which in other organisms is required for numerous essential functions, such as cell-cycle progression (Chafoules et al. 1982), signal transduction (Braam and Davis 1990), microtubule assembly (Charles et al. 1983), cell motility, cyclic nucleotide metabolism, and protein phosphorylation (for review, see Wang and Weissman 1979). It is a unique gene in yeast (Davis et al. 1983; Clayton 1985; Tschudi et al. 1985; Michels et al. 1986; Dragon et al. 1987).
There is reason to suspect (Straus and Straus 1976) that the same cellular machinery is involved in this gene duplication and the original homologous recombination event that generated the gene deletion.

In conclusion, the ability to target T. brucei genes for recombination, to observe the effect of this on cellular function, and to select and study revertants of such changes should be of great value in further understanding the unusual molecular processes of this interesting organism. In particular, the observation that a growth phenotype results from reducing the number of calmodulin genes to one-half the normal chromosomal complement offers promise that analogous deletion of one of the two allelic sets of other genes will similarly show a phenotype, permitting a study of the effect of T. brucei genes without necessitating the mutation of both chromosomal loci.

Materials and methods

Cell lines

Proclive T. brucei cultures [strain TREU 667] were stably transformed by electroporation with the CNeoC plasmid, cleaved with Xbal and NotI (Eid and Sollner-Webb 1991) or with Xbal or NotI alone (see Fig. 1A). The 18-kb pBR322-based CNeoC plasmid contains 7.2 kb upstream of the T. brucei calmodulin gene cluster and the first 30 nucleotides beyond the translation start site of the calmodulin a gene, a 1-kb segment from pSV2neo containing the neomycin phosphotransferase gene (Beck et al. 1982) inserted in-frame, the last 420 bp of the coding region of the calmodulin c gene, and 5.6 kb of 3′-flanking sequence (Eid and Sollner-Webb 1991), see Fig. 1A). All transformants have been maintained for the past year in regular growth medium supplemented with 50 μg/ml of G418 (Eid and Sollner-Webb 1991). To measure growth rates, logarithmic phase cells were seeded at 2 x 10^6 cells/ml in regular medium without G418 and cell density was measured every 2 days (doubling time ~24 hr).

DNA and RNA analyses and hybridization probes

Preparation of genomic DNA and total cellular RNA and their analyses by agarose gel electrophoresis and Southern and Northern transfers, were described previously (Eid and Sollner-Webb 1991). The neo probe (Fig. 1B) is the 1.3-kb HindIII–Avai fragment of pSV2neo (Beck et al. 1982) containing the coding region of the neomycin-resistance gene. The calmodulin-coding region probe (cal, Fig. 1B) is the 450-bp EcoRI–HindIII fragment from the genomic calmodulin genes (also present in the CNeoC fusion gene), which contains the bulk of the calmodulin-coding region (Tschudi et al. 1985). The pBR probe (Fig. 1B) is the 2.3-kb EcoRI–PvuII fragment from pBR322, which corresponds to the vector sequences of the CNeoC plasmid (Eid and Sollner-Webb 1991).

Primers, PCR, and DNA sequencing

The region of the recombination junction of transformant line 1 (event E) was cloned following PCR amplification of total genomic DNA by standard methods (Saiki 1990). The 3′ primer (5′-CCGATTCGGCCCAAGCGCCGAGAACCCTGC-3′) is complementary to a 5′ region of the neo sequence (Beck et al. 1982), the 5′ primer (5′-CCGATTCGGCCCAAGCGCCGAGAACCCTGC-3′) corresponds to a 3′ portion of the calmodulin-coding region (Tschudi et al. 1985). This allows amplifica-
tion of the region between the bz gene and the neomycin gene (event E, Fig. 1B). The PCR product was cleaved at the EcoRI and BamHI sites present in the 5’ and 3’ primer sequences, respectively, and was cloned in EcoRI-BamHI digested M13 phage for dyeoxy sequencing (Sambrook et al. 1989).

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