Epigenetic self-regulation of developmental excision of an internal eliminated sequence in *Paramecium tetraurelia*

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Differentiation of the somatic macronucleus of ciliates after sexual events involves the programmed excision of thousands of single-copy internal eliminated sequences (IESs) from the germ-line genome. We have studied two cell lines of *Paramecium tetraurelia* that have identical germ-line genomes but differ in their macronuclear genomes. In the IES- cell line, a 222-bp IES interrupting a coding sequence is reproducibly excised during macronuclear differentiation, whereas it is not in the IES+ cell line. In a cross between the two lines, the developmental alternative is maternally inherited, suggesting that it is epigenetically controlled by the old (prezygotic) macronucleus in each cell. Transformation of the macronucleus of both lines with plasmids carrying fragments of either version of the gene shows that the presence of the IES sequence in the old macronucleus results in retention of the IES in the new macronuclear genome of sexual progeny. This could be attributable to (1) inhibition of excision, or (2) repair of a double-strand gap left in the genomic sequence after constitutive excision of the IES, by a polymerization mechanism using a homologous IES+ template from the old macronucleus. The latter possibility is ruled out by experiments showing that modified IESs can inhibit excision without being copied in the new macronuclear genome. Possible mechanisms are discussed in the light of a quantitative analysis of excision inhibition by the maternal IES sequence.

[Key Words: Genomic rearrangements; IES excision; maternal inheritance; macronuclear differentiation; ciliates]

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Alternative DNA rearrangements have been shown to play a role in the developmental regulation of gene expression in a broad range of organisms but only in isolated cases, such as sporulation in *Bacillus subtilis*, mating-type switching in yeast, or the generation of immunoglobulin diversity in vertebrates. In no other group of organisms have programmed genomic rearrangements become such common practice as in the ciliates. The unique nuclear dimorphism characterizing these unicellular eukaryotes allows them to keep copies of the germ-line genome in the transcriptionally silent micronuclei, for use in sexual events (meiosis and fertilization), while expressing their genes from the extensively rearranged version of the genome found in the macronuclei. Macronuclei are highly polyploid somatic nuclei that divide amitotically and are destroyed at the end of the vegetative phase of the life cycle, during sexual events; after fertilization, new macronuclei and micronuclei differentiate from mitotic products of the zygotic nucleus. In addition to the amplification of the genome, the differentiation of a macronucleus from a diploid nucleus involves two types of reproducible rearrangements: the fragmentation of germ-line chromosomes in specific regions, followed by the addition of new telomeres, and the elimination of numerous internal sequences, either single-copy or repetitive, from most genes [for review, see Prescott 1994]. This complete developmental remodeling of the genome could be used by ciliates to produce functionally diverse vegetative clones from a given germ-line genotype and maintain useful polymorphisms. This would require the capacity to regulate specifically individual rearrangements.

The study of *Paramecium* strains with entirely homozygous germ-line genomes has revealed that an epigenetic, sequence-specific mechanism is involved in the developmental choice of alternative telomere addition regions in at least three events of germ-line chromosome fragmentation. These alternative rearrangements can lead to the loss from the somatic genome of genes normally located near macronuclear telomeres [Meyer 1992; Kim et al. 1994; Scott et al. 1994a] or to the modulation of their expression [Keller et al. 1992]. Specific regulation may also occur for the other type of developmental ge-
some fragmentation patterns, the excision and nonexcision characters follow a maternal pattern of inheritance in crosses between the IES+ and IES− lines, confirming that the developmental alternative is controlled by the old macronucleus in each cell and does not depend on any germ-line difference. Transformation of the vegetative macronucleus of both lines by plasmids containing fragments of the G gene shows that during nuclear reorganization, the presence of the IES sequence itself in the old macronucleus inhibits its own excision in the developing macronucleus.

**Results**

**Maternal inheritance of the IES+ and IES− characters at conjugation**

To study the inheritance of the IES+ and IES− developmental alternatives during conjugation, an IES+ clone of mating type E was crossed with an IES− clone of mating type O. After the pairing of the cells, the micronuclei undergo meiosis. An additional mitosis of the single surviving meiotic product yields two identical haploid gametic nuclei in each cell. The two paired cells then exchange one of these nuclei; karyogamy thus results in the formation of genetically identical F1 zygotic nuclei in the two mates (Sonneborn 1974). If the IES+ and IES− characters were determined genetically in the zygotic nucleus, the phenotype of the F1 macronucleus would be the same in the progeny of both mates.

Three pairs were analyzed. After their separation, the two exconjugants from each pair were isolated. Each exconjugant produces two F1 caryonides (after two divisions of the zygotic nucleus, two of the products differentiate into new macronuclei, which segregate without division to the two daughter cells); one of the two from each exconjugant was cultivated. The maternal inheritance of mating types in *P. tetraurelia* provides an independent means to identify the parental origin of the F1 caryonides (see Materials and methods). Figure 1 shows a

![Figure 1](https://genesdev.cshlp.org/)

Figure 1. Southern blot analysis of a cross between IES+ and IES− cell lines. Total DNA was digested with *PstI*, and the blot was hybridized with probe b (see Fig. 2). (Lane 1) The IES+ parent; (lane 2) the IES− parent. (Lanes 3, 5, 7) F1 caryonides derived from the IES+ parents of three different pairs; (lanes 4, 6, 8) F1 caryonides derived from the IES− parents of the same pairs.
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Southern blot of PstI-digested DNA from the IES+ and IES− parents of the cross (lanes 1 and 2, respectively) and from the F1 caryonides (lanes 3–8). In the IES+ macronuclear version of the G gene, the IES is contained in a 1.2-kb PstI fragment (see map in Fig. 2); the corresponding fragment of the IES− version is only 1 kb long. Only macronuclear DNA can be detected on such a blot, because it is ~250 times more abundant than micronuclear DNA. Hybridization of the blot with a probe specific for this fragment (see Fig. 2, probe b) shows that the three F1 caryonides derived from the IES+ parents are IES+ themselves (lanes 3,5,7), whereas the three F1 caryonides derived from the IES− parents are IES− (lanes 4,6,8). A small amount of the IES− form is detectable in the IES+ caryonide 5, as is sometimes observed in individual IES+ caryonides obtained through autogamy. Thus, although conjugation yields identical germ-line genomes in all F1 caryonides, in each one developmental processing of this genome produces the same macronuclear G-gene structure as in its cytoplasmic parent. The alternative processing therefore cannot be determined genetically in the zygotic nucleus. Rather, it appears to be controlled epigenetically by the old macronucleus in each cell.

**Transformation of the IES+ macronucleus with an IES− plasmid does not restore developmental excision after autogamy**

The maternal inheritance of the IES+ and IES− macronuclear versions may indicate that the mechanism of developmental excision is inhibited by the IES+ old macronucleus or that it requires positive activation by the IES− old macronucleus. Alternatively, it is possible that excision occurs constitutively during development in both lines, leaving double-strand breaks in the G-gene coding sequence, at both junctions with the IES. Repair of the genomic sequence by polymerization, using a homologous template originating from the old macronucleus, would result in the copying of the maternal G-gene version. This “cut-and-repair” model is similar to the mechanisms involved in the repair of genomic sequences after excision of the P element in *Drosophila* (Engels et al. 1990) and the Tcl transposon in *Caenorhabditis* (Plasterk 1991). It could account both for precise excision in IES− cells, and for the apparent lack of excision of one particular IES in genetically wild-type IES+ cells, which presumably contain all trans-acting factors necessary for IES excision. Although the transfer of sequences from the old macronucleus to the developing one has not been demonstrated, such a process has been postulated previously to account for other effects of transforming plasmids on developmental rearrangements (see Discussion). To test this hypothesis, we first asked whether transformation of the vegetative macronucleus of IES+ cells by a plasmid containing a portion of the IES− macronuclear version of the G gene would promote excision in the postautogamous progeny of transformed clones. The plasmid, p629, contains a 629-bp PstI–SpeI fragment located in the central part of the ~8-kb coding sequence (Fig. 2). An Asp718 site located 150 bp downstream from the PstI site is created by the excision of the 222-bp IES (Fig. 2). Plasmid p851, which

**Figure 2.** Restriction maps of the IES− and IES+ genomic sequences, and of the derived IES− and IES+ plasmids. The positions of probes a and b are indicated at top. The IES is shown as a shaded box. In the plasmid maps, the thin line represents the pUC18 vector sequence. The restriction site abbreviations are (A) Asp718; (P) PstI; (Pvu) PvuII; (S) SalI; (Sp) SpeI.
contains the 851-bp IES+ version of the same fragment, was used as a control. To minimize the nonhomologous recombination observed between monomers after microinjection of circular molecules into the macronucleus, the plasmids were linearized with a restriction enzyme cutting once in the pUC18 vector (when linear molecules are microinjected into the macronucleus, telomeric repeats are added to the extremities of monomers or end-to-end multimers; the resulting minichromosomes replicate autonomously during vegetative growth). Injected cells were grown individually for eight to nine fissions; four clones containing various amounts of each plasmid, as well as two uninjected controls, were then selected by a quick dot-blot procedure (see Materials and methods) and cultivated further for DNA extraction and Southern blot analysis. Figure 3A shows a BgIII digest of these samples, hybridized with probe b. The BgIII fragment containing the endogenous IES+ macronuclear G gene is seen as a thin band near the top of each lane. Neither plasmid contains any BgIII site; free-replicating monomers show up as somewhat smeared bands migrating at 3.5 kb [p629, lanes 1–4] or 3.7 kb [p851, lanes 5–8]. Dimers give a band at ~7 kb. Plasmid copy numbers ranged from <1 per haploid genome [undetectable in clones 1 and 5; cf. with uninjected controls, lanes 9 and 10] to >100 copies per haploid genome in clones 4 and 8 (i.e., 100,000 copies per macronucleus; see the quantitative analysis section). Lane 11 is an IES− wild-type control.

At the time of DNA extraction, ~1000 cells from each transformed clone were saved and starved to induce autogamy. In this self-fertilization process, the two identical gametic nuclei produced in each cell fuse together, yielding an entirely homozygous zygotic nucleus. Autogamy was checked by staining 100 cells from each clone to verify the fragmentation of the old macronucleus. The remainder of the starved cells was then transferred to rich medium and grown for DNA extraction. Figure 3B shows a Southern blot of a PstI digest of the postautogamous populations (each containing several hundred independent caryonidal clones), hybridized with probe b. Sample numbering is the same as in Figure 3A. In all samples except the IES− control (lane 11'), the 1.2-kb band shows that the IES was still maintained in the new macronuclear genome. The higher molecular weight fragments seen in some samples [lanes 3', 4', 8'] are PstI fragments from the plasmid monomers and dimers, attributable to the fact that a small fraction of cells had not undergone autogamy in these clones. The much smaller ratio of plasmid fragments to endogenous genomic fragment, compared with that of the preautogamous samples [Fig. 3A], is consistent with a fraction of nonautogamous cells <1%. Contaminating plasmid fragments were not always present after mass autogamies [lanes 2', 6', 7'], and were never observed when individual postautogamous caryonides from the same transformed clones were analyzed (not shown). Thus, transformation of IES+ cells with varying amounts of IES− plasmid p629 does not promote excision of the IES in the new macronucleus developing after autogamy.

Figure 3. Effects of transformation of IES+ cells with plasmids p629 and p851. (A) Southern blot of BgIII-digested DNA from IES+ clones transformed with p629 (lanes 1–4) or p851 (lanes 5–8), or uninjected controls (lanes 9, 10). Lane 11 is an IES− control. The blot was hybridized with probe b. The thin band near the top of each lane is the high molecular weight fragment containing the endogenous IES+ G gene. The sizes of plasmid monomers and dimers are indicated. (B) Southern blot of PstI-digested DNA from cultures obtained after autogamy of the transformed clones and uninjected controls, hybridized with probe b. The sizes of the IES+ (1.2 kb) and IES− (1 kb) G-gene fragments are indicated. Higher molecular weight fragments in lanes 3', 4', and 8' are contaminating plasmid fragments (see text). (C) Southern blot of PstI-digested DNA from cultures obtained after a second autogamy, hybridized with probe b.

**Decreased endogenous G-gene copy number after autogamy of clones transformed with G-gene plasmids**

In Figure 3B, the amount of the 1.2-kb genomic fragment appears to be inversely correlated to the amount of plasmid in preautogamous clones, although an approximately equal quantity of DNA was loaded onto each lane. (A quantitative analysis is presented in a later sec-
tion; note that in lanes 4' and 8', the intensity of the 1.2-kb band significantly overestimates the amount of the genomic G-gene fragment, because of the presence of multiple smeary bands arising from PstI digestion of the contaminating plasmid monomers and dimers.) The presence of high copy numbers of both p629 and p851 in the old macronucleus thus results in a reduced copy number of the endogenous homologous genomic region after autogamy. This global reduction, observed in post-autogamous cultures containing many different caryonid clones, does not represent an averaging of clones totally depleted in this genomic region and clones containing a normal copy number: Cultivation and analysis of individual caryonides from the autogamous cell populations showed that each of them presents a similar copy number reduction (not shown). No copy number reduction was observed for the unlinked C gene in the same samples (not shown). This is reminiscent of the effect of circular plasmids microinjected into the macronucleus of Paramecium primaurelia: Following autogamy of clones transformed with high plasmid copy numbers, specific deletions of the genomic sequences homologous to the plasmid inserts have consistently been observed in the new macronuclear genome (Meyer 1992; E. Meyer, A. Butler, and F. Caron, in prep.). In the present case, the size of the genomic region showing a decreased copy number is unknown but appears to be much larger than the plasmid inserts. Hybridization of Southern blots with different probes shows that it extends at least 6 kb upstream and 5 kb downstream of the IES; no discrete telomeric fragments or internal deletions were detected within this region (not shown).

**Excision can be restored in IES+ cell lines**

To study possible consequences of the reduction in IES+ G-gene copy number at nuclear reorganization, ~1000 cells from each postautogamous culture, which had been saved before DNA extraction, were again starved to induce autogamy. The results of a Southern blot analysis of the macronuclear G gene after this second mass autogamy is shown in Figure 3C. The 1.2-kb PstI fragment representing the IES+ form is revealed with probe b in all samples except the IES− control (lane 11'). However, the 1-kb fragment indicative of IES excision is now also observed in samples 2', 3', 4', 7', and 8', which derive from the parental cells with the smallest endogenous G-gene copy number (Fig. 3B). All of the 1-kb fragment in these samples could be recut by Asp718, showing that excision of the G-gene IES had recreated the wild-type junction (not shown). Thus, precise excision does not require the presence of the IES− G gene in the old macronucleus. The total G-gene copy number, as estimated by the sum of both bands, was rather homogeneous. In contrast, the fraction of excised molecules appeared to be inversely correlated with the IES+ G-gene copy number in the parental macronucleus. This suggests that developmental excision of the IES in the developing macronucleus is quantitatively inhibited, whatever the mechanism, by IES+ G-gene copies in the old macronucleus.

**IES+ plasmid p851 can transform the IES− cell line into a permanent IES+ line**

If this interpretation is correct, transformation of IES− cells with the IES+ sequence may result in retention of the IES in postautogamous macronuclei. To test this possibility, plasmids p629 and p851 were microinjected into the vegetative macronucleus of cells of wild-type strain d4.2. Transformed cell lines were selected and cultivated as described previously. After DNA extraction, samples were digested with PvuII, which does not cut within the plasmid inserts, but cuts the pUC18 sequence 110 bp to the left and 200 bp to the right of the inserts, yielding a 0.9-kb fragment from the IES− plasmid p629 and a 1.1-kb fragment from the IES+ plasmid p851 (Fig. 2). Figure 4A shows a Southern blot of PvuII-digested DNA from two uninjected control clones (lanes b, c), one clone transformed with p629 (lane d), and three clones transformed with p851 (lanes e–g). Lane a is an IES+ control. Plasmid copy numbers were estimated to be ~2 copies per haploid genome for p629 (clone d), and between 1 (clone e) and 10 (clone g) for p851 (see quantitative analysis). In addition to the plasmid fragments, hybridization with probe b revealed the IES− endogenous G gene as a 1.6-kb fragment in all samples, except the IES+ control (lane a), where the corresponding IES+ fragment is 1.8-kb long. A 1.8-kb fragment is also visible in lane g, it is unlikely that it arises through homologous recombination between the IES+ sequence of plasmid p851 and the endogenous G gene, because homologous recombination of plasmid DNA microinjected into the vegetative macronucleus of Paramecium has never been observed. This fragment probably originates from a fraction of cells in the preautogamous culture that had already undergone autogamy at the time of DNA extraction (see below).

Transformed clones were allowed to undergo a mass autogamy as in the previous experiment. A Southern blot of PstI-digested DNA from postautogamous cultures, hybridized with probe b, is presented in Figure 4B. The IES+ (lane a') and IES− (lanes b' and c') controls showed the same pattern as their respective parents; so did the postautogamous culture from the p629-transformed clone, in which the IES was excised from all macronuclear copies (lane d'). In all postautogamous cultures derived from the p851-transformed clones, however, the IES+ version was observed (lanes e'–g'). The fraction of G-gene copies retaining the IES in the new macronuclei appeared to correlate with the amounts of p851 plasmid in the parental macronuclei, reaching close to 100% in the g' culture. Thus, the presence in the macronucleus of an 851-bp fragment of the G gene containing the IES results in the maintenance of the IES in the macronuclear genome of the following sexual generation. The IES− version of the same fragment does not appear to have the same effect: The p629-transformed clone d had about twice as many plasmid copies as the p851-transformed clone e, but the d’ postautogamous culture did not show any detectable IES+ molecule, whereas e’ clearly did. The g’ sample also shows a decrease in the total G-gene copy number, similar to that
Figure 4. Effects of transformation of IES− cells with plasmids p629 and p851. (A) Southern blot of PvuII-digested DNA from an IES+ control (lane a), uninjected IES− controls (lanes b,c), and IES− clones transformed with p629 (lane d) or p851 (lanes e−g). The blot was hybridized with probe b. The sizes of the IES+ (1.8 kb) and IES− (1.6 kb) endogenous G-gene fragments are indicated. The 0.9- and 1.1-kb fragments arise from plasmids p629 and p851, respectively. (B) Southern blot of PstI-digested DNA from cultures obtained after autogamy of the transformed clones and uninjected controls, hybridized with probe b. The sizes of the IES+ (1.2 kb) and IES− (1 kb) G-gene fragments are indicated. (C) Southern blot of PstI-digested DNA from cultures obtained after a second autogamy, hybridized with probe b. The blot was hybridized with probe b. The sizes of the IES+ (1.2 kb) and IES− (1 kb) G-gene fragments are indicated. The 0.9- and 1.1-kb fragments arise from plasmids p629 and p851, respectively. [B] Southern blot of PstI-digested DNA from cultures obtained after autogamy of the transformed clones and uninjected controls, hybridized with probe b. The sizes of the IES+ (1.2 kb) and IES− (1 kb) G-gene fragments are indicated. (C) Southern blot of PstI-digested DNA from cultures obtained after a second autogamy, hybridized with probe b.

observed after autogamy of IES+ clones transformed with a high plasmid copy number (Fig. 3).

Some of the postautogamous cultures were allowed to undergo a second mass autogamy. Figure 4C shows a similar Southern blot analysis of the resulting cultures. The IES+ [lane a′′] and IES− [lane b′′] controls presented the same patterns as their respective parents. Cultu...
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Figure 5. Effects of transformation of IES− cells with plasmids p851S, pIES, and p851D. (A) Southern blot of PvuII-digested DNA from an un.injected IES− control (lane A), an IES+ control (lane B), and IES− clones transformed with p851S (lanes S1−S3), pIES (lanes I1−I8), and p851D (lanes D1−D4). The blot was hybridized with probe b. The sizes of the IES+ (1.8 kb) and IES− (1.6 kb) endogenous G-gene fragments are indicated. The 1.1- and 1.0-kb fragments arise from plasmids p851S and p851D, respectively. (B) The same blot as in A was stripped and rehybridized with pUC18 probe a, which allows a direct comparison of the amounts of different plasmids. The 0.5-kb fragment arises from plasmid pIES. (C) Southern blot of PstI-digested DNA from cultures obtained after autogamy of the transformed clones and uninjected controls, hybridized with probe b. The sizes of the IES+ (1.2 kb) and IES− (1 kb) G-gene fragments are indicated.

formed clones [Fig. 5B]. Copy numbers ranged from <1 to ~330 copies per haploid genome (in S3).

As in previous experiments, transformed clones were allowed to undergo a mass autogamy. Figure 5C shows a Southern blot of PstI-digested DNA from the postautogamous cultures, hybridized with probe b. In samples S1', S2', and S3', deriving from p851S-transformed clones, the 1.2-kb fragment indicates that the new macronuclear genome contained the IES. As with p851, the fraction of IES− molecules in the new macronucleus decreases with increasing plasmid copy number in the old macronucleus. A Southern blot of SalI-digested DNA from these samples showed that the IES+ form does not contain a SalI site (not shown). The IES in the new macronucleus therefore originates from the germ-line genome rather than from the plasmid of the old macronucleus. Samples derived from pIES-transformed clones (lanes I1'−I8') also show the 1.2-kb IES+ fragment. Thus, physical linkage of the IES sequence to the adjacent G-gene sequences is not required for the effect. Again, the fraction of IES+ copies in the new macronucleus correlates with the plasmid copy number in preautogamous clones; however, for an equal copy number, plasmid pIES seems to be much less efficient than plasmid p851S (e.g., cultures S2' and I8' have a similar IES+ /IES− ratio, although clone I8 contained ~17 times as many plasmid copies as clone S2, see quantitative analysis). In contrast, the 1.2-kb fragment was never observed in postautogamous cultures derived from p851D-transformed clones, even when the plasmid copy number was very high (up to 100 copies per haploid genome). The deletion of a 147-bp fragment internal to the IES thus abolishes the effect. Finally, it should be noted that for all three plasmids, and as already described for p851 and p629, the total G-gene copy number appeared to be smaller in postautogamous cultures derived from clones with the highest plasmid copy numbers.

Quantitative analysis of plasmid-induced reduction of G-gene copy number

A quantitative relationship was observed in the autogamy of all transformed clones, whether IES+ or IES−, between high G-gene plasmid copy number in the old macronucleus and reduced endogenous G-gene copy number in the new macronucleus. These variables were quantified for each clone by measuring the intensity of the corresponding bands on the Southern blots; to correct for differences in the amount of DNA loaded in each lane, the same blots were reprobed with a synthetic oligonucleotide specific for macronuclear telomeric repeats (see Materials and methods). The relative endogenous
G-gene copy number after autogamy (the sum of IES+ and IES− forms when both occur) was then plotted as a function of the relative plasmid copy number in preautogamous clones. Excluded from the analysis were samples 4/4′ and 8/8′, because a significant amount of contaminating plasmid fragments in 4′ and 8′ precluded an accurate determination of the G-gene copy number in the new macronucleus (Fig. 3B). The plot is shown in Figure 6. Relative copy numbers on both axes are expressed as number of copies per haploid genome, the reference value of one copy per haploid genome (∼1000 copies per macronucleus) being arbitrarily fixed as the average relative G-gene copy number in four postautogamous cultures derived from uninjected controls, which showed a 9% variation between the two extreme values. In spite of this variability, which may be ascribed to measurement errors or natural variation of the G-gene copy number or amount of telomeric repeats, a clear correlation appears: the endogenous G-gene copy number after autogamy decreases regularly with increasing plasmid copy numbers in preautogamous clones, in a seemingly logarithmic way. The most outlying point (indicated by an arrow) corresponds to sample g/g′ (Fig. 4); the plasmid copy number in clone g is probably underestimated because a fraction of the cells in this preautogamous culture had already undergone autogamy at the time of DNA extraction. It can be noticed that the postautogamous progeny of clones transformed with a small plasmid copy number (less than six to seven copies) appear to have a G-gene copy number larger than one, that is, larger than that of the postautogamous progeny of uninjected controls; the reason for this slight bias is unclear. Overall, the effects of different plasmids are not significantly different. A logarithmic function fitted among all points indicates that a 50% reduction in G-gene copy number is obtained on average with ∼70 plasmid copies per haploid genome in the old macronucleus.

Quantitative analysis of the induction of IES retention by the IES sequence in the old macronucleus

The experiments presented here have shown that transformation of the macronucleus of IES− cells with plasmids bearing the IES sequence simulates the behavior of the IES+ cell line, in which the IES is maintained in the genomic sequence during macronuclear differentiation. The effect is strongly dependent on plasmid copy number. To quantify the difference noted in the efficiency of different plasmids, the proportion of IES+ and IES− molecules was determined for all postautogamous cultures by measuring the relative intensities of the 1.0- and 1.2-kb PstI fragments revealed with probe b on the Southern blots. In Figure 7, the percentage of excised macronuclear G-gene copies is plotted as a function of the relative plasmid copy number in preautogamous clones, using the same reference value (one copy per haploid genome) as in Figure 6. p851D serves as a negative control: This plasmid had no effect on excision, even at very high copy numbers (up to 100 copies per haploid genome, i.e., ∼100,000 copies per macronucleus). In contrast, about five copies of the p851 plasmid per haploid genome resulted in only 11% of IES− G-gene copies in the new macronucleus. Plasmid p851S, which only differs from p851 by the presence of an additional SalI site, appears to have the same efficiency. As noted above, plasmid pIES was much less efficient: With ∼60 copies per haploid genome, 44% of the G-gene copies in the new macronucleus were still IES−.

Figure 6. Quantitative analysis of plasmid-induced reduction of G-gene copy number. The relative G-gene copy number in the new macronucleus is plotted as a function of the plasmid copy number in preautogamous clones, on a logarithmic scale. Both variables are expressed as number of copies per haploid genome, with the 1.0 reference value being the average relative G-gene copy number in postautogamous cultures derived from uninjected controls (see text and Materials and methods). The arrow points to the g/g′ sample; the plasmid copy number is probably underestimated in g (see text). Symbols for plasmids and recipient cells: (▲) p629/IES+; (■) p851/IES+; (○) p851/IES−; (●) p850S/IES−; (△) pIES/IES−; (□) p850D/IES−.
The complete absence of IES− G-gene in the uninjected IES+ cell line indicates that the effect of the endogenous IES+ genomic sequence in the old macronucleus must be stronger than that of IES+ plasmid p851, because one copy per haploid genome should be sufficient to maintain the IES on 100% of copies in the new macronucleus. The effect of subgenomic copy numbers of the chromosomal IES sequence can be studied quantitatively, using the cultures with various proportions of IES+ and IES− copies [lanes e− g’ in Fig. 4B], which were obtained by autogamy of p851-transformed IES− clones, and those with a reduced IES + copy number but no IES− copies [Fig. 3B samples], which were obtained by autogamy of transformed IES+ clones. The copy number of the IES + G gene in these cultures was determined by measuring the intensity of the 1.2-kb PstI fragment revealed with probe b and normalizing with the telomeric signal. After these cultures had undergone an additional autogamy, the percentage of excision in the new macronucleus [Fig. 4C, lanes e− g’; Fig. 3C samples] was measured as the relative intensities of the 1.0- and 1.2-kb PstI fragments and plotted as a function of the IES + G-gene copy number in the old macronucleus [expressed as a number of copies per haploid genome, using the same reference value]. Thus, in Figure 7, the effect of various copy numbers of the chromosomal IES can be compared directly to the effect of various copy numbers of the IES sequence carried by plasmids p851 and pIES. Samples 4'/4'' and 8'/8'' were again excluded from the analysis, for the reason mentioned above. Here, again, the proportion of IES− copies in the new macronucleus decreases with increasing IES+ copy numbers in the old macronucleus. The fact that the genomic plot is not monotonously decreasing is probably attributable to a slight overestimation of the IES+ G-gene copy number in the preautogamous sample 3’ (indicated by an arrow in Fig. 7), which contained a small amount of contaminating plasmid fragments [Fig. 3B]. As expected, the decrease is more rapid than with any of the plasmids. With 0.4 copy per haploid genome of the IES+ G gene in the old macronucleus, only ~6% of G-gene copies are excised in the new macronucleus.

**Discussion**

**Epigenetic regulation of developmental processing of the G-gene IES**

IES+ cell lines were initially obtained as mtF+/mtF+ homozygotes by the autogamy of an mtF+/mtF+ heterozygote in the mutant cytoplasmic lineage and were therefore inferred to have a completely wild-type germ-line genome (E. Meyer and A.-M. Keller, in prep.). Conjugation with an IES− wild-type cell line confirmed that the alternative processing leading to the IES+ or IES− macronuclear versions of the G gene is not genetically determined in zygotic nuclei. It was also shown that excision of the G-gene IES can be experimentally induced in the IES+ line by decreasing the macronuclear IES+ G-gene copy number [which follows from the massive injection of plasmids p629 or p851]. Because neither microinjection of plasmids into the macronuclear nor autogamy of a homozygous cell can modify the germ-line genome, this provides an independent demonstration that the IES+ cell line is genetically competent for excision of this IES. Conversely, we have shown that transformation of the macronucleus of wild-type IES− cells with IES+ plasmid p851 results in the retention of the IES in the new macronucleus differentiating after autogamy. This effect depends on the presence of the IES sequence in the plasmid insert, because it is not shared by plasmid p629, which contains the IES− version of the...
same restriction fragment, or by plasmid p851D, in which about two-thirds of the IES sequence was deleted. Although the germ-line genome was not modified in this experiment, the p851-induced IES+ character was itself transmitted to sexual progeny, like that induced by the mtB mutation. This demonstrates that nothing other than the presence of a fragment of the IES + G gene in the old macronucleus is required to turn an IES- wild-type strain into a stable IES+ cell line. Thus, during macronuclear differentiation in wild-type cells, excision or retention of the G-gene IES only depends on the structure and copy number of the G gene in the old macronucleus.

Developmental excision is inhibited by the presence of the IES in the old macronucleus

The cut-and-repair model, proposed to account for the maternal inheritance of the IES+ and IES- characters, postulates that a double-strand gap is created in the G gene by constitutive excision of the IES during macronuclear development in both IES+ and IES- cell lines and then repaired by the copying of a homologous template from the old macronucleus. The failure of IES-plasmid p629 to activate excision in the IES+ cell line, as well as the reappearance of precise excision after autogamy of clones with a reduced IES + G-gene copy number but devoid of IES- G gene, indicate that such a mechanism is not involved in the production of the correct IES- junction. Can the model still apply to the production of the IES + macronuclear version? If excision is initiated by a cut at each end of the IES, as proposed in hypotrichous ciliates for both IESs and transposon-like elements (Klobutcher et al. 1993; Jaraczewski and Jahn 1993; Williams et al. 1993), and shown for a programmed deletion event in the more closely related Tetrahymena (Saveliev and Cox 1995), a prediction of the model is that retention of the G-gene IES in the new macronucleus will critically depend on the physical linkage of IES and flanking sequences in the old macronucleus. This is not borne out by the transformation of IES- cells with plasmid pIES, which results in IES retention. The effects of modified IESs also run contrary to predictions of the model. The IES+ macronuclear G gene recovered after autogamy of p851S-transformed IES- cells contains the germ-line IES sequence, not the modified IES sequence present on the plasmid. Furthermore, an internal modification of the IES sequence in plasmid p851D abolishes the effect, which would not be expected.

Thus, the maternal inheritance of the IES+ and IES- characters appears to be attributable to an inhibition of excision of the G-gene IES in the developing macronucleus of the IES+ cell line, caused by the presence of the IES in the old macronucleus, no effect of the IES- G gene in the old macronucleus was evidenced. The results obtained with transformed cells are consistent with those obtained from un.injected cells with various endogenous IES+ and IES- G-gene copy numbers. The quantitative analysis shows that the percentage of excision in the new macronucleus is inversely correlated with the number of copies of the IES + G gene in the old macronucleus, as observed after injection of IES + plasmids in IES- cells (Fig. 7). However, excision appears to be more sensitive to the genomic IES+ sequence than to the IES+ plasmids: 90% inhibition is obtained with only 0.3-0.4 IES+ genomic copies per haploid genome in the old macronucleus or ~6 copies of plasmid p851, whereas plasmid pIES only results in 56% inhibition with 60 copies.

Possible mechanisms of excision inhibition by the maternal IES

During meiosis and fertilization, the old macronucleus breaks up into ~30 fragments in which DNA synthesis is progressively inhibited; these fragments remain transcriptionally active for the whole period of development of new macronuclei (Berger 1973). The mechanism through which the presence of the IES + G gene in the old macronucleus inhibits the developmental excision of the G-gene IES in the developing macronucleus has to operate through the cytoplasm because fusion of the fragments of the old macronucleus with the developing macronuclei is not observed under normal conditions. Two different types of mechanisms can be proposed. First, the IES + G gene in the old macronucleus could compete with the germ-line sequence for a protein factor necessary for IES excision, by titrating it out of the cytoplasm. Alternatively, a trans-acting factor could be actively produced from the IES + G gene in the old macronucleus and be exported to the developing macronuclei to inhibit excision. In both cases, the putative trans-acting factor would have to show some specificity for the G-gene IES. The vast majority of germ-line IESs are correctly excised in the IES+ cell lines obtained by p851 transformation, as shown by their wild-type phenotype. Given the very large number of IESs in the genome, an excision defect affecting a significant fraction of them would almost certainly be lethal. A PCR analysis further confirmed the correct excision of four other IESs, located in the A and B surface antigen genes.

The first type of mechanism would require that the putative excision factor, after its synthesis in the cytoplasm, be able to circulate freely in and out of the fragments of the old macronucleus before it reaches the developing macronucleus, to explain its total lack of action in IES+ cells. Furthermore, the complete titration of the factor by a small amount of IES+ G gene in the old macronucleus (90% inhibition of excision with only 0.4 copy per haploid genome, i.e., ~400 copies per macronucleus) would imply a high degree of specificity for the G-gene IES because there is no evidence for any other unexcised IES. However, highly specific protein factors are certainly not a general feature of IES excision. IESs appear to be noncoding, so that such factors would have to be encoded elsewhere in the genome. Moreover, the absence of any effect of the p851D plasmid on excision indicates that the binding site for the putative factor would have to be located within the internal 147-bp IES segment that is deleted in this plasmid. This is difficult
to reconcile with the quantitatively different efficiencies of plasmid pIES, plasmid p851, and the genomic IES + sequence, which all include the 147-bp segment.

In the second type of mechanism, an excision-inhibiting factor would be produced from the IES + G gene in the old macronucleus. Because the G-gene IES is apparently not coding (it is 80% A + T and does not contain any open reading frame) and because inhibition is observed after transformation with pIES, which only contains 214 bp of the IES, the inhibiting factor is not likely to be a protein. We propose that the IES sequence itself, or an RNA copy of it, is exported to the developing macronucleus, where it acts by pairing with the homologous germ-line sequence. Such a sequence-specific mechanism would readily explain why the excision of other IESs in the A and B genes is not affected by the G-gene IES in the old macronucleus. In this model, the stepwise increase in the efficiency of excision inhibition by the IES sequence, as one goes from pIES to p851 and to the IES + genomic sequence, might be explained by the different lengths of flanking-sequence homology to the germ-line sequence, which could lead to differences in pairing efficiency. The absence of excision inhibition by plasmids p629 and p851D would indicate that the pairing of the 147-bp segment (or simply pairing over a minimum length of the IES sequence) is required. However, all plasmids induce a quantitatively similar reduction of the G-gene copy number after autogamy. The fact that some of them, such as p629 and pIES, do not bear any common sequence, suggests that the copy number reduction effect also involves pairing rather than the titration of multiple sequence-specific proteins factors, as has been proposed for yet other effects of injected plasmids: the general and sequence-specific induction of deletions in P. primaurelia (Meyer 1992; E. Meyer, A. Butler, and F. Caron, in prep.), and the rescue of mutant telomere addition patterns in P. tetraurelia (Kim et al. 1994; You et al. 1994).

Although the data do not rule out the titration of a specific excision factor, the above arguments lead us to favor a pairing mechanism. Because it appears that retention of the IES is not attributable to repair synthesis, pairing could affect an earlier step of the excision process, which could be the recognition of the sequence to be excised, the formation of a functional excision complex, or an initial endonucleolytic cleavage. It would be of great interest to know whether a similar epigenetic regulation of excision is observed for other IESs in the genome. Experiments are now in progress to test this point and to determine the full extent of the sequence specificity of excision inhibition. Whatever the mechanism may be, its study is likely to bring new insight into the general process of IES excision.

Biological significance of epigenetic self-regulation of excision

The observation that IES excision can be epigenetically regulated in wild-type cells, like alternative telomere addition regions, highlights the great developmental plas-

ticity of the Paramecium macronuclear genome. Both types of genomic rearrangements may therefore help to explain the old observation that individual caryonides from entirely homozygous strains can display a number of stable alternative phenotypic characters that are irreversibly determined during macronuclear differentiation from a totipotent germ-line genome, with or without maternal inheritance (Sonneborn 1977; Sonneborn and Schneller 1979; Epstein and Forney 1984; Nyberg 1986).

The quantitative analysis presented in this study shows that any clone containing between 0.1 and 1 copy per haploid genome of the IES + G-gene version in its macronuclear genome will give rise to postautogamous progeny with a larger fraction of IES + copies. Only two alternative macronuclear states are therefore expected to be stably maintained through successive autogamies (0% and 100% IES + copies), explaining the stability of IES − and IES + cell lines. It should be pointed out that the dynamics of the epigenetic regulation system described here could fully account for puzzling features of the mating-type determination system in P. tetraurelia, the molecular mechanism of which is unknown. In addition to the maternal inheritance of two alternative differentiated types (Sonneborn 1977), these include the occasional and transient occurrence of mixed macronuclei in selfer clones (Nanney 1957) and the genetic restriction to one of the developmental alternatives by the pleiotropic mtFE mutation (Brygoo and Keller 1981b; E. Meyer and A.-M. Keller, in prep.).

Materials and methods

Cell lines and cultivation

P. tetraurelia wild-type strain d4.2 is a well-characterized derivative of stock 51 carrying the 29A allele of the A surface antigen gene, and gene k (Sonneborn 1974). Cells were grown in a wheat grass powder (Pines International Co., USA) infusion medium bacterized the day before use with Klebsiella pneumoniae and supplemented with 0.8 mg/liter of β-sitosterol (Merck, Darmstadt FRG), at 18°C or 27°C. Basic methods of cell culture have been described (Sonneborn 1970).

Autogamy and conjugation

Autogamy was induced by starving the cells after they had reached the appropriate clonal age (30 vegetative divisions) and assessed by staining with a 15:1 (vol/vol) mix of carmine red (0.5% in 45% acetic acid) and fast green (1% in ethanol). For caryonidal analyses, cells were isolated from depressions showing 100% autogamous cells. After the first cellular division, the two caryonides were isolated and cultivated separately. For mass autogamies, a whole depression (~1000 autogamous cells) was transferred to bacterized medium and grown collectively. Conjugation was induced by starving two clones with complementary mating types. Pairs firmly engaged in conjugation were transferred to individual depressions. The two exconjugants from each pair were isolated after their separation; After the first division, the two caryonides from each exconjugant were again isolated and grown separately. Their parental origins were determined by mating type testing of derived F2 postautogamous cells, using standard d4.2 tester strains (mating type is maternally inherited).
Dot-blot analyses

For each sample, ~500 cells were pipetted from depression slide cultures and transferred to 400 μl of 0.4 N NaOH, 50 mM EDTA. The lysates were incubated for 30 min at 68°C and loaded on a Hybond N + membrane (Amersham, UK) using a dot-blot apparatus. The membrane was kept wet with 0.4 N NaOH for 15 min, washed in 2× SSC (0.15 M NaCl, 0.015 M sodium citrate), and treated further as a Southern blot.

Microinjection of plasmids

Young cells (less than eight fissions after autogamy) were injected in Volvic mineral water (Volvic, France) containing 0.2% BSA, under an oil film (Nujol) while being visualized with a phase-contrast inverted microscope (Axiovert 35M, Zeiss). CsCl-purified plasmid DNA was linearized within the pUC18 vector sequence with ScaI (p629, p851, p851D) or XmnI (pIES, p851S), extracted with phenol, filtered on a 0.22-μm Millipore filter (Millipore), and precipitated with ethanol. Approximately 5 μl of a 5 mg/ml solution in water were delivered into the macronucleus.

Genomic DNA extraction

Cultures of exponentially growing cells (400 ml) at 1000 cells/ml were centrifuged. After being washed in Volvic mineral water, the pellet was resuspended in one volume of mineral water and added quickly to four volumes of lyss solution (0.44 M EDTA at pH 9.0, 1% SDS, 0.5% N-laurylsarcosine [Sigma], and 1 mg/ml of protease K [Merck]) at 55°C. The lysate was incubated at 55°C for at least 5 hr, gently extracted once with phenol, and dialyzed twice against TE (10 mM Tris-HCl, 1 mM sodium phosphate, 1% BSA, and 1 mM EDTA (pH 7.2) at 61°C (Church and Gilbert 1984). Probes were labeled using a random priming kit (Boehringer Mannheim, Germany) to a specific activity of 3·106 cpm/μg. Membranes were then washed for 30 min in 0.2× SSC and 0.5% SDS at 60°C prior to autoradiography or PhosphorImager quantification.

PCR amplification

Oligonucleotides were designed in the flanking sequences of IESs according to published sequences for the 51A IESs (Steele et al. 1994) and 4404 (Steele et al. 1994), and the 51B IESs 1417 (CTTGTGT-51B, IESs 1417 (CTTGTGT-) and 4404 pair, where the 51B 4404 pair, where the

Quantification of endogenous G gene and plasmid copy numbers

The endogenous G-gene copy number was determined by measuring the intensity of the PstI fragments revealed with probe b, using a PhosphorImager (Molecular Dynamics). To correct for differences in the amount of DNA loaded in each lane, the figures obtained were normalized by reprobing the same blots with a 30-bp degenerate oligonucleotide specific for macronuclear telomeric repeats, [5'-(C/A)AACCC-3']5; the telomeric signal, counted on the whole length of the lane, can be expected to a more reliable estimate of the total amount of macronuclear DNA on the blots than the signal from any genomic probe, because it does not depend on copy number variation of a particular macronuclear chromosome. Furthermore, neither macronuclear telomere length nor total amount of telomeric repeats vary significantly with clonal age (Gilley and Blackburn 1994). The linearity of the telomeric probe response was checked by quantifying the signal obtained with various amounts of the same genomic sample. Plasmid copy numbers were determined for all preautogamously transformed lines by hybridizing PvuII digests with the PUC18 probe a, as shown in Figure 5B. The correspondence between probe a and b signals is given by transformed clones such as d and e (Fig. 4A), in which the same PvuII plasmid fragment contains the sequences of both probes. The ratio can then be used to convert the probe signal from each transformed clone into the equivalent probe b signal, and furthermore, after normalization with the telomeric signal, into the relative plasmid copy number. This method could lead to an underestimation of high plasmid copy numbers if the presence of plasmid telomeres significantly increases the telomeric signal from transformed cells; however, this would only underestimate the efficiency difference noted between IES + plasmids and IES + endogenous G gene in excision inhibition.

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