

3,4) it was interesting to find out whether the feature of behavior of mutation *sn* on the homoeotic structures is retained at different temperatures. For this purpose homoeotic mutations *Ns* (3-48.0) and *aristopedia* (*ss<sup>a</sup>*, 3-58.5) (its three alleles: *ss<sup>ak</sup>*, *ss<sup>ax</sup>* and *ss<sup>a40a</sup>*) (transform antennae to the legs of mesothoracic type) were used. The number of large bristles was significantly more in mutants *Ns*. The double mutants *sn;Ns* and *sn;ss<sup>a</sup>* obtained at 17, 25 and 28°C and single mutants *sn*, *Ns* and *ss<sup>a</sup>* were studied by binocular microscope (increase 12.5 x 4.0). 100 individuals of every genotype were tested to estimate penetrance, 64 to estimate expressivity.

The effect of temperature on the degree of twisting of bristles in mutants *sn* was not detected. Twisting of bristles in mutants *Ns* and *ss<sup>a</sup>* was not found at different temperatures. Twisting of bristles on thoracic legs and antennal homoeotic limbs was observed at the interaction of homoeotic mutations *Ns* and *ss<sup>a</sup>* (its three alleles) with the mutation *sn* at all temperatures (16, 25 and 28°C). The complete penetrance of this effect was observed. The degree of its expression on homoeotic structures was identical to that on the corresponding segments of thoracic legs. In particular, twisting of bristles was significantly more marked in double mutants *sn;Ns* on the homoeotic structures than in mutants *sn;ss<sup>a</sup>*. It can be explained by the predominance of large bristles on homoeotic limbs in mutants *sn;Ns*. The effect of temperature on the degree of twisting of bristles in double mutants was not found. The coincidence of expressivity of mutation *sn* on homoeotic structures with that on the corresponding segments of thoracic legs independent of temperature points out the morphogenetic relationship of elements (bristles) of homoeotic and normal structures.

References: (1) Braun, W. 1940, *Genetics* 25:143-149; (2) Ouweneel, W. 1970, *Genetics* 41: 1-20; (3) Kaurov, B.A., V.I. Ivanov and V.A. Mglinetz 1976, *Genetika* (USSR) 12:75-81; (4) Kaurov, B.A., V.I. Ivanov and V.A. Mglinetz 1978, *Genetika* (USSR) 14:306-312.

Kemphues, K.J. and T.C. Kaufman. Indiana University, Bloomington, Indiana. Two-dimensional gel analysis of total proteins from X/O, X/Y, X/Y/Y, X/Y<sup>s</sup> and X/Y<sup>L</sup> testes from *D. melanogaster*.

An attempt to identify proteins specific to the Y chromosome of *D. melanogaster* utilized the NEPHGE two-dimensional gel electrophoresis system of O'Farrell (1977), as modified by Waring (1978), to display total proteins extracted from testes. These gels offer the best one-step method for visualizing a large array of proteins. In

these experiments, resolution of more than 300 spots from total proteins of [<sup>35</sup>S] methionine labeled adult testes was possible.

Careful comparisons of autoradiographs of gels of labeled proteins from X/O, X/Y, X/Y/Y, X/Y<sup>s</sup> and X/Y<sup>L</sup> testes were made. These comparisons revealed no consistent differences in the two-dimensional pattern of proteins from testes of the various genotypes, other than an increase in the intensity of labeling of five spots from X/Y/Y testes relative to other genotypes. No protein differences, either qualitative or quantitative, were found between the autoradiographs of X/O and X/Y testes.

One basic protein of 55,000 daltons, however, did fail to appear in gels of testes from two of the four X/O stocks examined. A two-dimensional gel analysis of protein from X/Y testes at various developmental stages (24 hour intervals, starting at 72 hours after egg deposition) showed that the 55,000 dalton protein was not synthesized until late in development (220 hours; late pupa). The late synthesis of this protein suggests that some triggering event is necessary prior to synthesis. Because X/O spermatids degenerate before maturation (Kiefer 1970), it is possible that the 55,000 dalton protein appears in some X/O testes and not in others because this degeneration could begin either before or after this triggering event due to differences in genetic backgrounds or culture conditions.

These results indicate that if the Y chromosome fertility factors are expressed as proteins, the detection of these proteins will require more elaborate techniques than those described here. These experiments also demonstrate that the Y chromosome functions in a subtle manner which does not involve the regulation of synthesis of the major protein components of spermiogenesis. However, it may be possible that the Y specific proteins contain no methionine and that other labeling procedures will allow for their detection.

References: Kiefer, B.I. 1970, *J. Cell Sci.* 6:117-194; O'Farrell, P.Z. et al. 1977, *Cell* 12:1133-1142; Waring, G.I. et al. 1978, *Dev. Biol.* 66:197-206.