These data provide further evidence for w* isoalleles and locus-specific action of w mutations in D. melanogaster. Comparison of Tables 1 and 2 shows that the phenotypic action of isoalleles in question is somewhat influenced by the genetic background. This influence, however, is not sufficient to cancel out the differences in the action of the w* isoalleles themselves and of the w mutations studied.


The present investigation was performed in order to study the effect of an electric field to somatic recombination and/or chromosome loss in larvae of Drosophila melanogaster. Previous observations on root tip cells of Allium have shown that the segregation of the chromosomes was disturbed by electric shocks.

The treatment was given to female larvae, heterozygous for yellow. The larvae were collected at random to all groups. The abdomen of hatched females was mounted on a slide in a drop of Euparal. The slides were coded and the bristles inspected under microscope. The size of the yellow spots was also recorded. In a preliminary test the females were also heterozygous for singed. As the classification of the singed character turned out to be unreliable, however, only the yellow character could be scored.

The electric treatment involved 250 volts, given to third instar larvae as a one second shock between two titan-electrodes in a bath of aq. dest. As a control that the larval age used was sensitive to the induction of somatic recombination and/or chromosome loss, one group of larvae received 1800r of X-ray at an intensity of 90r/min. A third group consisted of a control without any treatment.

In Fig. 1, the number of spots per fly is presented. The irradiated group (R) shows a significantly higher number of yellow spots than the control (C), indicating that the treated stage of the larvae was sensitive for induction of yellow spots. There is no difference between the control and the group receiving the electric shock (E).

An effect by the electric treatment is indicated, however, by the larger size of the yellow spots as compared to the control, as shown in Fig. 2, showing the number of bristles per yellow spot. The difference between groups C and E is highly significant both concerning the distribution (X^2 = 22.4, P < 0.0005) and the mean (F_1,84 = 11.7, P < 0.001). A slight tendency in the same direction occurs for the irradiated group, the difference versus the control being at the border of significance (P = 0.05). Further investigations are being performed in order to reveal the biological significance of these observations.