DROSOPHILA

Information Service

36

January 1962

Material Contributed by
DROSOPHILA WORKERS

and arranged by
E. NOVITSKI

Material presented here
should not be used in publications
without the consent of the author.

Prepared at the
DEPARTMENT OF BIOLOGY
UNIVERSITY OF OREGON
EUGENE, OREGON
### Stock Lists

**United States**

- Ames, Iowa .......................................................... 34:9
- Amherst, Massachusetts ........................................ 36:19
- Austin, Texas ....................................................... 35:16
- Baltimore, Maryland .............................................. 36:19
- Berkeley, California ............................................ 35:17
- Buffalo, New York ................................................. 35:18
- Chapel Hill, North Carolina ................................... 35:9
- Chicago, Illinois ................................................. 36:11
- Cleveland, Ohio .................................................. 35:10
- Cold Spring Harbor, New York .................................. 34:11
- DeKalb, Illinois .................................................. 36:11
- East Lansing, Michigan ......................................... 34:12
- Gainesville, Florida ............................................. 34:13
- Lafayette, Indiana ............................................... 35:11
- Lawrence, Kansas ................................................ 35:11
- Le Mars, Iowa ..................................................... 36:12
- Lexington, Kentucky ............................................. 35:11
- Los Angeles, California ......................................... 34:14
- Minneapolis, Minnesota ......................................... 35:11
- New Haven, Connecticut ......................................... 36:12
- New York, New York ............................................... 34:26
- Norman, Oklahoma ................................................ 34:15
- Oak Ridge, Tennessee ............................................ 35:11
- Pasadena, California ............................................ 36:13
- Philadelphia, Pennsylvania ..................................... 35:17
- Salt Lake City, Utah, Department of Genetics ............... 36:14
- Salt Lake City, Utah, Department of Surgery ............... 34:26
- Syracuse, New York ............................................... 35:32
- Tucson, Arizona .................................................. 35:32
- University Park, Pennsylvania ................................ 34:27
- Urbana, Illinois, Department of Genetics .................... 34:27
- Urbana, Illinois, Department of Psychology .................. 36:14

**Argentina** .......................................................... 35:33

**Australia**

- Adelaide, South Australia ...................................... 36:14
- Brisbane ............................................................ 35:34
- Hobart, Tasmania .................................................. 36:15
- Melbourne, Victoria .............................................. 34:28
- Sydney, New South Wales ........................................ 36:15

**Austria** ............................................................. 35:34

**Belgium** .............................................................. 34:29

**Brazil**

- Curitiba ............................................................. 35:34
- Porto Alegre ....................................................... 36:15
- São Paulo ............................................................ 34:63

**Canada**

- Toronto ............................................................... 36:16
- Vancouver ............................................................ 36:16

**Chile** ................................................................. 34:30

**Colombia** ............................................................ 36:17

**Denmark** ............................................................. 34:30

**Finland** ............................................................ 36:17

**France**

- Gif-sur-Yvette ..................................................... 36:19
- Lyon (Rhône) ....................................................... 36:19
- Strasbourg ........................................................... 34:32
<table>
<thead>
<tr>
<th>Country</th>
<th>Cities</th>
<th>Page(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Germany</td>
<td>Berlin-Buch</td>
<td>36:20</td>
</tr>
<tr>
<td></td>
<td>Berlin-Buch and Berlin-Dahlem</td>
<td>36:20</td>
</tr>
<tr>
<td></td>
<td>Heidelberg</td>
<td>34:33</td>
</tr>
<tr>
<td></td>
<td>Göttingen</td>
<td>35:35</td>
</tr>
<tr>
<td></td>
<td>Hamburg</td>
<td>36:21</td>
</tr>
<tr>
<td></td>
<td>Hamburg-Eppendorf</td>
<td>36:21</td>
</tr>
<tr>
<td></td>
<td>Karlsruhe</td>
<td>34:33</td>
</tr>
<tr>
<td></td>
<td>Marburg/Lahn</td>
<td>34:33</td>
</tr>
<tr>
<td></td>
<td>Mariensee</td>
<td>36:21</td>
</tr>
<tr>
<td></td>
<td>Münster/Westf.</td>
<td>36:22</td>
</tr>
<tr>
<td></td>
<td>Tübingen</td>
<td>36:22</td>
</tr>
<tr>
<td>Ghana</td>
<td></td>
<td>34:35</td>
</tr>
<tr>
<td>Great Britain</td>
<td>Bayfordbury</td>
<td>35:36</td>
</tr>
<tr>
<td></td>
<td>Birmingham, England</td>
<td>34:35</td>
</tr>
<tr>
<td></td>
<td>Cambridge, England</td>
<td>36:23</td>
</tr>
<tr>
<td></td>
<td>Edinburgh, Scotland</td>
<td>34:36</td>
</tr>
<tr>
<td></td>
<td>Glasgow, Scotland</td>
<td>35:36</td>
</tr>
<tr>
<td></td>
<td>Harwell, Berks, England</td>
<td>35:36</td>
</tr>
<tr>
<td></td>
<td>London, England            ,</td>
<td>34:36</td>
</tr>
<tr>
<td></td>
<td>Manchester, England</td>
<td>34:37</td>
</tr>
<tr>
<td></td>
<td>Sheffield, England</td>
<td>34:37</td>
</tr>
<tr>
<td>India</td>
<td>Calcutta, Indian Statistical Institute</td>
<td>34:37</td>
</tr>
<tr>
<td></td>
<td>Calcutta, University of Calcutta</td>
<td>35:38</td>
</tr>
<tr>
<td></td>
<td>Hyderabad</td>
<td>36:24</td>
</tr>
<tr>
<td>Israel</td>
<td></td>
<td>36:24</td>
</tr>
<tr>
<td>Italy</td>
<td>Milano</td>
<td>36:26</td>
</tr>
<tr>
<td></td>
<td>Naples</td>
<td>34:39</td>
</tr>
<tr>
<td></td>
<td>Pavia</td>
<td>34:39</td>
</tr>
<tr>
<td></td>
<td>Roma</td>
<td>36:26</td>
</tr>
<tr>
<td>Japan</td>
<td>Anzyo, Aichi</td>
<td>36:28</td>
</tr>
<tr>
<td></td>
<td>Chiba-Shi</td>
<td>34:40</td>
</tr>
<tr>
<td></td>
<td>Hiroshima</td>
<td>35:38</td>
</tr>
<tr>
<td></td>
<td>Kyoto</td>
<td>36:28</td>
</tr>
<tr>
<td></td>
<td>Misima</td>
<td>34:42</td>
</tr>
<tr>
<td></td>
<td>Mitaka, Tokyo</td>
<td>36:29</td>
</tr>
<tr>
<td></td>
<td>Osaka</td>
<td>34:43</td>
</tr>
<tr>
<td></td>
<td>Sapporo</td>
<td>34:44</td>
</tr>
<tr>
<td></td>
<td>Tokyo</td>
<td>34:44</td>
</tr>
<tr>
<td>Korea</td>
<td>Kongju, Chung Cheong Do</td>
<td>34:45</td>
</tr>
<tr>
<td></td>
<td>Kwangju</td>
<td>36:30</td>
</tr>
<tr>
<td></td>
<td>Seoul, Chung-Ang University</td>
<td>36:30</td>
</tr>
<tr>
<td></td>
<td>Seoul, National University</td>
<td>36:30</td>
</tr>
<tr>
<td></td>
<td>Seoul, Sung Kyun-Kwan University</td>
<td>36:31</td>
</tr>
<tr>
<td></td>
<td>Seoul, Yonsei University</td>
<td>36:31</td>
</tr>
<tr>
<td>Netherlands</td>
<td>Leiden, Rijksuniversiteit, Laboratorium Voor Stralengentick</td>
<td>35:38</td>
</tr>
<tr>
<td></td>
<td>Leiden, Rijksuniversiteit, Genetisch Laboratorium</td>
<td>35:39</td>
</tr>
<tr>
<td></td>
<td>Utrecht</td>
<td>35:39</td>
</tr>
<tr>
<td>Norway</td>
<td></td>
<td>34:46</td>
</tr>
<tr>
<td>Spain</td>
<td></td>
<td>35:41</td>
</tr>
<tr>
<td>South Africa</td>
<td>Johannesburg</td>
<td>36:33</td>
</tr>
<tr>
<td></td>
<td>Pretoria</td>
<td>36:34</td>
</tr>
</tbody>
</table>
January 1962  Table of Contents--continued  36:3

Sweden
  Stockholm  ..........................................................  35:42
  Uppsala ..............................................................  36:34
  United Arab Republic ............................................  34:47

New Mutants
  Report of A. Chovnick and A. Schalet .........................  36:37
  Report of K. S. Gill ...............................................  36:37
  Report of E. R. Grell ............................................  36:37
  Report of T. Inamumi .............................................  36:38
  Report of P. T. Ives .............................................  36:38
  Report of P. T. Ives .............................................  36:38
  Report of Shanta V. Iyengar ....................................  36:38
  Report of E. B. Lewis ............................................  36:39
  Report of V. Maeda ................................................  36:39
  Report of E. Ortiz ................................................  36:39
  Report of Verena Rohr ............................................  36:39

Linkage Data
  Report of M. M. Green ...........................................  36:40
  Report of Afton Hansen and Eldon Gardner ....................  36:40

Other Drosophila Species

Stock

  United States
    Ames, Iowa ...........................................................  34:56
    Amherst, Massachusetts ........................................  36:42
    Baltimore, Maryland ............................................  36:42
    Chicago, Illinois ................................................  36:42
    Cold Spring Harbor, New York ..................................  34:56
    Dayton, Ohio ....................................................  34:57
    DeKalb, Illinois ................................................  36:42
    Lexington, Kentucky ............................................  34:57
    Lincoln, Nebraska ...............................................  36:42
    Los Angeles, California ........................................  36:42
    New Haven, Connecticut .........................................  36:43
    New York, New York ..............................................  34:59
    Pasadena, California ............................................  34:59
    Philadelphia, Pennsylvania .....................................  36:44
    Pittsburgh, Pennsylvania .......................................  35:48
    Raleigh, North Carolina ........................................  36:45
    Richmond, Virginia .............................................  34:60
    Rochester, New York .............................................  35:48
    St. Louis, Missouri .............................................  34:60
    Tucson, Arizona ................................................  34:61

Australia
  Melbourne, Victoria ...............................................  34:62
  Sydney, New South Wales ..........................................  36:46

Austria ..............................................................  34:62

Belgium ..............................................................  34:63

Brazil
  Porto Alegre ........................................................  36:46
  Sao Paulo ..........................................................  34:63

Canada ..............................................................  34:63

Chile .................................................................  34:63

Colombia ..............................................................  36:47

Finland ...............................................................  34:64

France ...............................................................  36:47

Germany
  Berlin-Buch ........................................................  36:48
  Berlin-Buch and Berlin-Dahlem ..................................  36:48
Research Notes

Alderson, T., and M. Pelecanos. The mutagenic activity of ethylating agents by the larval feeding method in the presence and in the absence of ribonucleic acid ........................................... 36:53

Band, H. T. Comparison of viabilities and variances for homozygous chromosomes and heterozygous combinations using different matings in the test cross generation .................................................. 36:55

Band, H. T. Preliminary evidence that variation in temperature affects viability of heterozygous wild type flies .................................................. 36:56

Barigozzi, C. Inheritable melanotic tumours induced implanting ovaries of a tumourless stock in tumourous larvae ........................................... 36:58

Bennett, Jack, and Irene Y. L. Wei. Cross resistance between DDT and "Dri Die" in D. melanogaster .................................................. 36:58

Bentvelzen, P. A. J. Interdependence in "natural" selection of eye color mutants v and cn in D. melanogaster ........................................... 36:59


Castiglioni, M. C., and Raimondi G. Reszonico. Cultivation of Drosophila cells in synthetic medium .................................................. 36:61


Di Pasquale, A. New observations on the transmission of "brown spots" in D. melanogaster .................................................. 36:62

Druger, M. Development of D. pseudobscura at low temperature .................................................. 36:63

Frost, J. N. A test for nullo-II and nullo-III eggs in free X triploids .................................................. 36:63

Garcia-Bellido, A. Correlation between cytological stages in the spermatogenesis of D. melanogaster and their sensitivity to X-rays .................................................. 36:63

Gardner, E. J., and A. M. Hanson. Further studies on the transfer of the tumorous head material effect in D. melanogaster .................................................. 36:65
<table>
<thead>
<tr>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glass, H. Bentley. The mutagenic effect of a 5-r dose of X-rays</td>
<td>36:65</td>
</tr>
<tr>
<td>Glassman, E. Some observations on the prune-killer gene</td>
<td>36:66</td>
</tr>
<tr>
<td>Glassman, E., J. D. Karem, E. C. Keller, Jr., and J. McLean. Gene dosage relations at the ma-1 and ry loci</td>
<td>36:66</td>
</tr>
<tr>
<td>Goldberg, A., A. Schalet, and A. Chovnick. On the lethality of double mutants of Hm-3 and various ry mutant alleles</td>
<td>36:67</td>
</tr>
<tr>
<td>Goldschmidt, Elisabeth. The effect of silver nitrate on a melanotic tumor stock</td>
<td>36:68</td>
</tr>
<tr>
<td>Gottschewski, G. H. M., and W. Querner. Spreading of injected fluorochromes in explanted cephalic complexes of different larval stages</td>
<td>36:69</td>
</tr>
<tr>
<td>Greenberg, Rayla. Two new cases of SD found in nature</td>
<td>36:70</td>
</tr>
<tr>
<td>Grell, R. F., and E. H. Grell. A correction to the cytology of the rearrangement associated with Glazed</td>
<td>36:71</td>
</tr>
<tr>
<td>Hadorn, E., and I. Faulhaber. Range of variability in cell number of larval salivaries</td>
<td>36:71</td>
</tr>
<tr>
<td>Hanks, G. D. Selection for reduced recovery of yellow, white attached X females in D. melanogaster</td>
<td>36:72</td>
</tr>
<tr>
<td>Hansen, A. M., and E. J. Gardner. New eye phenotype in D. melanogaster expressed only at high temperature</td>
<td>36:72</td>
</tr>
<tr>
<td>Heed, W., J. Russell, and D. Harrington. Diversity and density of Drosophila in the immediate vicinity of Tucson with special reference to D. pseudoobscura</td>
<td>36:73</td>
</tr>
<tr>
<td>Hess, Oswald. Scute8 as Y suppressed lethal factor</td>
<td>36:74</td>
</tr>
<tr>
<td>Hildreth, P. Influence of different Y chromosomes on secondary nondisjunction in D. melanogaster</td>
<td>36:75</td>
</tr>
<tr>
<td>Hiraizumi, Y. Low viability induction by the segregation distorter (SD) locus; preliminary note</td>
<td>36:77</td>
</tr>
<tr>
<td>Hoenigsberg, H. F., Y. Garcia Cortés, and D. Ortíz Rubio. The male and female choice in studies of sexual preference in D. melanogaster mutants</td>
<td>36:78</td>
</tr>
<tr>
<td>Hoenigsberg, H. F., Y. Garcia Cortés, and D. Ortíz Rubio. The degree of sexual preference in D. melanogaster Cy BL and the fitness associated with it</td>
<td>36:78</td>
</tr>
<tr>
<td>Hollander, W. F. Two mosaics</td>
<td>36:78</td>
</tr>
<tr>
<td>Hollander, W. F., and Michael F. Festing. Equational exceptions from roughex males</td>
<td>36:79</td>
</tr>
<tr>
<td>Hunter, Alice S. Abnormal sex ratio in wild D. pseudoobscura</td>
<td>36:79</td>
</tr>
<tr>
<td>Imazumi, T. On a strain of XXY of D. melanogaster with two translocations</td>
<td>36:80</td>
</tr>
<tr>
<td>Iyengar, Shanta V. A male Drosophila mosaic for the Ycw+ chromosome</td>
<td>36:80</td>
</tr>
<tr>
<td>Kang, Yung Sun, and Hei Yung Lee. On Hirtodrosophila macromaculata sp. nov. from South Korea; with 7 text-figures</td>
<td>36:81</td>
</tr>
<tr>
<td>Kaplan, William D., V. E. Tindermole, and D. H. Gugler. The number of sperm present in the reproductive tracts of D. melanogaster females</td>
<td>36:82</td>
</tr>
<tr>
<td>Kikkawa, H. Strain differences in proteolytic enzyme activities in D. melanogaster</td>
<td>36:83</td>
</tr>
<tr>
<td>King, R. C. Vitellogenesis in Drosophila</td>
<td>36:83</td>
</tr>
<tr>
<td>Koch, R., and H. Buria. Dispersal rates in D. subobscura and D. obscura in relation to factors of environment, sex, and age</td>
<td>36:83</td>
</tr>
<tr>
<td>Koref-Santibañez, Susi. A comparative study of courtship behavior in some species of the mesophragmatica group of Drosophila</td>
<td>36:84</td>
</tr>
</tbody>
</table>
Lefevre, G., Jr., and Ulla-Britt Jonsson. Sperm relationships in twice-mated D. melanogaster females .......................... 36:85
Lefevre, G., Jr., and Ulla-Britt Jonsson. The effects of cold shock on D. melanogaster sperm ........................................ 36:86
Lewis, E. B. Salivary gland chromosome analysis of segregation distorer lines ................................................... 36:87
Lovelette, E., and F. Ratty. Comparisons of inbred and random-bred larval survival to 1200r .............................. 36:87
Lüönd-Luchsinger, S. The riboflavin content in Malpighian tubules of D. hydei ...................................................... 36:88
Malogolowkin, Ch. A new sibling species of the D. willistoni group ................................................................. 36:88
Malogolowkin, Ch. A new transitional race in D. paulistorum ........................................................................ 36:88
Mettler, Lawrence E. Fertility relationships of recombination-hybrid males from the cross of D. melanogaster and D. arizonensis .................................................. 36:89
Mettler, Lawrence E. Locating mutants by crossing species with chromosome differences ................................. 36:90
Mettler, Lawrence E. Drosophila pachea ........................................................................................................... 36:90
Milkan, R. D. cve phenodeviants in the progenies of wild inseminated females .................................................. 36:90
Milkan, R. D. Protection against phenocopying by pre-treatment at high temperatures .................................... 36:91
Momma, E., A. Kaneko, and T. Shima. Rate of emergence of pupae irradiated at various stages in D. virilis .......... 36:91
Moree, Ray. Relative fecundity involving the e locus in D. melanogaster ......................................................... 36:92
Moriwaki, D. A shift of sex-ratio in the progeny from irradiated males in D. melanogaster ........................................ 36:92
Moriwaki, D., and H. Ikeda. Disturbance of "sex ratio" condition by X-ray irradiation ........................................... 36:93
Mukai, T., and S. Chigusa. Radiation-induced mutation rates of polygenes controlling the number of sternopleural bristles in D. melanogaster .................................................. 36:93
Mukherjee, A. S., and R. C. Strohman. A preliminary study on the chromatographic behavior of the heterozygous and homozygous conditions of a mutant and that of wild type D. melanogaster .................................................. 36:94
Munz, P. Xanthindehydrogenase activity in D. melanogaster (Oregon-R) ......................................................... 36:94
Narain, P. Effect of age of female on the rate of egg production in D. melanogaster .................................................. 36:96
Narise, T. Genetic studies on migrating activity in D. melanogaster ................................................................. 36:97
Nash, D. Selection for changes in the manifestation of the Hairless mutant ......................................................... 36:99
Nash, Donald J. Fertility studies involving miniature-dominant ................................................................. 36:100
Nöthiger, R. Sepiapteridine and riboflavine in Drosophila .............................................................................. 36:101
Novitski, E. A comment on the accumulation of inversions in natural populations ........................................... 36:101
Novitski, E. A note on Sturtevant and Beadle's 1936 inversion paper ............................................................. 36:102
Ogaki, M. Inheritance of heat tolerance in D. melanogaster ............................................................................. 36:103
Ogita, Z. Genetic control of all-esterase activity in D. melanogaster ............................................................... 36:103
Okada, T. "Speed index" shown by the apodemes of drosophilid flies ............................................................. 36:104
Okada, T. "Compensatory adaptation" of the ejaculatory apodeme of drosophilid flies ........................................... 36:104
Oksala, T. A. The effect of autosomal inversion heterozygosity on crossing-over frequency in the X chromosome of D. melanogaster  36:104
Oshima, C. The persistence of deleterious genes in natural populations of D. melanogaster  36:105
Oshima, C. Dieldrin resistance in D. pseudoobscura  36:105
Parsons, P. A. A biochemical polymorphism in D. melanogaster  36:106
Pelecanos, M. Induced oögonial lethals in Drosophila  36:106
Pipkin, S. B. Spontaneous mosaics in the progeny of triploid females  36:110
Pozzi, L. V., S. Giavelli, G. P. Sironi, and E. Gallucci. Frequency of recessive sex-linked lethals in D. melanogaster spermatogenesis, in O2, N2 and air, with 600 r and 1200 r  36:110
Reitan, P. J., and M. E. Anan. The effects of dehydration on the frequency of irradiation induced embryonic abnormalities in Drosophila  36:111
Roberts, Paul. Autonomy of a claret nondisjunctional ovarian transplant in D. melanogaster  36:112
Ronen, Amiram. Induced somatic recombination in the third chromosome of D. melanogaster  36:112
Schepers, A. M. An interaction in Pteridine metabolism between garnet and brown genes in D. melanogaster  36:114
Schuitlen, G. G. M. A case of aberrant sex-ratio in D. melanogaster  36:114
Schwinck, I. Drosopterin formation and semi-lethality of the mutant rosy in temperature experiments  36:114
Seki, T. Absence of beta-alanine in hydrolyzate of the pupal sheathes of ebony mutant in D. virilis  36:115
Sherwood, Eva R. All-male offspring from heatshocked cultures  36:115
Shima, T., A. Kaneko, and E. Momma. Hatchability of eggs during varying lapses from the time of mating in D. virilis  36:115
Snyder, L. A. The effect on TEM-induced mutations and translocations of storing treated spermatozoa in the female  36:116
Sondhi, K. C. Selection for an invisible pattern of macrochaetes in Drosophila  36:116
Sperlich, D. Hybrids between D. melanogaster and D. simulans in nature  36:118
Stern, Curt, and Eva Sherwood. Can primordial germ cells of the genotype XXY produce functional sperm?  36:118
Stevenson, Richard. Altitudinal distribution of inversion heterozygotes in D. robusta  36:118
Stone, L. E. Structure and variation of the salivary gland chromosomes in D. affinis  36:119
Strangio, V. A. Recessive lethals, sex chromosome loss, and non-disjunction followed simultaneously  36:120
Strangio, V. A. Pseudo-allelism at spineless-aristapedic locus  36:121
Thompson, Peter E. The basis for high "nondisjunction" from maroon-like females  36:123
Tsukamoto, M. Comparative studies on the oxidation of DDT in D. melanogaster  36:124
Toyofuku, Y. Non-random association of inversions in D. nigromaculata  36:124
Valencia, Ruby M. Sex Ratio after irradiating fertilized eggs  36:126
Wedvik, Hans. The effect of low temperature on fertility of D. melanogaster males  36:127
Once again the pressure of the ever-increasing number of contributions has made it necessary to take measures to keep the size within the limits imposed by the method of reproduction and the binding, and the funds available for this work. This has been done by postponing those stock lists which have not materially changed since they appeared in either DIS-34, or DIS-35.

It is my opinion that the usefulness of DIS might be increased if the scope of the notes were broadened somewhat to include categories which may be of considerable interest to Drosophila workers, but which have no other outlet at the present time. I have therefore included several notes which may be taken as illustrations of the sort of thing I have in mind. The note in this issue on inversions in D. pseudoobscura covers a theoretical point relevant to the entire argument concerning the distribution of inverted sequences among the chromosomes of that species, and others as well, which might be important to anyone planning to do experimental work along this line. A second note treats in detail a calculation made in 1936 by Sturtevant and Beadle which I think should be in the record somewhere. In DIS-34 I have a note which represents a complete and unequivocal retraction of a previous note and in DIS-35, one which extends and explains a regularly published paper of my own. Undoubtedly all Drosophila workers can think of similar uses to which DIS might be put, and their suggestions and contributions are welcome.

Of course this once again raises the thorny issue as to whether DIS is or is not a publication, one which the editor prefers to avoid but, when pressed, must answer in the negative. Many of the early objections to considering this a publication have now been removed: copies are generally available to anyone who wishes them; complete sets are on file in the libraries of a large number of universities, both U. S. and foreign, as well as in the Library of Congress, and in many instances the notes are referred to in publications just as casually as if DIS were a publication itself. On the other hand, there is no strictly organized reviewing system, proofs are not returned to the writer, and notes are not to be quoted without permission of the author. (Continued on page 179)
Corrections and additions to list of Stocks in DIS 34:10.

1. Oregon-R: inbreeding, generation 370 on 6117
8. Samarkand 204-55: from $ #7, inbreeding, generation 55 on 6117
33. lost
51a. ras dy
51b. ras2 m/y f:
55a. sn oc/y f:
86. change to: y g53d sd/+ =
86a. y ras2 f
117a. bv
138. lost
142a. ci^D / ey^D
145. change to Multiple Chromosomes--the se is se50k
145a. +/y f := bw; sp Pol
150a. vg; se50k e 60k
162. lost
164a. sn' oc ptg3/Fm1, y31d sc8 wA Lz8 B & Fm1
164b. v m/Fm1, y A sc8 wA Lz8 B & Fm1

BALTIMORE, MARYLAND: THE JOHNS HOPKINS UNIVERSITY

Note: # = to be selected

Wild Stocks

<table>
<thead>
<tr>
<th>Number</th>
<th>Stock Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>b1</td>
<td>Amherst-34</td>
</tr>
<tr>
<td>b2</td>
<td>Canton-S</td>
</tr>
<tr>
<td>b3</td>
<td>Crimea</td>
</tr>
<tr>
<td>b4</td>
<td>Florida (inbred)</td>
</tr>
<tr>
<td>b5</td>
<td>Formosa</td>
</tr>
<tr>
<td>b6</td>
<td>Kyoto, Japan</td>
</tr>
<tr>
<td>b7</td>
<td>Lausanne-5</td>
</tr>
<tr>
<td>b8</td>
<td>Oregon-R150 (mass culture from 150th generation of sib. pair matings)</td>
</tr>
<tr>
<td>b9</td>
<td>Salta, Argentina</td>
</tr>
<tr>
<td>b10</td>
<td>Seto, Japan</td>
</tr>
<tr>
<td>b11</td>
<td>St. Louis-7 (bw)</td>
</tr>
<tr>
<td>b12</td>
<td>Stephensville</td>
</tr>
<tr>
<td>b13</td>
<td>Swedish-b</td>
</tr>
<tr>
<td>b14</td>
<td>Tuscaloosa, Alabama</td>
</tr>
<tr>
<td>b15</td>
<td>Urbana-3</td>
</tr>
<tr>
<td>b15a</td>
<td>Varese, Italy</td>
</tr>
<tr>
<td>b16</td>
<td>Woodbury, New Jersey</td>
</tr>
</tbody>
</table>

Chromosome 1

<table>
<thead>
<tr>
<th>#c1</th>
<th>br we ec rb t4/ Ins (1) sc8</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>In49, B Lz8 wA y31d</td>
</tr>
<tr>
<td>#c2</td>
<td>ec ct6 (s) car/ CLB</td>
</tr>
<tr>
<td>c3</td>
<td>f B</td>
</tr>
<tr>
<td>c4</td>
<td>f' y</td>
</tr>
<tr>
<td>c4a</td>
<td>g53k</td>
</tr>
<tr>
<td>c5</td>
<td>Lz50.d/ y</td>
</tr>
<tr>
<td>c6</td>
<td>sc cv dx v f</td>
</tr>
<tr>
<td>c7</td>
<td>sc t2 v f Tu car/ y f :=</td>
</tr>
<tr>
<td>c7a</td>
<td>v1 (suppressible)</td>
</tr>
<tr>
<td>c8</td>
<td>w</td>
</tr>
<tr>
<td>c8a</td>
<td>wA</td>
</tr>
<tr>
<td>c9</td>
<td>w m f (normal)</td>
</tr>
<tr>
<td>c9a</td>
<td>w m f (xxy)</td>
</tr>
<tr>
<td>c9b</td>
<td>y2 cho2</td>
</tr>
<tr>
<td>c9c</td>
<td>y ot f</td>
</tr>
<tr>
<td>c10</td>
<td>y ct6 ras2 f</td>
</tr>
<tr>
<td>c10a</td>
<td>y sn3 y36f (y36f - unsuppressed)</td>
</tr>
<tr>
<td>c10b</td>
<td>y2 sn51c15 ras2 v1 f</td>
</tr>
<tr>
<td></td>
<td>(v1 - suppressed)</td>
</tr>
</tbody>
</table>
**Dropping**

c11 Dp(1) scs1, y w f

**Inversions**

c12 In (1) In49, y fa^n
c13 In (1) rst3
c14 In (1) scs4 Ins scs1, y

c16 In (1) scs8, B

c17 Ins (1) scs1 scs8, B w a

c18 Ins (1) scs1 In48, B f v y/ y f =
c20 In (1) y3F, B

**Lethals**

#c21 car l(C1 + 1)/ Ins (1) scs1 scs8, B w a

c22 car l(B2 + 6)/ Ins (1) scs1 scs8, B w a

c23 car l(A3 + 3)/ Ins (1) scs1 scs8, B w a

**Closed X's**

c28 Xc17, y f =
c29 Xc2/ y f =
c30 Xc2, y v
c31 In (1) Xc2 wvc/ y w lsz4fx

**Chromosome 1**

d1 a1 b c sp2
d2 a1 dp b pr cn c px sp/ Cy sp
d2a a1 dp b pr cn c px sp/ Cy pr cn sp
d2b a1 dp b pr cn vg c a px bw mr sp/
S2 Cy lt3 pr+ Bl cn2 L4 sp2
d3 a1 Sp b L24/ Cy
d4 ap(49)/ Cy
d4a b pr cn

d4b b Tft vg/ b vg
d5 b vg
d6 BL L/ Cy
d6a BL L/ Sm5, a12 Cy ltY sp2
d7 bw (ex.49)
d7a on Su-Pm
Cy on vg Pm
d7b on Su-Pm Tac
Pm (dp b c? )
d7c c px sp
d8 dp32
d9 L2
d9a l(2) me
d10 M10.29.29/ Cy
d10a mi/ Pm2
d11 net S ho/ Cy E-S
d11a pvs
d11b Pfd/ Ins (2L,2R) Cy S2
d11c px sht sp
d12 rn/ Cy
d13 S Sp Bl L/ Cy cn2 sp
d14 stw2 (ex.51.6.12a.3)/ Cy
d14a Tac sp/ Cy sp
d15 Tft/ Cy

**Deficiencies**

d16 Df (2) bw5/ Cy sp
d17 Df (2) dpY51/ Cy

**Inversions**

d18 In (2R) bw8/ Cy
d19 Ins (2L,2R) Cy bwV2/ al dp b pr
ccn c px sp
d20 In (2LR) bwV29/ Cy
d21 In (2LR) bwV30k1/ Cy
d22 In (2R) bwV30k10/ Cy

d23 Ins (2L,2R) Cy bwV34/ b vg
d24 In (2) b bwVDe 1/ b ltl cn mi sp
d25 In (2) bwVDe 2/ Rev l

d26 In (2) bwV13/ Cy

**Chromosome 3**

e1 bar-3
e1a e11
e2 g1 bxD/ Inv LVM
e2a g1 Sb/ LVM
e3 Ly Sb/ Inv LVM
e3a l (3) tr/ M6 Sb
e4 M(3) y g1/ Inv LVM
e5 M(3) y Sb/ Inv LVM
e6 p ss bx/ T(2;3) Xa
e6a red
e7 ru h th st cu sr es ca
e8 Sb bxD/ T(2;3) Xa
e9 se
e10 se e
e11 stbrk
e11a st(ex.b8)

**Chromosome 4**

#f1 Ce/ ci eyR
f2 spa
f3 svn

**Multichromosomal**

g1 b(Su-er+)/ bw; st er
g1a Swedish b erupt
g1b b(Su-er)+ Pfd bw; st er
g1c b(Su-er)+ Tft bw; st er
g1d b(Su-er)+ bw; st er bx/ st er Pr
g1e cn bw; e
g1f ctg2v; bw; e; (ey)2+
g2 Cy/ Pm dsJ3k; H/ Sb-C
g3 Cy pr cn/ Pm dsJ3k; H/ Sb-C
January 1962

Melanogaster - Stocks - Baltimore

36:11

Aberrations

#g10 v; In(2R) bw$2/ v; +
g11 Ins (1) sc8L S. sc8R, wa B

Chromosome 1

Ins (1) sc8L S. sc8R, wa B
Ins (1) sc8L S. sc8R, wa B/yf: = sc cv y f
sc8 Y(y+)/ yB & yf: = y

Chromosome 2

sf2
tu 503

#g11a T(y; 2) j/ px bw sp
#g11b T(y; 2; 3) F; st/ ri pP
#g12 T(2; 3) bw$2/ st
#g13 T(2; 3) bw$5 st/ T(2; 3) pGr st
#g14 T(2; 3) bw$De/ Cy
#g15 T(2; 3) M6/ ru h th st cu sr e's
#g16 T(2; 3) pGr/ Cy
#g17 T(2; 3) rw Cy sp
#g18 T(2; 3; 4) bw$ok18 Ins (2LR)/ Cy
#g19 T(2; 3) Cy/ pr cn; by
#g20 T(2; 3)05 Sp (L$4)+/ pr cn; by
#g21 T(2; 3)05 Sp (L$4) DL (Pr)+/ pr cn; by
#g23 T(2; 3) Sp; Dl Pr/ pr cn by
#g24 T(2; 4)/ pr cn; ci eyR

Tumor Stocks
tu A2 cito-pl-st
tu B3 (Italy)
tu-53# Jacobs (on 2R)
g8a see Multichromosomal listing

CHICAGO, ILLINOIS: UNIVERSITY OF CHICAGO
Department of Zoology

Stocks listed in DIS 35 with the following numbers are no longer kept in culture:
11, 13, 36, 37, 41, 44, 46, 49, 50, and 64.

DeKALB, ILLINOIS: NORTHERN ILLINOIS UNIVERSITY
Department of Biological Sciences

In addition to that listed in DIS 34.

Wild Stock

Oregon R.

Chromosome 1

Ins (1) sc8L S. sc8R, wa B

Chromosome 2

sf2
tu 503

bw tu
cn bw
ar dp b pr px sp

Chromosome 3

tx
ru h th st pP cu sr e's

Multichromosomal

Cy al$2 sp$2/Pm; Ubx 130/ Sb
Cy/Pm ds J3$k; H/In3R mo sr
y f: =; bw; e; ci yR
bw; st
y sc 5I In 49 sc8'; bw; st pP
Wild Stocks

a-1 Oregon-R

Chromosome 1 (X)

b-1 f
b-2 
In(1)dl-49, y w
b-3 
Ins(1)sc8, dl-49, sc8 v f/y f:=
b-4 se ec cv ptg3 v/y v f car:=
b-5 w
b-6 y se cv v f car/y f:=

Altered Y Chromosomes

c-1 yBS (BS YL'bb+YS)/y v c YBS
(BS YL'bb+YS)/y f:=
c-2 ybw+ (YL bw+bb+YS)/y v; bw c
 y v; bw f

Chromosome 2

d-1 b cn e bw
d-2 bw

d-3 bw81
d-4 bwAm

d-5 bwM58

d-6 bwM59

d-7 bw75

d-8 bw59

d-9 cn su-Pm/S1, al2 Cy sp2

d-10 Df(2) bw5 sp2/Ya

d-11 
Inst(2L+2R)Cy, bw45a sp2

ox45a/B1
d-12 px
d-13 px bw sp
d-14 sp
d-15 vg
d-16 vg/Inst(2L+2R)Roi, bw45a sp2

ox45a

Chromosome 3

e-1 e
e-2 ry2

Multichromosomal

f-1 y:=/Y; bw; In(3LR)Ubx130, Ubx130 e/st (1;2;3)
f-2 y:=/Y; bw81; In(3LR)Ubx130, Ubx130 es/st (1;2;3)
f-3 y:=/Y, bw75; In(3LR)Ubx130, Ubx130 es/st (1;2;3)
f-4 y; bw; e; ci eyR (1;2;3;4)
f-5 y:=/Y; In(3LR)Ubx130, Ubx130 es/
In(3R)Vno, Vno (1;3)
f-6 bw; st (2;3)
f-7 bw81; st (2;3)
f-8 bwAm; st (2;3)
f-9 bwM58; st (2;3)
f-10 bwM59; st (2;3)
f-11 bw75; st (2;3)
f-12 bw59; st (2;3)
f-14 bw59; In(3LR)Ubx130, Ubx130 es/st (2;3)
f-15 In(2L)Cy, Cy px bw sp/b55; st
(2;3)
f-16 In(2L)Cy, Cy px bw81 sp/b55; st
(2;3)
f-17 In(2L)Cy, Cy px bw75 sp/b55; st
(2;3)
f-18 In(2L)Cy, Cy px bw59 sp/b55; st
(2;3)
f-19 In(2L)Cy, Cy/b55; st (2;3)
f-20 Ins(2L+2R)S1, al2 Cy sp2/Bl;
In(3LR)Ubx130, Ubx130 es/
In(3R)Vno, Vno (2;3)
f-21 px bw sp; st (2;3)
f-22 px bw81 sp; st (2;3)
f-23 px bw75 sp; st (2;3)
f-24 px bw59 sp; st (2;3)
f-25 px sp; st (2;3)
f-26 px; st (2;3)
f-27 sp; st (2;3)
f-28 vg; e (2;3)

NEW HAVEN, CONNECTICUT: YALE UNIVERSITY
Department of Zoology

Stock list remains essentially as in DIS-74-14 except for the following corrections and additions:

58 y2 w6 cv sn55a v +f /M-5
60 w6 c460c/M-5
114 sc8 In S w6 sc8; In SM1, al Cy sp2/
dp h Pm ds33k; C Sb/Ubx130 es (H-40)

136 Df(1) w258-21, see No. 153
154 trp/In(3LR) Ubx130 ( FM43/w3 v )
155 y2 w3 m f
Note: The following is a list of additions, losses, and corrections to the list of stocks from this laboratory in DIS 34. The convention for listing new stocks can be illustrated by an example; the new stock, w sn3 m, is given the number 143b and should be inserted after 143 in the Pasadena DIS 34 stock list. Some minor typographical errors in the DIS 34 list will be corrected the next time the full stock list is reprinted.

Stock Additions to DIS 34 list:

**Chromosome 1**
- 6b. amx 1s8 v/ y f : =
- 31b. ec ot6 s car/ FM6, y31d sc8 dm B
- 60b. 1z36 / y f : =
- 110b. sn3 la3yh v/ y f : =
- 135b. v f Bx49k car/ y f : =
- 143b. w sn3 m

**Chromosome 2**
- 200b. al S ast ho/ SM1, al2Cy sp2
- 200c. alpha-1 (pP)
- 319b. lt std/ SM2, al2Cy ltV sp2
- 346b. pd l1
- 381b. SD-5/ SM5, al2Cy2ltV sp2
- 381c. SD-72 / SM5, al2 Cy ltV sp2

**Multichromosomal Stocks**
- 641b. b (Su-er*) bw; st er (2;3)
- 643b. cn; ry2
- 647b. Su-er tu bw; st er su-tu (2;3)

**Attached-X**
- 652b. Y DM / FM6, y31d sc8 dm B

**Closed-Y**
- 659b. Y0, bw / X+; bw (fb "MIR")

**Inversions-X**
- 720b. In(1) dl-49, y Su-Hw Hw m2 g4 / y f w : =
- 721b. Ins(1) dl-49, BM1, y sc v cu-x BM1

**Stock Losses:**

**Chromosome 1**
- 20 Bxr49k/ y f : = (replaced by 135b)
- 33 ec dx/ dl-49, y Su-iw Hw m2 g4 (replaced by 32 and 720b)

**Chromosome 2**
- 200 al S ast ho/ Cy, En-S (replaced by 200b)
- 327 M(2)p/ Cy, al2 lt3 1/4 sp2 (replaced by 395 and 274)
- 394 Sp j/ In(2L) Cy-t, Su-S dp2 pr (replaced by 395 and 274)

**Chromosome 3**
- 550 ry2 (replaced by 643b)

**Corrections to DIS 34 list:**

For:
- 653 y2 su-wa bb
- 718 dl-49, ty-2 bbl
- 749+ Ins(2L+2R)Cy, (2R)bwY34 (314, 315, etc.)
- 749++ Ins(2L)Cy + (2R)NS (333)
- 759+ with st 1(3)W ca in)

Read:
- y2 su-wa w4 bb dl-49, ty-1 bbl
- Ins(2L+2R)Cy, (2R)bwY34 (333)
- Ins(2L)Cy + (2R)NS (345)
- with st 1(3)W ca (in 578)
Note: Only unusual stocks are listed.

**Wild-type**

1. Salt Lake City
2. Solway
3. Chromosome 1
   3. lix
4. Chromosome 4
   4. ar/eyD
   5. bt
   6. btD/ciD
   7. Ce2/spaCat
8. ci ey1 eyR svn
9. ci +
10. ci +4
11. ci +5
12. ci +/eyD
13. ciD/spaCat
14. eyD/Scn
15. 1 (4) PT-1/eyD
16. 1 (4) PT-2/eyD
17. 1 (4) PT-3/eyD
18. 1 (4) 4d/ciD
19. 1 (4) 5e/ciD
20. 1 (4) 6f/eyD
21. 1 (4) 7g/eyD

**Deficiencies**

19. y/z57j
20. Xc2/scS1
21. X.r3/YLC
22. l (4) 10k/ciD pol
23. l (4) 14o/ciD
24. l (4) 25x/eyD
25. spa
26. spa pol
27. pr; Mal
28. y; bw; e; ci eyR
29. Df (4) M-4/eyD
30. vg
31. b vg
32. ca
33. e4 wo ro

**Behavioral stocks**

1. Positive geotaxis over 120 generations of selection for performance in mass
2. Negative geotaxis screening maze
3. Also other stocks used in geotaxis experiments on reversed and relaxed selection as well as the foundation population.

**Australia**

Adelaide, South Australia: University of Adelaide, Department of Genetics

**Wild**

1. Canton S

**Chromosome 1**

2. B
3. sd
4. car
5. ct v f
6. 42
7. Muller-5
8. rb cx
9. sc cv v f
10. v
11. w
12. wa55b
13. w sat
14. w m f
15. y
16. y
17. y w spl
18. y wD sc ec
19. y/iz57j
20. Xc2/scS1
21. X.r3/YLC
22. al
23. al dp b pr c px
24. cn
25. b j
26. bw
27. dp
28. fj wt/Xa
29. ho
30. vg
31. b vg
32. ca
33. e4 wo ro
34. Ly/D3
35. ss
36. ci eyR
37. ey2

**Chromosome 4**

38. bw; st
39. v; bw
40. y; CY/pr;
41. y w; dp
42. e; bw
43. e; vg
44. e; dp

**Multichromosomal**

45. Df (4) M-4/eyD

**Multichromosomal**

46. Df (4) M-4/eyD

**Chromosome 3**

47. Df (4) M-4/eyD
Hobart, Tasmania: University of Tasmania, Department of Zoology

Wild Stocks

Canton-S

Several strains from different places in Tasmania.

Chromosome 1

101 ct v f^5
102 y / 1z-7j
103 Bag/od^5g
104 sc^51 In-S B apr sc^8 (Basc)
105 C1B / y^2 apr ec cv ct v f
106 y / B
107 Xc2 / sc^51
108 Xc2 Y v f
109 Xc2 v f / y
110 y v f / w
111 B
112 y^W / y / sc^8.Y
113 sc^51 In-S apr sc^8
114 y apr
115 y^¥^5g
116 y apr / sc^8.YB-S

Chromosome 2

201 cn bw
202 Cy L / Pm ds^33k

203 S / Cy L^4
204 r-j wt / Xa T 2:3
205 Cy pr / al dp b pr px sp
206 b cn c bw

Sydney, New South Wales: Sydney University, CSIRO Animal Genetics Laboratory

Three stocks of sc are held which may be of interest -

1. homozygous for sc w^, average scutellar bristle number of something over 4.
2. homozygous for sc w^(1), has a scutellar bristle number of 2, this number having been rendered rather invariable by selection.
3. sc w^ x sc w not balanced. In this stock the mean scutellar bristle number of + w^(1)
   + males is about 7 and ++ females about 8 1/2.

These stocks are referred to as follows: sc w High, sc w^(1) LV, sc w High + w^(1)

For more complete list, see DIS 34.

BRAZIL

Pôrto Alegre: Universidade do Rio Grande do Sul, Departamento de Genética, Instituto de Ciencias Naturais

Chromosome 1

yellow vermilion
scute - crossveinless - vermilion - forked

miniature
forked
carnation
vermilion
white
honey
blood
eosin
prune
carmine

Chromosome II
clot
purple
cinnabar

DIS: 35 remains the same, with the following additions:

Chromosome I
B
car
g
m g f
w
y w
y v f
y
w^a
y w m
y v f car
Basc
sr
st
se h

Chromosome II
bw
b vg
c
cp
px
vg

Chromosome III
bar^-3
ss
es

Chromosome IV
ey2

Multichromosomal
w, e
w, e, pol
Cy, e
bw, e
B w^a
bw pol
al dp b pr Bl c px sp/SM al_2 Cy sp^2

Inversions
In (1)y^4 y
In II CQ
Cy

Translocations
T (2,3) Xa/l(3) Xa R
ri p^F/st T (y,2,3) F

Vancouver, British Columbia: The University of British Columbia, Department of Biology and Botany

Wild Stocks

<table>
<thead>
<tr>
<th>Chromosome 1</th>
<th>4 m</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>5 w</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>1 Urbana-S</th>
<th>2 B</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3 la/ClB</td>
</tr>
</tbody>
</table>
## Chromosome 2
- 6 b pr c px sp
- 7 bw
- 8 dp
- 9 L/Cy
- 10 vg

## Attached X's
- 13 Y and w

## Chromosome 3
- 11 e

## Chromosome 4
- Multichromosomal

## COLOMBIA

**Bogotá: Universidad de Los Andes, Departamento de Genética**

### Wild Stocks
1. Sao Paulo
2. Oregon-K
3. Pacific
4. Canton-S
5. Oregon-R
6. Pavia
7. Varese
8. Anzyo-Aichi
9. Canton Special
10. Chausuyama-Aichi
11. Hachijijima
12. Hiroshima
13. Hita-Kyushu
14. Omogo-Shikoku
15. Oregon
16. Shibuya-Hokkaido
17. Suzuka-Mie
18. Takagicho-Tokyo
19. Yonekawa-Yamaguchi
20. African Strains

### Selection Stock
- Lobe (Artificial selection of 1 eye)

### Inbred Lines
- Edinburgh
- Oregon-R (320 generations)
- b pr (77 generations)
- Lobe la. (3 generations)

## FINLAND

**Helsinki: University of Helsinki, Institute of Genetics**

### Wild Stocks
1. Berlin
2. Canton-S
3. Oregon-K
4. Oregon-R-S
5. Porvoo
6. Swedish-b

### Multichromosomal
36: 18 Melanogaster - stocks - Finland

9 f
10 fu/CLB
11 g2 ty & X
12 In(1)dl 49, y fa
13 In(1)rst3, rst3
14 In(1)sc4, y sc4
15 In(1)wM4
16 Ins(1)scS1L S, sc8R, scS1 w B
17 In(1)sc8, y31d sc8 dm B 1
18 ras2
19 s
20 sc
21 sc cv v f
22 sd1 (se)
23 sn3
24 sp
25 w
26 wS sn/CLB
27 Xw v B & y
28 y ac v
29 y sn3 bb
30 y v f
32 r

Chromosome Y

33 f, Y5/yL
34 X, Y3/yS (Neuhaus)
35 In(1)wm4 and extra Y
36 In(1)wm4; rl and extra Y

Chromosome 2

37 al dp b pr c px sp
38 a12 Cy, InL 1t3/b pr Bl 1t3 cn2
39 Bl L2/Cy
40 bw
41 cp2 InCyR cg sp2/InsNS px sp
42 D3/Payne
43 dp2 ab2 pr Bl rn NSR mr/a12 Cy
44 dp2x sp cn2/S2 Cy cn2 (homozygous, InCyR)
45 f1
46 f1 px
47 In(2L)Cy, a12 ast3 b pr (Cy not present)
48 Ms, b mr/Cy
49 rl
50 rn/Cy
51 rn/Cy cn2 sp2
52 rn/Cy Bl cn2 L4 sp2
53 rn In(2R)W/Cy cn2 sp2
54 stw
55 vg

Chromosome 3

56 Bg0/In(3R)C, l(3)a
57 D3 Sb/InLP Dfd InRp ca
58 e
59 e11
60 Gl Sb/LVM
61 In(3R)DF, st DLB/In(3R)pW, st 1(3)w ca
62 In(3R)pFLA (homozygous)
63 Ly Sb/LVM
64 Me, InL Sb/ru h D InsCxF
65 R Ly/In(3L)P, gm
66 se
67 se app
68 se rt2 th/me, InL
69 tra/me, T23
70 W Sb/InsCxF

Chromosome 4

71 ci
72 ciW
73 ey
74 spa
75 svn

Multichromosomal

76 Cy/Pm; D/Sb
77 vg; e
78 wM4; Cy/ap4 vg
79 wM4; Cy/blt

Deficiencies

80 Df(2)MS-4/SML, a12 Cy sp2
81 Df(2)MS-8/SML, a12 Cy sp2
82 Df(2)MS-10/SML, a12 Cy sp2
83 Df(2)rl10a 1t cn/Cy
84 Df(2)rl10a 1t cn/Pm ds3k

Translocations

85 T(Y;2)B/b c
86 T(Y;2)c/cn3
87 T(2;3)rn/Cy sp
88 T(2;3)Xa/Sb Ubx
FRANCE

Lyon: Rhône Laboratoire de Zoologie expérimentale, Faculté des Sciences,
16, quai Claude Bernard

Wild Strains

Oregon R.
Lyon
Champetières (inbred)
Algérie

Gif-sur-Yvette (Seine et Oise): Centre National de la Recherche scientifique,
Laboratoire de Génétique formelle

<table>
<thead>
<tr>
<th>Chromosome 1</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1 - B</td>
<td></td>
</tr>
<tr>
<td>2 - car bb</td>
<td></td>
</tr>
<tr>
<td>3 - car od f</td>
<td></td>
</tr>
<tr>
<td>4 - cv</td>
<td></td>
</tr>
<tr>
<td>5 - cv v f</td>
<td></td>
</tr>
<tr>
<td>6 - rb g</td>
<td></td>
</tr>
<tr>
<td>7 - v</td>
<td></td>
</tr>
<tr>
<td>8 - v f</td>
<td></td>
</tr>
<tr>
<td>9 - Basc : sc^S^1 B In S wa sc^8</td>
<td></td>
</tr>
<tr>
<td>10 - X^c^2t^2 ; y f : =</td>
<td></td>
</tr>
<tr>
<td>11 - y f</td>
<td></td>
</tr>
<tr>
<td>12 - y f : = ; y Z we</td>
<td></td>
</tr>
<tr>
<td>13 - y v m f d1.49</td>
<td></td>
</tr>
<tr>
<td>14 - y w</td>
<td></td>
</tr>
<tr>
<td>15 - y w spl sn^3</td>
<td></td>
</tr>
<tr>
<td>16 - y wa cv v f</td>
<td></td>
</tr>
<tr>
<td>17 - y wa spl rb</td>
<td></td>
</tr>
<tr>
<td>18 - y^2su wa^ab / d1.49 yw 1z / Y B S</td>
<td></td>
</tr>
<tr>
<td>19 - d y^2 LL / y^2 oc 1z Y^S ; y^2 LL / y ac sc pn co rb cm ct6 sn^3 oc ras v m g^2 f car / sc^S^1 B In 49 1z^5 sc^8</td>
<td></td>
</tr>
<tr>
<td>20 - Y : y bw^+ / yv bw</td>
<td></td>
</tr>
<tr>
<td>21 - Y^v X In E N v y . LL se^8 y^+</td>
<td></td>
</tr>
<tr>
<td>22 - wa</td>
<td></td>
</tr>
<tr>
<td>23 - wa^B / wa^B^+ : = f ; d^B</td>
<td></td>
</tr>
<tr>
<td>24 - wa</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 2</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>25 - al</td>
<td></td>
</tr>
<tr>
<td>26 - al dp b pr</td>
<td></td>
</tr>
<tr>
<td>27 - al dp b pr c px sp</td>
<td></td>
</tr>
<tr>
<td>28 - BL L / Cy</td>
<td></td>
</tr>
<tr>
<td>29 - b pr</td>
<td></td>
</tr>
<tr>
<td>30 - bw</td>
<td></td>
</tr>
<tr>
<td>31 - cn</td>
<td></td>
</tr>
<tr>
<td>32 - dp</td>
<td></td>
</tr>
<tr>
<td>33 - dp bw</td>
<td></td>
</tr>
<tr>
<td>34 - J^3_4e^3</td>
<td></td>
</tr>
<tr>
<td>35 - L</td>
<td></td>
</tr>
<tr>
<td>36 - Pm / Cy</td>
<td></td>
</tr>
<tr>
<td>37 - Tft / Cy</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 3</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>38 - c3g</td>
<td></td>
</tr>
<tr>
<td>39 - D C X F / Dfd</td>
<td></td>
</tr>
<tr>
<td>40 - ri pp</td>
<td></td>
</tr>
<tr>
<td>41 - st</td>
<td></td>
</tr>
<tr>
<td>42 - th st cp</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 4</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>43 - ey^2</td>
<td></td>
</tr>
<tr>
<td>44 - ey^2ci</td>
<td></td>
</tr>
<tr>
<td>45 - ey^ci</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Multichromosomal</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>46 - c^3_g ; cn b</td>
<td></td>
</tr>
<tr>
<td>47 - Cy / Pm ; H / Sb</td>
<td></td>
</tr>
<tr>
<td>47 - cv ; e</td>
<td></td>
</tr>
<tr>
<td>48 - cv , f ; e</td>
<td></td>
</tr>
<tr>
<td>49 - Cy , v f ; e</td>
<td></td>
</tr>
<tr>
<td>50 - sc^S^1 B In S wa sc^8 ; Cy / Pm ; H / Sb</td>
<td></td>
</tr>
<tr>
<td>51 - d sc LL / sc WB Y^S ; Cy In / S Sp ; ab^2lid</td>
<td></td>
</tr>
<tr>
<td>52 - Sifter : S Sp - T ( 2-3 ) P-. In S D X F / Sm1. al^2 Cy ; Dl H e p^1</td>
<td></td>
</tr>
<tr>
<td>53 - T ( 2-3 ) E / Cy - R</td>
<td></td>
</tr>
<tr>
<td>54 - tu ; e</td>
<td></td>
</tr>
<tr>
<td>55 - tu ; w</td>
<td></td>
</tr>
<tr>
<td>56 - v f ; e</td>
<td></td>
</tr>
</tbody>
</table>
Wild Stocks
1 normal (Berlin wild)
2 normal (England)

Chromosome 1 (X)
3 w
4 w sn^3
5 wbf
6 wa
7 we
8 wco sn^2
9 wch wy
10 wmb
11 gt w^a
12 y
13 y^303
14 y w
15 y pn
16 y cv v f
17 y w bb
18 y fa wy^2 g^2
19 f
20 sc
21 sc rb cv
22 sc ec ct
23 spl
24 m
25 B
26 car bb Y; bb
27 v
28 cv
29 car
30 fa^n
31 ct
32 fu^f/C1B
33 vv^+/
34 vv/w^e
35 vv/x^c
36 v w f/+
37 v w f/B
38 f/C1B
39 v/C1B
40 y w/C1B
41 w^e bb^l/C1B
42 sc ec ct^6 v s^2 f
43 17/de 49, yMw y^lz
44 sc^31 In S w^a sc^8
45 sc^1B In S w^a sc^8 = M-5
46 sc ec ct v g f
47 sc ec ct v g

Chromosome 2
48 j
49 bw
50 bwpp
51 b cn vg
52 L^2/Cv
53 S Sp ab^2 ltd/NS px Sp
54 b pr vg a sp
55 vg
56 bw cn
57 al dp

Chromosome 3
58 e^11
59 st
60 p^D

Chromosome 4
61 Dfd^r-L
62 r h st Dfd p^p ss e^8
63 ri a
64 ss^a
65 jv se

Multichromosomal
66 ey^2
67 ci ey^L

Tumor Stocks
75 sc e^11 tu 49n
76 tu^b

Ringchromosome
77 sc^8. Y/y B c
78 Xc^2, yw d
79 y In 49 v f d
80 Cy al^2 1t^3 L^4 sp^2/+;
C M^e Sb C/+ 81 fj px sp; p^D/C M^e Sb C
83 Df(4)M^4/e^y^D
### Hamburg, Von-Melle-Park 10: Zoologisches Staatsinstitut und Zoologisches Museum

**Wild Stocks**

<table>
<thead>
<tr>
<th>Chromosome 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Oregon-S</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. w</td>
</tr>
<tr>
<td>3. v/f</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>7. cu</td>
</tr>
<tr>
<td>8. ss</td>
</tr>
<tr>
<td>9. Sb/H Payne</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>10. ey^2</td>
</tr>
</tbody>
</table>

### Hamburg-Eppendorf: Universitäts-Frauenklinik, Strahlenbiologische Abteilung

**Wild Stocks**

1. normal (Berlin wild)

<table>
<thead>
<tr>
<th>Chromosome 1 (X)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. CIB/+</td>
</tr>
<tr>
<td>3. sc^S1 B InS w^a sc^8</td>
</tr>
<tr>
<td>4. sc^8 Y/y f x sc^8 Y/xc2 y v</td>
</tr>
<tr>
<td>5. w</td>
</tr>
<tr>
<td>6. X^2/CIB</td>
</tr>
</tbody>
</table>

### Mariensee: Max-Planck-Institut für Tierzucht und Tierernährung

**Wild Stocks**

Berlin
Oregon-R
Canton-S

<table>
<thead>
<tr>
<th>Chromosome 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. gl</td>
</tr>
<tr>
<td>2. ru</td>
</tr>
<tr>
<td>3. ro</td>
</tr>
<tr>
<td>4. v</td>
</tr>
<tr>
<td>5. ry^2</td>
</tr>
<tr>
<td>6. Chromosome 1+3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>7. S/Cy, E-S</td>
</tr>
<tr>
<td>8. pc^2</td>
</tr>
<tr>
<td>9. cn</td>
</tr>
<tr>
<td>10. bw</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chromosome 1+2</th>
</tr>
</thead>
<tbody>
<tr>
<td>11. ma-1; cn</td>
</tr>
<tr>
<td>12. cu, kar</td>
</tr>
</tbody>
</table>
Melanogaster - stocks - Germany

DIS: 36

Wild Stocks

a1 + Crimea
a2 + Oregon R

Chromosome 1 (X)

b1 In(4) snx2 & y f:=
b2 ras^4 n/C1b
b3 v
b4 w
b5 y2 wa sn5 B & y

Combinations of scute or similar inversions

d1 ("Binsc") sc51 B In49 sc8 & y f:=
d2 ("new Binscy") y sc51 B In49 sc8

Altered Y's sometimes with mutants in X

f1 sc8 Y/y51 sc8 B f.In49 v
f2 sc8 Y B sc8/y2 Y f sc. 6 & y f:=

Sterilizer ("sz") Stocks

f3 ("sz +") YLo/X.YS
f4 ("sz bw") YLo/X.YS; bw
f5 ("sz c") YLo/X.YS & y v f:=; c
f6 ("sz e") YLo/X.YS & y v f:=; e

Chromosome 2

g1 bri
g2 bw (iso 2, 1959)
g3 bwD

Wild Stocks

1 Sevelen (Zürich)
2 Berlin (Marburg)
3 Oregon R

Tübingen: Max Planck-Institut für Biologie

Chromosome 1

4 B
5 Df(1)bb In(1)bb-, y sl2 bb+/Fm4,
   y31d sc8 dm B (Extra Y's)

Münster/Westf.: Institut für Humangenetik der Universität Münster

Chromosome 2+3

cn; ry2
(Sb; Ubx¥Xa
(Cy SM1; Ubx¥Xa

Chromosome 4

ey
eyR

Wild Stocks

1 + Crimea
2 + Oregon R

Chromosome 1 (X)

b1 In(4) snx2 & y f:=
b2 ras^4 n/C1b
b3 v
b4 w
b5 y2 wa sn5 B & y

Combinations of scute or similar inversions

d1 ("Binsc") sc51 B In49 sc8 & y f:=
d2 ("new Binscy") y sc51 B In49 sc8

Altered Y's sometimes with mutants in X

f1 sc8 Y/y51 sc8 B f.In49 v
f2 sc8 Y B sc8/y2 Y f sc. 6 & y f:=

Sterilizer ("sz") Stocks

f3 ("sz +") YLo/X.YS
f4 ("sz bw") YLo/X.YS; bw
f5 ("sz c") YLo/X.YS & y v f:=; c
f6 ("sz e") YLo/X.YS & y v f:=; e

Chromosome 2

g1 bri
g2 bw (iso 2, 1959)
g3 bwD

Wild Stocks

1 Sevelen (Zürich)
2 Berlin (Marburg)
3 Oregon R

Tübingen: Max Planck-Institut für Biologie

Chromosome 1

4 B
5 Df(1)bb In(1)bb-, y sl2 bb+/Fm4,
   y31d sc8 dm B (Extra Y's)
**Chromosome 1**

1. flp
2. ptg2
3. ras2
4. y.f.car

**Chromosome 2**

5. al, dp, b, pr, cn, px, sp.
6. CyI4/d,b.

**Chromosome 3**

8. by, cu.
9. cu, kar.
10. h, ri.
11. Mé/Sb.
12. th, ri, kar, Sb.
13. th, st, cp, Sb.
14. ve, h, eyg, cp.

---

**Note:** Only stocks not generally available in this country are listed.
INDIA

Hyderabad: Osmania University, Radiation Genetics Project
aided by Department of Atomic Energy
(Government of India)

Wild Stocks
1. Oregon-K
2. Madras

Chromosome 1
1. sc$^{Sl}$ B In S w$^{a}$ sc$^{B}$. Muller-5
2. yvf (XX) Attached X
3. yvf

Chromosome 2
1. Cy/ Bl L$^{2}$
2. dp b cn bw
3. Cy Bl L$^{2}$/ LVM
4. dp
5. bw
6. cn
7. b
8. dp b cn
9. dp b cn
10. dp b bw

ISRAEL

Jerusalem: Hebrew University of Jerusalem

Wild Stocks
Berlin
Canton - S
2 wild strains from various parts of Israel
4 isogenic strains derived from wild populations in Israel

Chromosome 1
Basc (Muller -5)
B
BB
B/y
f
fB
f$^{yS}$/Yf (Finland)
$^{E^S}$
Hw$^{y2c}$/Basc

11. b cn
12. cn bw
13. dp bw

Chromosome 3
1. Gl So/D
2. st

Chromosome 4
1. ey

Multichromosomal
1. y sc$^{Sl}$ In 49 sc$^{B}$; dp b cn bw -
0.1. dp b cn bw X:II
2. y sc$^{Sl}$ In 49 sc$^{B}$; Cy Bl L$^{2}$ -
0.1. Cy Bl L$^{2}$ X; II
3. y sc$^{Sl}$ In 49 sc$^{B}$; bw st
0.1 bw st X:II:III
4. bw st II&III
5. yy bw e ey X:II:III:IV:
January 1962  Melanogaster - Stocks - Israel  36:25

10 second chromosome lethal balanced over Cy L
melanotic tumor strain (a^144) homozygous for a wild second chromosome

Chromosome 3

1;2
Bld w^2/ w; Cy
v; bw
y; Cy L/Pm

1;2;3
X,Y InEN In^49 y; cn bw; e (no free Y)
(Bloomington)
y In^49 v; bw; e (Bloomington)

2;3
bw; st
cn bw; ri e (Bloomington)
Cy L/Pm; H/Sb
Cy/Pm; D/Sb
fes ms cn sp/Cy O; h ri e^8/M.e ri
(Bloomington)
pr; st

2;3;4
bw; e; ci ey^(Bloomington)

Not located

D - like
### Milan: Università di Milano, Istituto di Genetica

#### ITALY

Wild Stocks

<table>
<thead>
<tr>
<th>Wild Stocks</th>
<th>33) net</th>
<th>Chromosome 3:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Canton - S</td>
<td>34) so</td>
<td>38) H/Sb sr In(3R) Mé</td>
</tr>
<tr>
<td>2) Chieti - v</td>
<td>35) so² b cn</td>
<td>39) ltr/ Sb sr In(3R) Mé</td>
</tr>
<tr>
<td>3) Crkwenika</td>
<td>36) SoC</td>
<td>40) R Ly/In(3R) P, gm</td>
</tr>
<tr>
<td>4) Gaiano</td>
<td>37) spt</td>
<td></td>
</tr>
<tr>
<td>5) Jaslo o. c.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6) Moltrasio</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7) Oregon - R</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8) Pavia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9) S. Maria</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10) Suna</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11) Urbana</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12) Valdagno</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13) Varese</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Chromosome 1

<table>
<thead>
<tr>
<th></th>
<th>41) obt</th>
</tr>
</thead>
<tbody>
<tr>
<td>14) B</td>
<td>42) ri-s se ss k e³ ro</td>
</tr>
<tr>
<td>15) fan</td>
<td>43) ru b st p³ ss e³</td>
</tr>
<tr>
<td>16) NBS</td>
<td>44) ru</td>
</tr>
<tr>
<td>17) ptg</td>
<td>45) ve</td>
</tr>
<tr>
<td>18) sd</td>
<td></td>
</tr>
<tr>
<td>19) w³</td>
<td></td>
</tr>
<tr>
<td>20) w²</td>
<td></td>
</tr>
<tr>
<td>21) w</td>
<td></td>
</tr>
</tbody>
</table>

#### Chromosome 2

<table>
<thead>
<tr>
<th></th>
<th>46) px¹3³ oo; ru jv se st ca</th>
</tr>
</thead>
<tbody>
<tr>
<td>22) a px sp</td>
<td>47) tg (formerly abab¹³⁹)</td>
</tr>
<tr>
<td>23) ab</td>
<td></td>
</tr>
<tr>
<td>24) ast³ dp cl</td>
<td></td>
</tr>
<tr>
<td>25) b cn vg</td>
<td></td>
</tr>
<tr>
<td>26) btl</td>
<td></td>
</tr>
<tr>
<td>27) btl³</td>
<td></td>
</tr>
<tr>
<td>28) bw ba</td>
<td></td>
</tr>
<tr>
<td>29) c wt px</td>
<td></td>
</tr>
<tr>
<td>30) cn</td>
<td></td>
</tr>
<tr>
<td>31) ft</td>
<td></td>
</tr>
<tr>
<td>32) 11²</td>
<td></td>
</tr>
</tbody>
</table>

#### Chromosome 3:

<table>
<thead>
<tr>
<th></th>
<th>58) H/Sb sr In(3R) Mé</th>
</tr>
</thead>
<tbody>
<tr>
<td>33) net</td>
<td>59) ltr/ Sb sr In(3R) Mé</td>
</tr>
<tr>
<td>34) so</td>
<td>60) R Ly/In(3R) P, gm</td>
</tr>
</tbody>
</table>

#### Multichromosomal:

<table>
<thead>
<tr>
<th></th>
<th>61) 1.₄ Cy sp/Pm ; H/Sb sr In(3R) Mé</th>
</tr>
</thead>
<tbody>
<tr>
<td>35) so² b cn</td>
<td>62) Df(2) Px² Df(2) Px, bw sp/SMI, al² Cy sp²</td>
</tr>
<tr>
<td>36) SoC</td>
<td>63) Df(2) bw³ Df(2) bw³ sp²/Xa</td>
</tr>
<tr>
<td>37) spt</td>
<td>64) Df(2) Px Df(2) Px/ Df (2)P ; Df(2;3) P/In (3R) Mo, sr ; w²</td>
</tr>
</tbody>
</table>

#### Deficiencies

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>65) tu A1</td>
<td>66) tu B1</td>
</tr>
<tr>
<td>67) tu B2</td>
<td>68) tu B3</td>
</tr>
<tr>
<td>69) tu B4</td>
<td>70) tu C1</td>
</tr>
<tr>
<td>71) tu C2</td>
<td>72) tu C3</td>
</tr>
<tr>
<td>73) tu C4</td>
<td>74) tu C5</td>
</tr>
<tr>
<td>75) tu D</td>
<td>76) tu Aspra</td>
</tr>
<tr>
<td>77) tu mwh</td>
<td>78) tu Oregon</td>
</tr>
<tr>
<td>79) tu SoC</td>
<td></td>
</tr>
<tr>
<td>80) tu w</td>
<td>81) tu y Hw</td>
</tr>
</tbody>
</table>

### ROMA: Istituto di Genetica Facolta di Scienze DELL Universita

#### Citta Universitaria

Wild Stocks

<table>
<thead>
<tr>
<th>Wild Stocks</th>
<th>3) pn</th>
</tr>
</thead>
<tbody>
<tr>
<td>A 1 Canton - S</td>
<td>4) sc cv v f B / y f : =</td>
</tr>
<tr>
<td>2 Oregon - R</td>
<td>5) sw</td>
</tr>
</tbody>
</table>

**Normal X Chromosome**

| B 1 car bb | 6) w³ |
| | 7) w³ w x |
| 2 n²/109 / In (1) d¹⁴⁹, y Hw m² g² | 8) w³l |
| | 9) w³f |
| | 10) w³f / y f : = |
| | 11) w³o |
January 1962  Melanogaster - Stocks - Italy

36:27

12 wcol
13 wcp
14 wdy / y v f car
15 wQ
16 wsat
17 y ac, sc, pn / y f =
18 y cv v f car
19 y fan sm3
20 y wcol spl f / In (1) rst3, rst3 f
21 y w sat rb
22 y259 / y2 su-wa wB scf
23 y2wcf

Chromosome 2

C 1 b cn c bw
  2 Bl L2 / SM5, al2 Cy ltY sp2
  3 bw
  4 bwD
  5 cn bw
  6 Sb J L2 Pin / SM5, al2 Cy ltY sp2
  7 Cy Bl L / d l
  8 Cy Bl L / Sp Pin

Chromosome 3

D 1 ca K - pn
  2 C1 Sb / L V M
  3 H2 / In (3R) Vno, Vno
  4 ru h th st cu sr es ca
  5 sc ss K es ro
  6 st C J G ca / In (3LR) Ubx130,
    Ubx130 es (1;2;3)
  7 st sr es ro ca

Multichromosomal

E 1 bw; st (2;3)
  2 In (1) AM, y2 / FM6, y3id dm B;
    SM1, al2 Cy sp2 / Bl; In (3R)
    Vno, Vno / In (3LR) Ubx130
    Ubx130 es (1;2;3)
  3 y; suh (1;4)
  4 y; ru h th st pP cu sr es (1;3)
  5 y; bw; st (1;2;3)
  6 y; pol (1;4)
  7 yf = ci eyR (1;4)
  8 lys rc ; (2;3)

Triploid

F 1 y259 / FM4, w f / FM4, w f

Inverted X Chromosomes

G 1 In (1) d1-49, y w tf f
  2 In (1) d1-49, w ls
  3 In (1) d1-49, y Hv m2
  4 In (1) d1-49, y Hv m2g4

5 In (1) sc4L, sc8R, y sc4+6m4
  6 In (1) se AM
  7 In (1) se8 se8
  8 Ins (1) sc8, d1-49, 3C-4EF,
    15DE-20, y3id sc8 dm B (FM6)
  9 In (1) sc8, d1-49, y3id vo f
  10 In (1) w sat , w sat
  11 In (1) y sc+ wa m car
  12 In (1) 481 (12E-F; 14B),
    y bbl 481

Deficiency and Duplications

H 1 Df (1) N8/d1-49, y Hv m2
  2 Df (1) N8/d1-49, y Hv m2 g4
  3 Df (1) N26a-39 wch / FM4, y3id
  4 Df (Y) ybb-

Translocations

I 1 T (1;4) BS (16 A1), y2 cv v
    BS car / y f =
  2 T (1;4) wM5 (3C3), wM5
  3 T (2;3) bwyl4; bwyl4

Closed - X

L 1 Xc, y / y f =

X Chromosomes with a Y Arm Attached

K 1 X yL (C-2), y cv v f car
    bbr, yL
  2 X yL (A-3), sc cv v . yS
  3 ySx (FR-1), yS y cv v f
  4 yS X (P-7), In (1) EN, yS y f

Attached - XY

N 1 X yL xS (108-9 Parker), y2
    su-wa yL, yS
  2 X yS xL (110-8 Parker), y2
    su-wa yS yL, yL y+
  3 X yS, yL (129-16 Parker), y2
    su-wa yS yL, yS+y+
  4 ySx, yL, Ins (1) EN, d1-49,
    yS car f v y . yL

Altered Y

O 1 sc8 y:bw+ (yL bw+ bbyS ac y+)
  2 y sc8 y: y (acyl L, bb + yS)
  3 ySx y+ (BS y+ bb + y+)
  4 ySx yar
  5 y: bw+ (yL bw+ bb+ yS)
  6 y: bw+ (MYR )
  7 y: w+
  8 yY w+
  9 yY y wB
JAPAN

Anzyo, Aichi: Nagoya University, Department of Animal Breeding, Faculty of Agriculture

Wild Stocks

1. Anzyo-Aichi
2. Canton Special
3. Chausuyama-Aichi
4. Hachijojima
5. Hikosan-Kyushu
6. Hiroshima
7. Hita-Kyushu
8. Omogo-Shikoku
9. Oregon
10. Suzuka-Mie
11. Takagicho-Tokyo
12. Yonekawa-Yamaguchi

Chromosome 1

13. Bx
14. ec ct6 g2 bb1/C1B
15. f
16. m
17. m581
18. sc31 B Ins w Sc6 Muller 5
19. sc31 B Ins w Sc8 1(1)59/y w m
20. v
21. w
22. y521
23. y w m
24. y w m f
25. y w m f / y C1B

Chromosome 2

26. b
27. bw

28. bw (from population bell No. 33)
29. cn
30. Cy / bw (M)
31. Cy / bw (T)
32. Cy bw / bw
33. Cy / 1(2)50c
34. Cy bw / 1(2)50c
35. Cy / Pm
36. dp
37. dpV b
38. Pm b
39. Pm / 1(2)50c
40. vg
41. vg'Nw Hia / T(2,3) Sm Cy

Chromosome 3

42. cu
43. e
44. Sb

Multichromosomal

45. Cy / 1(2)50c ; Sb
46. Cy / 1(2)50c ; Sb cu / cu
47. Pm / 1(2)50c ; cu
48. v ; bw

Unanalysed

49. Dichaeote like
50. brown like
51. jaunty like

Kyoto: Kyoto University, Faculty of Science, Department of Zoology

Wild Stocks

South Africa (6)
Sweden (1)
Switzerland (1)
U.S.A. (30)

Common Stocks

Canton-S
Oregon-RS

From different natural populations

Formosa (1)
France (5)
Israel (3)
Italy (2)
Japan (29)
Spain (2)

Chromosome 1

1 B
2 car
3 ec ct6 g2bb1/C1B
4 f
5 Muller-5
6 v
7 w
8 w a
<table>
<thead>
<tr>
<th>Chromosome 2</th>
<th></th>
<th>Chromosome 3</th>
<th></th>
<th>Chromosome 4</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>(w^e)</td>
<td>13</td>
<td>(b)</td>
<td>25</td>
<td>(ca)</td>
</tr>
<tr>
<td>10</td>
<td>(w\ m)</td>
<td>14</td>
<td>(b\ gp)</td>
<td>26</td>
<td>(Confluent-3)</td>
</tr>
<tr>
<td>11</td>
<td>(y)</td>
<td>15</td>
<td>(bw)</td>
<td>27</td>
<td>(cu)</td>
</tr>
<tr>
<td>12</td>
<td>(y\ w\ m\ f)</td>
<td>16</td>
<td>(bw\ (Nanzenji))</td>
<td>28</td>
<td>(e^{11})</td>
</tr>
<tr>
<td></td>
<td></td>
<td>17</td>
<td>(ce)</td>
<td>29</td>
<td>(e^{14}se)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>18</td>
<td>(cn)</td>
<td>30</td>
<td>(ro)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>19</td>
<td>(cn\ bw)</td>
<td>31</td>
<td>(ru\ h\ tu\ st\ cu\ sr\ es\ ca)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>(dp)</td>
<td>32</td>
<td>(se)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>21</td>
<td>(dpx)</td>
<td>33</td>
<td>(se\ (Nanzenji))</td>
</tr>
<tr>
<td></td>
<td></td>
<td>22</td>
<td>(S/Cy\ E-S)</td>
<td>34</td>
<td>(Hn^3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>23</td>
<td>(vg)</td>
<td>35</td>
<td>(ss^a)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>24</td>
<td>(vg^{no})</td>
<td>36</td>
<td>(st)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>37</td>
<td>(wo)</td>
</tr>
</tbody>
</table>

| Multichromosomal |  | 38          | \(svn\) | 39          | \(v\; dp\) |

| Lethal Stocks |  | 48          | \(y; bw; e; ci\ ey^R\) | 49          | \(L-{1}\us-2\) |
|              |  |             |       | 50          | \(L-{1}\us-48\) |
|              |  |             |       | 51          | \(Df(2)px^2/Cy L^1sp^2\) |

| Special Stocks |  | 52          | \(st\; gr\; or\; ro\; ca\) | 53          | \(tu^{35e}\) |
|               |  | 54          | \(tu\; bw\) | 55          | \(tu\; st\) |

| XX with two translocations |  | 56          | \(Y/Basc/T(1;2), T(2;3)\) | 57          | \(Y/wm/T(1;2), T(2;3)\) |

**Mitaka, Tokyo: International Christian University, Biology Department**

**Wild Stocks**

<table>
<thead>
<tr>
<th>Tokyo</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(y, m, ywm^f, w, w^e, w^a)</td>
</tr>
<tr>
<td>2</td>
<td>(bw, vg)</td>
</tr>
<tr>
<td>3</td>
<td>(cu, e)</td>
</tr>
</tbody>
</table>
### KOREA

**Kwangju: Chonnam National University, Department of Biology**

#### Wild Stocks

1. Kwangju
2. Oregon-R
3. Swedish-C

#### Chromosome 1

<table>
<thead>
<tr>
<th>#</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>$l_{2}^{37h}$</td>
</tr>
<tr>
<td>5</td>
<td>$w$</td>
</tr>
<tr>
<td>6</td>
<td>$X^{62} v y_2$</td>
</tr>
<tr>
<td>7</td>
<td>$y$</td>
</tr>
<tr>
<td>8</td>
<td>$y w f$</td>
</tr>
<tr>
<td>9</td>
<td>$Y_{1c/y w Y^S} y v f$</td>
</tr>
<tr>
<td>10</td>
<td>Muller 5</td>
</tr>
</tbody>
</table>

#### Chromosome 2

<table>
<thead>
<tr>
<th>#</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>11</td>
<td>$B_1 L_2/Cy, sp^2$</td>
</tr>
<tr>
<td>12</td>
<td>$B_1 L_2/Cy, bw sp^2$</td>
</tr>
<tr>
<td>13</td>
<td>$cn$</td>
</tr>
<tr>
<td>14</td>
<td>$ds S G b pr/Cy, al^2 I^3 I^4 sp^2$</td>
</tr>
<tr>
<td>15</td>
<td>$l_1 std/SM1, al^2 Cy sp^2$</td>
</tr>
</tbody>
</table>

#### Chromosome 3

<table>
<thead>
<tr>
<th>#</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>16</td>
<td>$M(2)S7/SM5, al^2 Cy I^v sp^2$</td>
</tr>
<tr>
<td>17</td>
<td>$Pin Pfd/Cy, sp^2$</td>
</tr>
<tr>
<td>18</td>
<td>$Pfd/Ins(2L+2R)Cy, S^2$</td>
</tr>
<tr>
<td>19</td>
<td>$pr$</td>
</tr>
<tr>
<td>20</td>
<td>$Sp Bl L/Cy, sp^2$</td>
</tr>
<tr>
<td>21</td>
<td>$Sp bwD/SM5, al^2 Cy I^v sp^2$</td>
</tr>
<tr>
<td>22</td>
<td>$vg$</td>
</tr>
</tbody>
</table>

### Seoul: Chung-Ang University, College of Liberal Arts & Sciences, Department of Biology

#### Wild Stocks

1. Oregon-R
2. Urbana-S
3. Kongju (3 strains)
4. Seoul

#### Chromosome 1

<table>
<thead>
<tr>
<th>#</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$w$</td>
</tr>
<tr>
<td>2</td>
<td>$y$</td>
</tr>
<tr>
<td>3</td>
<td>$Y_{1c/y w Y^S} y v f$</td>
</tr>
</tbody>
</table>

#### Chromosome 2

<table>
<thead>
<tr>
<th>#</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>$B_1 L_2/Cy, bw sp^2$</td>
</tr>
<tr>
<td>5</td>
<td>$cn$</td>
</tr>
<tr>
<td>6</td>
<td>$j$</td>
</tr>
<tr>
<td>7</td>
<td>$vg$</td>
</tr>
</tbody>
</table>

#### Inversion

- Muller-5

### Seoul: Seoul National University, Department of Zoology

#### Wild Stocks

1. Canton-S
2. Kongjoo (Korea)
3. Oregon-R
4. Quilpart (Korea)
5. Seoul (Korea)
6. Swedish-C
7. Urbana-S

#### Chromosome 1

<table>
<thead>
<tr>
<th>#</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>BB</td>
</tr>
<tr>
<td>2</td>
<td>Bx</td>
</tr>
</tbody>
</table>

#### Chromosome 2

<table>
<thead>
<tr>
<th>#</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>$Bll_Cy$</td>
</tr>
<tr>
<td>4</td>
<td>$c$</td>
</tr>
<tr>
<td>5</td>
<td>$cn$</td>
</tr>
<tr>
<td>6</td>
<td>$J$</td>
</tr>
<tr>
<td>7</td>
<td>$vg$</td>
</tr>
</tbody>
</table>
January 1962  Melanogaster - Stocks - Korea  36:31

Chromosome 3  
ru  ca  Multichromosomal
ca  e se  
e11 se  Inversion
ro  Muller-5

Seoul: Sung Kyun-Kwan University, Department of Biology

Wild Stocks  
1) Oregon-R  
2) Seoul (Korea)  
3) Swedish-C  

Chromosome 3  
11) ca  
12) ro  
13) ru cu ca  
14) se h  
15) ve h th

Chromosome 1  
4) BB  
5) w  
6) y

Chromosome 4  
16) pol

Chromosome 2

Inversion  
7) c  
8) cn  
9) j  
10) pfd/Sm-5

Seoul: Yonsei University, Department of Biology

Wild Stocks  
1) Canton-S  
2) Oregon-R --- Isogenic  
3) Oregon-R-C --- Isogenic  
4) Oregon-S  
5) Samarkant (Japan)  
6) Seoul-1 (Korea) --- Isogenic  
7) Seoul-2 (Korea) --- Isogenic  
8) Suwon (Korea)  
9) Swedish-C  
10) Yangdong (Korea)

Chromosome 1  
11) B  
12) bi ct6 q2  
13) bo  
14) br  
15) Bx3  
16) cm  
17) ec  
18) ec dx  
19) fa  
20) rg  
21) sc cv v f

22) sc_cv v eq  
23) sn3  
24) svr  
25) t  
26) t2 v f  
27) v  
28) v car  
29) w  
30) wa  
31) wbf2  
32) wch  
33) wco sh2  
34) wcol  
35) w0 tbl/CLB  
36) y  
37) y ac v  
38) y sc mf2  
39) y2 cv v f  
40) M-5/y sc6 y  
41) M-5/y ac Sn3 cn  

Chromosome 2

42) a px or  
43) a px sp  
44) ab  
45) al
<table>
<thead>
<tr>
<th>Chromosome 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>95 ss</td>
</tr>
<tr>
<td>96 st</td>
</tr>
<tr>
<td>97 th</td>
</tr>
<tr>
<td>98 th st cp</td>
</tr>
</tbody>
</table>

**Chromosome 4**

<table>
<thead>
<tr>
<th>46 al bc sp²</th>
</tr>
</thead>
<tbody>
<tr>
<td>47 al b pr an vg a sp²/ In(2LR)Cy, L¹ sp²</td>
</tr>
<tr>
<td>48 al dp b pr blt bw/ SM5, a¹ 2 Cy ltv sp²</td>
</tr>
<tr>
<td>49 al dp b pr c px sp/Cy, pr</td>
</tr>
<tr>
<td>50 al dp b pr Bl c px sp/ SM¹ a² Cy sp²</td>
</tr>
</tbody>
</table>

**Chromosome 3**

| 51 b |
| 52 b lt wxt bw |
| 53 b vg |
| 54 bw |
| 55 bw ba |
| 56 Bl/Cy, bw¹⁵a sp² or¹⁵a |

**Multi-chromosomal Stocks**

| 99 bt |
| 100 ci |
| 101 ci gvl bt |
| 102 ey |
| 103 ci gvl eg²sv¹ |
| 104 pol |
| 105 spa |

**Attached-X**

| 118 br ec/y³d |
| 119 y/g² ty |

**Deficiencies**

| 120 Df(3)sbd¹⁰⁵/Xa |
| 121 Df(2)al Cy, En-S |

**Duplications**

| 122 Lp(2;3)S |

**Inversions**

| 123 Muller-5 |
| 124 Ins(1)s¹⁵¹ |

**Translocations**

| 125 Vg¹⁵/H¹a/SM⁵, a¹ ² Cy ltv sp² |
| 126 Vg¹⁵/R¹d, bw sp or |
| 127 A/¹n(3R) hp hp |

| 128 T(1;2) B¹d/C¹R |
| 129 T(2;3) Xa/Sb bx³ |
### South Africa

**Johannesburg:** University of the Witwatersrand, Department of Zoology

#### Wild Stocks

<table>
<thead>
<tr>
<th>Location</th>
<th>Chromosome 1</th>
<th>Chromosome 2</th>
<th>Multichromosomal</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Bethulie</td>
<td>rb cm g³</td>
<td>101 c px</td>
<td>142 bw; e; cl ey</td>
</tr>
<tr>
<td>2 Bloemfontein</td>
<td>rb cm car</td>
<td>102 cn</td>
<td>143 bw; cl ey</td>
</tr>
<tr>
<td>3 Canton-S</td>
<td>rb cx</td>
<td>103 cn35k</td>
<td>144 bw; st</td>
</tr>
<tr>
<td>4 Cape Town</td>
<td>rb g³</td>
<td>104 cn vg</td>
<td>145 Cy/Pa, ds33k;H/In(3R)</td>
</tr>
<tr>
<td>5 Cedara</td>
<td>sc</td>
<td>105 cl</td>
<td>Moist; sr</td>
</tr>
<tr>
<td>6 Drakensberg</td>
<td>sc ec cv ct³ v</td>
<td>106 cl50a</td>
<td>146 g³; bw</td>
</tr>
<tr>
<td>7 Florida</td>
<td>sc ec cv ct³ v g²f</td>
<td>107 dke c</td>
<td>147 g³; st</td>
</tr>
<tr>
<td>8 Graaff-Reinet</td>
<td>svr w³</td>
<td>108 dp</td>
<td>148 g³; st p³</td>
</tr>
<tr>
<td>9 Inhaca Island</td>
<td>v³6f</td>
<td>109 lt std/Cy sp²</td>
<td></td>
</tr>
<tr>
<td>10 Johannesburg</td>
<td>v w³</td>
<td>110 lt stwJ</td>
<td></td>
</tr>
<tr>
<td>11 Kalahari</td>
<td>w</td>
<td>111 ltd</td>
<td></td>
</tr>
<tr>
<td>12 Kariba Dam</td>
<td>w m</td>
<td>112 pd</td>
<td></td>
</tr>
<tr>
<td>13 Limpopo</td>
<td>w m f</td>
<td>113 pr</td>
<td></td>
</tr>
<tr>
<td>14 Nelspruit</td>
<td>w³</td>
<td>114 pr42d</td>
<td></td>
</tr>
<tr>
<td>15 Nyasa Lake</td>
<td>w³³</td>
<td>115 px</td>
<td></td>
</tr>
<tr>
<td>16 Oregon-R</td>
<td>w³³</td>
<td>116 sf²</td>
<td></td>
</tr>
<tr>
<td>17 Stanford Lake</td>
<td>w³³ rb</td>
<td>117 sp</td>
<td></td>
</tr>
<tr>
<td>18 Stellenbosch</td>
<td>w³³ l³</td>
<td>118 Su-H/Cy, pr</td>
<td></td>
</tr>
<tr>
<td>19 Tzaneen</td>
<td>w³³ c³</td>
<td>119 tk sf² abb</td>
<td></td>
</tr>
<tr>
<td>20 Umgazi River</td>
<td>w³³ c³</td>
<td>120 vg</td>
<td></td>
</tr>
<tr>
<td>21 West Rand</td>
<td>w³³ c³</td>
<td>121 vgdn</td>
<td></td>
</tr>
<tr>
<td>22 Western Province</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>23 Zoutpansberg</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Chromosome 1

<table>
<thead>
<tr>
<th>Location</th>
<th>Chromosome 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 bi ct³ g²</td>
<td></td>
</tr>
<tr>
<td>25 bo</td>
<td></td>
</tr>
<tr>
<td>26 B</td>
<td></td>
</tr>
<tr>
<td>27 car</td>
<td></td>
</tr>
<tr>
<td>28 car²</td>
<td></td>
</tr>
<tr>
<td>29 cm</td>
<td></td>
</tr>
<tr>
<td>30 cm g³ car</td>
<td></td>
</tr>
<tr>
<td>31 cm³ ct v</td>
<td></td>
</tr>
<tr>
<td>32 ct v dy g</td>
<td></td>
</tr>
<tr>
<td>34 ct³</td>
<td></td>
</tr>
<tr>
<td>35 cv ct</td>
<td></td>
</tr>
<tr>
<td>36 cv sc</td>
<td></td>
</tr>
<tr>
<td>37 ec</td>
<td></td>
</tr>
<tr>
<td>38 ec ct³ v g³</td>
<td></td>
</tr>
<tr>
<td>39 f²</td>
<td></td>
</tr>
<tr>
<td>40 f³ m</td>
<td></td>
</tr>
<tr>
<td>41 f³ v</td>
<td></td>
</tr>
<tr>
<td>42 g³</td>
<td></td>
</tr>
<tr>
<td>43 g³³</td>
<td></td>
</tr>
<tr>
<td>44 g³</td>
<td></td>
</tr>
<tr>
<td>45 m</td>
<td></td>
</tr>
<tr>
<td>46 pn²</td>
<td></td>
</tr>
<tr>
<td>47 ras dy</td>
<td></td>
</tr>
<tr>
<td>48 ras²</td>
<td></td>
</tr>
<tr>
<td>49 rb</td>
<td></td>
</tr>
<tr>
<td>50 rb car</td>
<td></td>
</tr>
</tbody>
</table>

#### Chromosome 2

<table>
<thead>
<tr>
<th>Location</th>
<th>Chromosome 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>91 albasp/Cy L⁴ sp²</td>
<td></td>
</tr>
<tr>
<td>92 al dp b pr Bl c px</td>
<td></td>
</tr>
<tr>
<td>93 a sp²</td>
<td></td>
</tr>
<tr>
<td>94 b</td>
<td></td>
</tr>
<tr>
<td>95 b pr cn</td>
<td></td>
</tr>
<tr>
<td>96 b pr cn a</td>
<td></td>
</tr>
<tr>
<td>97 bw</td>
<td></td>
</tr>
<tr>
<td>98 bw²b</td>
<td></td>
</tr>
<tr>
<td>99 bw¹ d</td>
<td></td>
</tr>
<tr>
<td>100 c</td>
<td></td>
</tr>
</tbody>
</table>

#### Multichromosomal

<table>
<thead>
<tr>
<th>Location</th>
<th>Multichromosomal</th>
</tr>
</thead>
<tbody>
<tr>
<td>142 bw; e; cl ey</td>
<td></td>
</tr>
<tr>
<td>143 bw; cl ey</td>
<td></td>
</tr>
<tr>
<td>144 bw; st</td>
<td></td>
</tr>
<tr>
<td>145 Cy/Pa, ds33k;H/In(3R)</td>
<td>Moist; sr</td>
</tr>
</tbody>
</table>
Melanogaster - Stocks - South Africa


eras²; st
150 rb; bw
151 rb; ry
152 rb; se
153 rb; st
154 car; ry
155 car; se
156 vg; se
157 w² rb; se
158 w²; cd
159 y; bw; e; ci ey

Attached-X
160 f B/ su²-v-pr v
161 Y¹+/²
162 Y² su-w a w² bb/
sc⁴L sc⁸R

Inversions

163 In(1)A99b
164 In(1)d1-49, y fa^n

Translocations

165 In(1)rst³, y rst³
car bb
166 In(1)w^n⁴
car bb
167 In(1)w^n⁴; st
168 Ins(1)sc⁵L, S, sc⁸L
w² B sc⁸

In(1)dl-49, Y fan

Chromosome 2
bw
cn
cn bw
vg

Chromosome 3
D/Gl
e
se

Attached-X

/y

Multichromosomal
bw; st
Cy/Pm; H/In(3R)
T(1;3)⁰⁴; D/G1B
y bw ci e ey

SWEDEN

Uppsala 7: University of Uppsala, Institute of Genetics

Wild Stocks

1. Algeria
2. Amherst-3
3. Bayfordbury
4. Boa Esperanca, Minas Gerais, Brazil
5. Canton-S
6. Crimea
7. Curitiba
8. Florida
9. Formosa
10. Gruta, Argentina
11. Hikone-R (resistant to BHC, DDT, parathione, nicotine)
12. Karsnäs
13. Kochi-R (resistant to parathione)
14. Oregon-R
15. Salvador, Bahia, Brazil
16. Samarkand
17. San Miguel, Buenos Aires, Argentina
18. Stäket
19. Tunnelgatan
20. Ultuna
21. Örebro
Chromosome 1 (X)

101. B
102. B/y:=
103. BB car; sc8 Y/y f:= ; sc8 Y
104. B
105. ct
106. cv
107. cv sn3
108. ec
109. ec ct v g
110. ec ct v f
111. f
112. f B od car/ y f:=
113. f BB; sc8 Y/ y f:= ; sc8 Y
114. f B1 B1/ y f:=
115. f od sy car
116. fu ff/ GlB
117. g (Sweden)
118. g2
119. In (1) w4
120. la/ GlB
121. ma-1/ y f:=
122. od car
123. sc z w7G2 ec/ y w f:=
124. sc z ec
125. sc sc mottled
126. sc sc B InS wA sc8 (Muller-5)
127. sc sc InS wA sc8
128. sn3
129. sp-w
130. sp-w2
131. su-wa wa
132. v g
133. w
134. w cv
135. w cv sn3
136. w sn3
137. wa su-f/ y f:=
138. wa4/ y f:=
139. wbf f5
140. wbf2
141. wbl
142. wBwx
143. wch wy/ y f:=
144. wco
145. wco sn2
146. we
147. we2
148. we2 en-w6/ y f:=
149. wh
150. wh ct
151. wi yb
152. wst
153. y
154. y ec ct v f
155. y rest3 car
156. y f Eb/ sc S1 B InS wA sc8
157. y2 sc wA wch fa/ y w f:=
158. y2 sc wA
159. y2 sc wA wch/ y w f:=
160. y2 su-wA wA2 wch spl/ y f:=
161. y2 wa
162. y2 wa w
163. y2 wa sc
164. y2 wbf spl sn3/ y f:=
165. z
166. z wA/ y w f:=
167. z wA3/ y w f:=
168. z wA Bf

Chromosome 2

201. bw
202. bwD
203. Cy/ Pm
204. Cy/ S
205. feq Alu 1/ al2 Cy 1t3
206. nw2/ Cy RNS
207. pr
208. S/ NS, px sp
209. vg
210. vg bw

Chromosome 3

301. ca
302. cd
303. D3/ InP
304. D/ Stb
305. e11
306. kar
307. ri ss
308. ri2
309. ri2 ss
310. ro
311. ru h et pP ss ss ss
312. ry
313. ry2
314. ry2 cd
315. se
316. ss
317. st
318. st p
319. st ry
320. st ss e11

Chromosome 4

401. ey
402. svn

Multichromosomal

501. wch; Su-wch/ Cy (1;2)
502. wcol; bw (1;2)
503. we: cr-w/ Cy 
504. yS1 sc8 InS y3P; al2 Cy 
   lt3 sp2/ dp b Pm1; 
   ru h D3 InCXF ca/ Sb 
   In (3R) (1;2;3) 
505. y' w spl; Cy; Ubx130/ Xa 
506. bw; cd (2;3) 
507. bw; st (2;3) 
508. Cy/ S; D/ InP (2;3) 
509. L sp; th (2;3) 
510. L2/+; sp; th (2;3) 
511. sp; th (2;3) 

Definitions and Duplications

601. sc z Df(1)w258-45/ FM4 
602. y2 Df(1)w258-45/ FM4 

603. XY', w2 R; Y'/ y / Y' (Stern) 
604. Df(1)w258-45; y w spl dm; 
   Dp(1;3)wco/ y w f:= 
605. Dp(1)is/ y f:= 
606. Dp(1;2R)w5107 
607. Dp(1;2R)w51c20 
608. Dp(1)w2/ y w fi:= 
609. Dp(wa2/ w4)/ y f:= 
610. Dp(wa2/ w4) ec/ y f:= 
611. sc Dp(1)259d15/ y f:= 
612. z Dp(wa2/ w4)/ y f:= 

Translocations

701. T(1;4) w5 
702. T(2;3) bwYDe4/ Cy
Report of A. Chovnick and A. Schalet

A large number of X-ray induced rosy mutants are under study in this laboratory (see notice DIS 34, page 122). Some of these mutants have been reported in prior notes (Schalet and Chovnick, DIS 34; Chovnick, Schalet and Kernaghan, 1961), and others will be discussed in future reports. We are using the designations ry⁴ through ry⁷³ for existing mutants. In view of Hubby's report of a new mutant, ry³ (DIS 35), we have changed the designation of our ry³ to ry³a.

Report of K. S. Gill


Report of E. H. Grell

nw³: narrow-Dominant. E. H. Grell, 59f. 2-83⁻. From an X-rayed Canton-S male. The wings tend to be longer and more narrow than normal. The phenotype is highly variable and except for slightly squared wing tips sometimes approaches wild type. The viability of heterozygotes (nw³/+ ) is below normal and homozygotes are entirely inviable; nw³/nw is also lethal. RK2.
Report of Afton M. Hansen and Eldon J. Gardner

**scrap** Hansen and Gardner 1960. 2-747. Pub. Hansen and Gardner, 1961, Genetics 46:1809. Appeared in a "wild" Cockaponsett stock that had been subjected for several days of high temperature. This stock was apparently homozygous for the **scrap** gene at the time the phenotype was detected. Expressivity is variable. Usually the ventral one-third of the eye is flattened and differentiated from the dorsal two-thirds by a horizontal depression. Tufts of vibrissae are often present on the anteroventral border of the eye. Occasionally the eyes are reduced. Growths may be present. Either or both eyes are affected. The ommatidia are shorter in the area of depression. The expression completely overlaps the wild-type at 25°C, with an average penetrance of 80 per cent at 30°C. Temperature effective period at 30°C is from the forty-second to sixty-eighth hour of development, with the entire period at 30°C necessary for maximum penetrance. Viability good. RK4.

Report of T. Imaizumi

**Cf-3** : Confluent-3 60j. 3-66-2. Dominant, homozygous lethal; arose spontaneously as I male in the cross of Oregon-RS x b vg, an allele of Delta, phenotype of heterozygote very likely to that of Delta, good viability; homozygote die at late stages of embryo and partially at early larval stage. Perhaps with deficiency.

Report of P. T. Ives

**61e** : miniature61e Ives, 61e23. Like m. Induced by 1 kr γ radiation in an Oregon-R/rucuca sperm which was deposited on day 4 of an exhaustive mating schedule.

Functional allelism to m established by A. B. Burdick, who is also investigating its pseudoallelic properties.

Report of P. T. Ives

**60k** : ebony60k. Ives, 60k25. Like e. Spontaneous in vg; se50k.

Report of Shanta V. Iyenyar

**ballet** : ballet, X-chromosome, not localised; X-ray induced in young male, recessive, wings one third the normal length stretched outwards and slightly upwards, wing tip broadened, venation markedly altered as in fused; viability of the male impaired but not to the same extent as females since only one female homozygous for ballet has been found but died one hour after emergence. Mutation does not seem to be affected by temperature.

**narrow wing-rough eyes** , X-chromosome not localised, X-ray induced. Wings like tapered on II chromosome, eyes smaller, oval in outline and uniformly more rough. Both sexes have been noticed, recessive, does not seem to be affected by temperature.
Report of E. B. Lewis

Correction to DIS 34:51. The mutant alpha-1 of H. W. Lewis and its description should appear under a separate heading entitled "Report of H. W. Lewis."

Report of Y. Maeda

uex: unexpanded Maeda, Y., 5813. 2-. Spontaneous in local stock (Kobe) of wild. Wings are wrinkled and crumpled as those of newly emerged flies, about 1/2 length of normal size, like pu, often inflated as balloon. Tibia and tarsi in 3rd legs are irregularly shortened and gnarled, so the mode of creeping is abnormal. Posterior scutellas bent toward median. The data from the preliminary linkage analysis by crossing with cn or bw indicated that the approximated position of 2-55+ for uex (+ 724, cn 27, uex 14 and cn uex 663 from cn uex/+ x cn uex, and + 535, bw 367, uex 233 and uex bw 258 from uex bw/+ x uex bw). Male viability is somewhat low. RK2.

Report of E. Ortiz

w57b: white57b Ortiz, 57b. From Canton-S wild type male, treated with ethyl-urethane, as 8 males. Behaves as an allele of our stock of w. Eye colour snow white. Good viability and fertility. RK1.

w58a: white58a Ortiz, 58a. Spontaneous as 14 males in our laboratory stock of vg. Behaves as an allele of our stock of w. Eye colour snow white. Good viability and fertility. Kept in stock as w58a; vg. RK1.

d61g: lozenge61g Ortiz, 61g. From Oregon-R wild type male, X-ray induced, as 22 males. Eye oval, smaller than wild type, all facets run together into smooth surface. Eye colour dark brown, with darker rim. Homozygous female highly infertile. RK2.

Report of Verena Rohr

rS: rudimentary Swiss Hadorn, 59d. Spontaneous in cross +/+ x ma-1/. Wings obliquely truncated, often arclike and blistered with medial and lateral marginal bristles sparse and ruffled. Often more than one bristle on one socket. L4 and L5 shortened. Males fertile, females sterile. Wing size much more reduced in homozygous females than in hemizygous males. Expressivity similar to r39k, but with great variability. rS/ male offspring from different crosses much influenced by genotypic milieu. Compounds rS/r39k, rS/R9 semilethal with abnormal marginal wing bristles. Compound rS/r12 subvital, some individuals with normal lateral marginal wing bristles. RK2.
D. MELANOGASTER

**LINKAGE DATA**

**Report of M. M. Green:** Gene sequence at the X chromosome tip in *Drosophila melanogaster*.

From a cross of ♀ y² sc w³ ec/y² su-w³ a¹ X +♂ a a single y⁺ sc⁺ w⁺ a a was found. Subsequent crosses showed that the a a was y² sc w³ and carried a short duplication of the X tip inserted in chromosome 3. This duplication in all probability arose as a spontaneous X-3 translocation in a parental +♂ of which only the X part translocated to chromosome 3 was found. In addition to y² and sc, the duplication covers su-w³ but not dor, thereby establishing the probable order as y², sc, su-w³, dor. Inadvertently this duplication was lost.

**Report of Afton M. Hansen and Eldon J. Gardner**

Scarp-eyed (scrp) flies were mated with curly, plum, stubble, dichaete flies. All individuals in the F₁ generation had wild-type eyes. Flies from this mating with curly wings and stubble bristles were then back-crossed with scrp flies and their progeny allowed to develop at 30°C. The progeny with scarp eyes had either stubble bristles or wild-type bristles and wild-type wings. Scarp eyes did not occur in combination with curly wings. Thus, scrp was located in the second linkage group.

Scarp-eyed flies were mated with aristateless, dumpy, black, purple, curved, plexus, speck flies and the wild-type F₁ females were back-crossed with al + dp + pr c px sp males. Flies of each of the 12 possible single crossover gametes were collected and mated with scrp males or females. Their progeny developed at 30°C. When scarp-eyed flies appeared in the progeny, the parent was scored as a carrier of scrp. A summary of data from two experiments is included in the following table:

<table>
<thead>
<tr>
<th>Genotype tested</th>
<th>Number of successful matings</th>
<th>Number that were carriers of scrp</th>
</tr>
</thead>
<tbody>
<tr>
<td>+ dp b pr c px sp</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>al ++ ++ ++ ++</td>
<td>17</td>
<td>17</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ++ b pr c px sp</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>al dp + ++ ++ ++</td>
<td>16</td>
<td>15⁺</td>
</tr>
<tr>
<td>al dp + pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ++ pr c px sp</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>al dp b + ++ ++</td>
<td>13</td>
<td>13</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ++ ++ + c px sp</td>
<td>53</td>
<td>2</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
(Hansen and Gardner, table--continued)

<table>
<thead>
<tr>
<th>Genotype tested</th>
<th>Number of successful matings</th>
<th>Number that were carriers of scrp</th>
</tr>
</thead>
<tbody>
<tr>
<td>al dp b pr + + +</td>
<td>23</td>
<td>22</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ + + + px sp</td>
<td>19</td>
<td>16\textsuperscript{b}</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>al dp b pr c + +</td>
<td>30</td>
<td>0</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ + + + + sp</td>
<td>13</td>
<td>11\textsuperscript{c}</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ + + + + +</td>
<td>13</td>
<td>12\textsuperscript{d}</td>
</tr>
<tr>
<td>al dp b pr c px sp</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{a} only 5 offspring  
\textsuperscript{b} only 8, 5, and 3 offspring 
\textsuperscript{c} only 2 and 19 offspring  
\textsuperscript{d} only 14 offspring

These data showed that scrp was located near \( c \), but to the left. Three of 76 crossovers between \( pr \) and \( c \) (21 map units) were also crossovers between \( c \) and scrp. Crosses with lobe-recessive showed that scrp is not an allele of \( L^r \). Scrp is located at approximately 74 units from the left end of the second chromosome.
AMHERST, MASSACHUSETTS: AMHERST COLLEGE

Correction to list in DIS 34:

D. simulans: Amherst, Mass. 1961

BALTIMORE, MARYLAND: THE JOHNS HOPKINS UNIVERSITY

D. funebris
D. hydei
D. simulans
D. simulans, La-3
D. simulans, La-4
D. simulans, Lima, Peru, a5 D. simulans, Lima, Peru, a10
D. simulans, New Orleans
D. simulans, South Africa
D. virilis

CHICAGO, ILLINOIS: UNIVERSITY OF CHICAGO

Department of Zoology

Stocks listed in DIS 34 with the following numbers are no longer kept in culture: 4, 37, and 39.

D. virilis

DeKALB, ILLINOIS: NORTHERN ILLINOIS UNIVERSITY

Department of Biological Sciences

Stock Lists, other species: Essentially as in DIS 34, but have added:

D. tripunctata - several local samples
D. immigrans - several local samples

LINCOLN, NEBRASKA: THE UNIVERSITY OF NEBRASKA

Zoology Department

D. affinis: Alabama, Georgia, Florida, Iowa, Kentucky, Louisiana, Minnesota, Mississippi, Missouri, Nebraska, New York, North Carolina, South Carolina, Tennessee, and Texas. All wild-type strains except for several with CO2 sensitivity (virus-caused).

D. algonquin: Massachusetts and Minnesota.

D. athabasca: Massachusetts, Maine, Alaska, Minnesota.


LOS ANGELES, CALIFORNIA: UNIVERSITY OF CALIFORNIA

Department of Botany

D. pseudoobscura

Lethal strains: A number of lethal strains of various gene arrangements of the third chromosome from Southern California and Guatemala, which are currently being tested. Lethal strains are currently being established from wild samples from Bogota, Colombia.
Wild strains (homozygous and isogenic for Chromosome 3 inversions):

Standard (1) San Jacinto Mountains
Arrowhead (1) San Jacinto Mountains
Arrowhead (1) Texas
Chiricahua (2) San Jacinto Mountains
Tree Line (1) San Jacinto Mountains
Tree Line (4) Guatemala
Pikes Peak (2) San Jacinto Mountains
Pikes Peak (1) Texas
Santa Cruz (2) San Jacinto Mountains
Santa Cruz (1) Guatemala
Oaxaca (2) Guatemala
Cuernavaca (3) Guatemala
Vandeventer (2) San Jacinto Mountains
Thomas Mountain (1) San Jacinto Mountains
Pinyon (1) San Jacinto Mountains (carries lethal factor, maintained heterozygously balanced with marked Standard chromosome)

Chromosome 1

Cnv (Convergent veins)
Pt w^s
Pt mg^2
y sn v co sh
sd (scalloped)
upd
er (erect bristles)

Chromosome 2

g_1^2
pcv^1
ubx
ga
upt gl
upt bx Ba gl (In) 1/1
upt bx Ba gl (In) 1/DL ubx gl2 bv
pcv^1 ubx cd gl2^2 bv

Chromosome 3

or (Standard)
or (Santa Cruz)
pr (Standard)
pr (Arrowhead)
or pr (Standard)
or pr (Santa Cruz)
or pr cv (Standard)
or pr spr (Standard)
or pr spr (Arrowhead)
or px (Arrowhead)
or px pr (Arrowhead)
or ru (Tree Line)
or pr vg (Arrowhead)
or Bl L pr cv (AR) / lethal (CU)
or Bl L Sc pr cv (ST) / lethal (CU)
or Bl px^D (AR) / or L (SC)
or Bl Sc ru pr cv (ST) / or L (SC)
or L Sc pr (ST) / lethal (CU)
or Bl px^D (AR) / lethal (CU)
or Bl L Sc (VA) / lethal (CU)

Chromosome 4

inc j hk
tg^c

Multichromosomal

D. hydei San Jacinto Mountains
D. tolteca Guatemala
D. biopaca Guatemala

NEW HAVEN, CONNECTICUT: YALE UNIVERSITY
Department of Zoology

D. americana americana: Independence, Ohio; Western
D. americana texana: Florida
D. ananassae: Cristobal
D. bifasciata: Pavia, normal and sex-ratio
D. busckii: Abingdon, Pa.
D. equinoxialis: Puerto Rico, normal and sex-ratio
D. flavomontana: Yampa River, Colo.
D. funebris: Rexburg, Idaho; Stockholm, Sweden; Upperville, Va.; Yucatan; white
D. giberosa: South Mexico
D. hydei: New Haven, Conn.; Zurich, Switzerland
D. laicola: Fairbanks, Minn.
D. littoralis: Switzerland
D. melanica
D. montana: Cottonwood Canyon, Utah; LU
D. nebulosa: Haiti, normal and sex-ratio
D. nigromelanica
D. novamexicana
D. paramelanica
D. paulistorum: Belem; Bucamaranga; Cantareiras; Lancetilla; Trinidad
D. pseudoobscura: Pinon Standard
D. simulans
D. virilis: Japan
D. willistoni: Barbados-3; Belem; Recife-3; Recife-6; Recife Pop. 168; ebony; pink;
white eye; sex-ratio
Zaprionis vittiger: South Africa

PHILADELPHIA, PENNSYLVANIA: WOMAN'S MEDICAL COLLEGE

D. robusta

A. Homozygous stocks (in addition to those listed in DIS-34):

<table>
<thead>
<tr>
<th>Stock designation</th>
<th>X</th>
<th>2</th>
<th>3</th>
<th>Origin of wild strain</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPK 22A7</td>
<td>11</td>
<td>SS</td>
<td>SS</td>
<td>Pokagon State Park,</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Steuben County, Indiana</td>
</tr>
<tr>
<td>IPK 61C5</td>
<td>11</td>
<td>SS</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>IPK 108B2</td>
<td>S1</td>
<td>3S</td>
<td>S1</td>
<td></td>
</tr>
<tr>
<td>IPK 108B6E</td>
<td>S1</td>
<td>3S</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>IPK 117B7D</td>
<td>11</td>
<td>3S</td>
<td>S1</td>
<td></td>
</tr>
<tr>
<td>IPK 122G1</td>
<td>S1</td>
<td>3S</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>IPK 122K3C</td>
<td>12</td>
<td>3S</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>IPK 209D2D</td>
<td>S1</td>
<td>SS</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>IPK 227F</td>
<td>11</td>
<td>S1</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>OBH 158A1</td>
<td>12</td>
<td>SS</td>
<td>SS</td>
<td>Broadview Heights,</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Cuyahoga County, Ohio</td>
</tr>
<tr>
<td>OBH 171B1D</td>
<td>11</td>
<td>SS</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>OBH 178C,D</td>
<td>1S</td>
<td>SS</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>OW 75C2,5</td>
<td>SS</td>
<td>SS</td>
<td>SS</td>
<td>Woorster, Wayne County, Ohio</td>
</tr>
<tr>
<td>OW 92R2A</td>
<td>12</td>
<td>SS</td>
<td>SS</td>
<td></td>
</tr>
<tr>
<td>OW 101E5</td>
<td>1S</td>
<td>SS</td>
<td>SS</td>
<td></td>
</tr>
</tbody>
</table>

*Following notation in Carson, H. L., 1953, Genetics 38:168.

B. Wild strains (not inbred): Ten or more each from Alabama, Indiana, Ohio,
Mississippi, and South Carolina.
D. arizonensis

4 strains from Tucson, Arizona:

50 isofemale strains collected October and November, 1961:

- Tucson, Arizona (2)
- Catalina Mountains (8,000 ft.), Arizona (1)
- Patagonia, Arizona (6)
- Magdalena, Sonora, Mexico (29)
- Cornelio, Sonora, Mexico (1)
- Hermosillo, Sonora, Mexico (10)
- Desembogue, Sonora, Mexico (1)

Mutant strains:

- White eye - chromosome 1
- Lobed eye - chromosome 3

D. psuedoobscura

Isofemale strains:

- Bryce, Utah (8)
- Ferron, Utah (6)
- Gunnison, Colo. (6)
- Lemon Cave (9)
- Mather, Calif. (8)
- Mono, Calif. (7)

(74 inbred lines from the above strains)

New isofemale strains:

- Patagonia, Arizona (1) 1961.
- San Felipe, B. Calif., Mexico (1) 1961
- Chiriaco Summit, California (2) 1961

Other species

- D. aldrichi - Hermosillo, Sonora, Mexico (1961)
- D. americana
- D. hamatofila
- D. hydei
- D. longicornis
- D. mojavensis - Chocolate Mts., California
- D. mulleri - Austin, Texas
- D. mulleri - (G-207 - 4th chromosome inversion)
- D. persimilis
- D. putrida - Raleigh, N. C.
- D. rita
- D. simulans
- D. texana
- D. virilis (3 strains)
AUSTRALIA

Sydney, New South Wales: The University of Sydney

Addition to list in DIS 34:62:

"D. husckii": Sydney; Melbourne

BRAZIL

Pôrto Alegre: Universidade do Rio Grande do Sul, Instituto de Ciencias Naturais

D. willistoni

Wild strains from: Praia do Leste (Paraná), Ilha das Cobras (Paraná), Col. São Pedro (Rio Grande do Sul), Itapeva (Rio Grande do Sul), Manaus (Amazonas), Tubarão (Sta. Catarina), Maranhuapé.

Chromosome I

w^ey sn In ru (analyzer stock)
yellow
w'
sepia
w^elz

Chromosome II

abb bw (analyzer stock)
SHkabbbw (In)/lethal (analyzer stock)
abbpx bw
Em abbww/abbbw
20 wing eye and other mutants from natural populations unirradiated or irradiated with Co60.

Chromosome III

pink (analyzer stock)
Δ pink (In)/lethal (analyzer stock)
ebony
ebony mixed with other mutant
30 other of the same origin as those listed for Chromosome II.
Several "semilethals" and "normal" viable strains, homozygous for the second and third chromosomes.

Other Species

D. ananassae - Cassarongongo (Bahia), Tabatinga (Amazonas), Sacavem, Boa Viagem (Maranhão), Belém (Pará).
D. bandeirantorum - Itatiaia (Rio).
D. bocainensis - Praia do Leste, Ilha das Cobras (Paraná), Pitanga (Bahia), Eldorado (R.G.Sul), Jacú (Sta. Catarina), Belém (Pará).
D. capricorni - Itatiaia (Rio), Eldorado (R.G.Sul), Ilha das Cobras (Paraná).
D. cardini - Costa Rica H 15-1, Panamá H 7918, Belém (Pará), Eldorado (R.G.Sul), Pedras (Bahia).
D. equinoxialis - Belém, Içana, Tefé (Amazonas).
D. fumipennis - Pedras (Bahia), Paranaí (Paraná).
D. gaúcha - Cordoba (Argentina) (from prof. Brncic).
D. immigrans - Cassarongongo (Bahia).
D. insularis - Islands of St. Kitts and Guadabep (from prof. Dobzhansky).
D. kikkaway - Eldorado (R.G.Sul).
D. nebulosa - Lima, Tingo Maria (Perú), Natal (R.G.Norte), São Luiz (Maranhão),
Angra dos Reis (Río), Eldorado (R.G.Sul), Guaraparí (Espírito Santo),
Pitanga, Pedras (Bahia).
D. neocardini - Angra dos Reis (Río).
D. neolaetina - Itatiaia (Río).
D. neomorpha - Trinidad.
D. paulistorum - Colônia S. Pedro (R.G.Sul), Ilha das Cobras, Praia do Leste (Paraná),
Marãus (Amazonas), Florianópolis (Sta. Catarina), Belém (Pará),
Sapé (Minas), Coroico (Bolivia), Tarapoto, Tingo Maria (Perú),
San Salvador (El Salvador), Bucaramanga (Colombia), Trinidad
(from prof. Dobzhansky).
D. pallidinemnis - Eldorado (R.O.S.).
D. pararepleta - Eldorado (R.O.Sul), Ilha das Cobras, Paranaí (Paraná), Jacú
(Sta. Catarina), Pedras (Bahia).
D. payani - Vallenar Vina del Mar (Chile).
D. polymorpha - Eldorado, Itapeva (R.O.Sul), Ilha das Cobras (Paraná), Jacú
(Sta. Catarina), Pedras (Bahia), Boa Viagem (Maranhão).
D. prosaltans - Eldorado (R.O.Sul), Pitanga (Bahia), Praia do Leste (Paraná),
Guaçará (Espírito Santo), Jacú (Sta. Catarina), Boa Viagem
(Maranhão), Belém (Pará).
D. saltans - Itatiaia (Río, Pitanga (Bahia).
D. simulans - Eldorado (R.O.Sul), Angra dos Reis (Río), Pedras (Bahia), Tabatinga
(Amazonas), Boa Viagem (Maranhão), Porto Platon, Serra do Navio
(Território do Amapá).
D. sturtevant - Ilha das Cobras, Praia do Leste, Paranaí (Paraná), Pedras (Bahia),
Italiaia (Río), Belém (Paraná), Boa Viagem, Sacavem (Maranhão).
D. tropicalis - Palma, Maranguape (Brazil), Trinidad 330.
D. tropicalis cubana - Townsend.
Drosophila Species - Stocks

GERMANY

Berlin-Buch: Deutsche Akademie der Wissenschaften zu Berlin
Institut für Experimentelle Krebsforschung, Genetische Abteilung, Lindenberger Weg 70

Other Species:

80 D. funebris
81 D. funebris ev : 1. Chromosom
82 D. funebris st : autosomal
83 D. funebris ci : I
84 D. funebris wy : III
85 D. simulans : v
86 D. virilis : wild
87 D. busckii

Note: Additions and corrections to the list in DIS 34 (p. 64)

Lost: 102 D. repleta: wild

Additions: D. funebris

Chromosome 1

105 ci

Chromosome 3

106 wy

Tübingen: Max Planck-Institut für Biologie

D. busckii
D. hydei
D. miranda
D. persimilis
D. pseudoobscura
D. simulans
D. virilis

ISRAEL

Jerusalem: Hebrew University of Jerusalem

D. immigrans Brisbane
D. immigrans Israel
D. simulans
D. subobscura Küsnacht
D. subobscura Elon (Israel)

ITALY

Milano: Universita' di Milano, Instituto di Genetica

Drosophila simulans

Wild type from Pavia
Wild type from Aspra

Stocks selected for tumor manifestation:

tu A

tu B1

tu B3

tu C

tu Aspra
JAPAN

Tokyo: Tokyo Metropolitan University
Department of Biology

D. ananassae

Wild stocks

1. Texas
2. TL1
3. TL3
4. TL4
5. TL3-4
6. TL3-11
7. Barro Colollarado, Panama 69 (Low elevation)
8. Barro Colollarado, Panama 74 (Low elevation)
9. Turrialba, Costa Rica 101 (High elevation)
10. Turrialba, Costa Rica 104 (High elevation)
11. Turrialba, Costa Rica 125 (High elevation)
12. Christobal, Panama
13. Baton Rouge, Louisiana
14. Calcutta, India
15. Hawaii-H
16. Meakan (1)
17. Nishi Tappu (2)
18. 2L-AH
19. 2L-BH
20. 2L-A; 3L-AH

Mutants

21. 2L-A; 3L-AM
22. 3L-AC104
23. 2L-B; 3L-AH

24. Barro Colollarado, Panama
25. st f
26. ru
27. pxd
28. st f ru²
29. st f se
30. y f
31. b
32. Bn-R
33. bw-R
34. fu
35. j
36. px
37. S
38. se
39. sn
40. wy

D. bifasciata

Wild

1. Akkeshi (3 strains)
2. Asakawa (1)
3. Gotokuji (1)
4. Hakkoda (3)
5. Kikokonagatake (1)
6. Kitazawatoge (1)
7. Kumotoriyama (3)
8. Meakan (1)
9. Mshitappu (2)
10. Ohkurayama (1)
11. Pavia, Italy (1)
12. Pfywmal, Switzerland (1)
13. Shibunoyu (1)
14. Taetsusuzan (2)
15. Tanigawadake (1)
16. Tsukuba-san (1)

Autosomal

ag
ar
ar ic
ar ob
ar ro
arp
bn
ca ro
cn
ic
ic cn
M ob
ob
orr
ps yh
vi
y (5 strains)

Mutant

Sex-linked

a y
f
y

Cytoplasmic

Sex-ratio, Italy
Sex-ratio, Japan (6 strains)
### Other Species

<table>
<thead>
<tr>
<th>Species</th>
<th>Origin</th>
<th>Strains</th>
</tr>
</thead>
<tbody>
<tr>
<td>D. americane</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. ambigua</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. auraria</td>
<td>Wild</td>
<td>6 strains (Type A)</td>
</tr>
<tr>
<td>D. buskii</td>
<td>Wild</td>
<td>3 strains</td>
</tr>
<tr>
<td>D. chinoi</td>
<td>Wild</td>
<td>4 strains</td>
</tr>
<tr>
<td>D. funebris</td>
<td>Wild</td>
<td>2 strains</td>
</tr>
<tr>
<td>D. hydei</td>
<td>Wild</td>
<td>4 strains</td>
</tr>
<tr>
<td>D. immigrans</td>
<td>Wild</td>
<td>5 strains</td>
</tr>
<tr>
<td>D. kikkawai</td>
<td>Wild</td>
<td>5 strains</td>
</tr>
<tr>
<td>D. lutea</td>
<td>Wild</td>
<td>17 strains</td>
</tr>
<tr>
<td>D. miranda</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. novamexicana</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. obscura</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. pseudoobscura</td>
<td>Wild</td>
<td>5 strains</td>
</tr>
<tr>
<td>D. pseudoobscura</td>
<td>Mutant</td>
<td>4 strains</td>
</tr>
<tr>
<td>D. pulchrella</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. simulans</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. suzukii</td>
<td>Wild</td>
<td>4 strains</td>
</tr>
<tr>
<td>D. takahashii</td>
<td>Wild</td>
<td>37 strains</td>
</tr>
<tr>
<td>D. tristis</td>
<td>Wild</td>
<td>1 strain</td>
</tr>
<tr>
<td>D. virilis</td>
<td>Wild</td>
<td>6 strains</td>
</tr>
<tr>
<td>D. virilis</td>
<td>Mutant</td>
<td>5 strains</td>
</tr>
</tbody>
</table>

### NETHERLANDS

**Groningen:** State University of Groningen, Genetical Institute, Haren (Gr)

**Stock List - Species:**

<table>
<thead>
<tr>
<th>Species</th>
<th>Origin</th>
<th>Strains</th>
</tr>
</thead>
<tbody>
<tr>
<td>D. pseudoobscura</td>
<td>various strains homozygous for AR. CH. ST.</td>
<td></td>
</tr>
<tr>
<td>D. immigrans</td>
<td>(wild 1959)</td>
<td></td>
</tr>
<tr>
<td>D. mercatorum</td>
<td>(wild 1961)</td>
<td></td>
</tr>
<tr>
<td>D. repleta</td>
<td></td>
<td></td>
</tr>
<tr>
<td>D. hydei</td>
<td>(wild 1961)</td>
<td></td>
</tr>
<tr>
<td>D. buskii</td>
<td>(wild 1961)</td>
<td></td>
</tr>
</tbody>
</table>

### SOUTH AFRICA

**Johannesburg:** University of the Witwatersrand, Department of Zoology

**D. persimilis**

Porcupine Flat

**D. pseudoobscura**

Several strains of Standard, Arrowhead, Chiricahua, Treeline, Pikes Peak, and Santa Cruz

**gl**

**or**

**se**

**tb b v se pp**

**v**
D. simulans

Drakensberg
Free State
Inhaca Island
Johannesburg
Kalahari
Limpopo

Mkuzi Game Reserve
Nyasa Lake
Stellenbosch
Umgasi River
West Rand
Zoutspansberg

Other species

D. busckii: Inhaca Island
D. funebris: Witwatersrand, Natal
D. hydei: Inhaca Island, Natal
D. nebulosa: Brazil
D. segui: Limpopo River
D. willistoni: Brazil

D. yakuba: Northern Transvaal,
Inhaca Island

Zapirion ghesquierei
Z. tuberculatus: various strains
Z. vittiger: various strains

SPAIN

Barcelona: Universidad de Barcelona, Centro de Genetica Animal y Humana

D. ambigua. Several Spanish stocks.
D. busckii. Barcelona.
D. buzzatii. Armentera (Spain).
D. funebris. Several Spanish stocks.
D. hydei. Barcelona.
D. immigrans. Barcelona.
D. mercatorum mercatorum. Barcelona.

D. mercatorum pararepleta. Jijuca (Brazil).
D. phalerata. Several Spanish stocks.
D. repleta. Barcelona.
D. simulans. Barcelona.
D. subobscura. Several Spanish stocks;
mutant stocks.
D. testacea. Barcelona.
Parascaptomyza disticha. Barcelona.

Madrid: Centro de Investigaciones Biologicas, Laboratorio de Genetica

D. busckii: Madrid, Santianes.
D. funebris: Madrid.
D. guvenotii: Santianes.
D. immigrans: Madrid, Santianes.
D. melanogaster: Madrid, Mallorca, Ribadeo, Rocafort, Ronda 10, Ronda 30, Santianes.
D. obscuroides: Santianes.
D. repleta: Madrid.
D. subobscura: Madrid, Santianes.

SWEDEN

Uppsala 7: University of Uppsala, Institute of Genetics

Drosophila littoralis
Drosophila hydei
Drosophila funebris
Drosophila subobscura
D. arizonensis

1: lobed eye. Coulson, K59. Third chromosome recessive, spontaneous in A7.7a stock. Eye reduced in size, with notch in anterior edge. Eye reduction variable from slight notch to nearly eyeless. Eye color slightly duller and darker than wild type.
It has been shown (Alderson, 1960a) that formaldehyde exhibits no mutagenic activity towards Drosophila melanogaster, by the larval feeding method, unless ribonucleic acid is present in the treatment medium, whereas the presence or absence of ribonucleic acid has no influence on the mutagenic activity of urethane. A 6-amino group alkylation of adenylic acid by formaldehyde in the treatment medium has been shown (Alderson, 1961) to be the responsible reaction for the mediation of the mutagenic activity of formaldehyde in Drosophila: adenylic acid may be present as any of its free mononucleotides or bound in the ribonucleic acid polynucleotide, but the mutagenic activity of formaldehyde is completely dependent on the presence of adenylic acid within the treatment medium.

The responsible mutagenic alkylation of adenylic acid by formaldehyde obviously does not occur to a sufficient extent in vivo for the reaction product to significantly increase the mutation rate. Yet, even under physiological conditions, alkylation would be expected to take place. One reason for lack of effect may be that the extent of the reaction required is not attained under physiological conditions; for example, in the case of formaldehyde, dimerisation by methylene bridging of adenylic acid is postulated as the effective mutagenic alkylation (Alderson, 1960b). In the case of other mutagenic alkylation agents, the alkylation is obviously attained in vivo. Szybalski (1961) has shown that the mutagenic activity of triethylene melamine in bacteria is probably mediated by in vivo alkylation of precursor thymidylic acid since the in vitro reaction product of triethylene melamine and thymidylic acid is found to be mutagenic.

In vitro alkylation of purine and pyrimidine bases is thus implicated as an important mechanism in chemically-induced mutagenesis, especially in view of the mutagenic activity of several N-methyl xanthines in bacteria (Novick and Szilard, 1952), and that of caffeine (1,3,7-trimethyl xanthine) in Drosophila (Andrews, 1959).

Since mutagenicity by alkylation agents may be mediated by both in vivo and in vitro reactions, an attempt to separate these reactions by culturing Drosophila larvae in media containing or lacking purine and pyrimidine components was carried out. Further, the recent chemical characterisation and isolation of the products of alkylation of the nucleic acids and their constituent nucleotides (Lawley, 1960) might reveal the essential in vitro alkylations concerned in mutagenesis.

Two ethylating agents, diethyl sulphate and ethyl methanesulphonate, were chosen for a preliminary study. Both ethylating agents were tested for mutagenic activity in the presence (0.4%) and in the absence of ribonucleic acid using a chemically defined and aseptic treatment medium (see Alderson, 1960a, for the composition). Diethyl sulphate (B.D.H. 99-101 per cent) and ethyl methanesulphonate (supplied as a 0.5 M saturated solution by the Chester Beatty Research Institute) were added when the temperature of the treatment medium was 60°C. In most experiments Oregon-K eggs were sterilised using Sang's (1956) method and transferred as newly hatched larvae to the treatment medium for their entire larval life; in one experiment the larvae were treated for the first 48 hours of larval life and then transferred onto ordinary laboratory food to complete their development. Hatching males were individually examined for sex-linked recessive lethal mutations by the Muller-5 technique; two broods were used, each with two females for three days.

Neither ethylating agent showed a significant difference in its mutagenic activity whether ribonucleic acid was present or absent from the treatment medium. (Table 1.)

References.

Table 1.

<table>
<thead>
<tr>
<th>Medium</th>
<th>Mutagen</th>
<th>Concentration</th>
<th>Length of treatment of larvae</th>
<th>Larval developmental time in days</th>
<th>Per cent survivors</th>
<th>First Broods</th>
<th>Second Broods</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Total</td>
<td>Lethals</td>
</tr>
<tr>
<td>No RNA</td>
<td>Diethyl sulphate</td>
<td>$3.2 \times 10^{-4}$M</td>
<td>0-48 hrs.</td>
<td>9-9.5</td>
<td>50.25</td>
<td>337</td>
<td>2</td>
</tr>
<tr>
<td>With RNA</td>
<td></td>
<td></td>
<td></td>
<td>8-8.5</td>
<td>81.0</td>
<td>364</td>
<td>1</td>
</tr>
<tr>
<td>No RNA</td>
<td>Diethyl sulphate</td>
<td>$2.0 \times 10^{-4}$M</td>
<td>Entire larval life</td>
<td>11-12</td>
<td>54.8</td>
<td>432</td>
<td>8</td>
</tr>
<tr>
<td>With RNA</td>
<td></td>
<td></td>
<td></td>
<td>9-9.5</td>
<td>57.5</td>
<td>434</td>
<td>15</td>
</tr>
<tr>
<td>No RNA</td>
<td>Diethyl sulphate</td>
<td>$2.6 \times 10^{-4}$M</td>
<td>Entire larval life</td>
<td>15-16</td>
<td>36.0</td>
<td>378</td>
<td>8</td>
</tr>
<tr>
<td>With RNA</td>
<td></td>
<td></td>
<td></td>
<td>11</td>
<td>73.0</td>
<td>406</td>
<td>18</td>
</tr>
<tr>
<td>No RNA</td>
<td>Ethyl methane-</td>
<td>$5.0 \times 10^{-6}$M</td>
<td>Entire larval life</td>
<td>14-16</td>
<td>48.5</td>
<td>1020</td>
<td>62</td>
</tr>
<tr>
<td>With RNA</td>
<td>sulphonate</td>
<td></td>
<td></td>
<td>10</td>
<td>61.25</td>
<td>1118</td>
<td>56</td>
</tr>
</tbody>
</table>
To investigate the genetic load in many species of Drosophila, it is possible to study the wild chromosomes in homozygous condition and in random combinations. Derivation of the chromosomes often presents the investigator with a choice, however. Matings in the test cross generation may be made such that two genotypes are produced in the resulting progeny of four genotypes. The latter method, introduced by Wallace (1956), produces a "control" genotype in every culture—a type free of the wild chromosome(s) being studied. Thus, chromosomes and combinations can be scored by the usual % wild type viability measurement, by ratio of wild type to "control"; or by other ratio measurements which can be devised.

In the latter method, however, Spassky et al (1960) found the ratio of wild type to control quite variable so used % wild type viability as the measurement of chromosome performance. Other investigators, Dobzhansky, Krimbas, and Krimbas (1960) and Greenberg and Crow (1960) have used other ratios obtainable among the four genotypes, though the former also used the ratio of wild type to control.

Counting four genotypes takes longer than counting two per culture, and the variability accompanying the +/control ratio limiting the usefulness of this measure. At the suggestion of Dr. P. T. Ives, chromosomes from the August 1960 collection from the S. Amherst D. melanogaster population were used to compare the two types of test crosses.

Since only 52 of the 274 wild males received proved fertile, data on the frequency of drastic (le + sre) chromosomes would be of questionable value, so is omitted. Derivations followed standard procedure. For the test crosses two types of matings were made for each chromosome line: Cy x Cy and Cy x Bl, with one culture being scored for each type of mating. Analysis was confined to second chromosomes only.

Of the nondrastic chromosomes recovered (%+/+% greater than 17% by Cy x Cy matings; %+/+% greater than 12 1/2% by Cy x Bl matings), 13 chromosomes were chosen and 7 combinations devised to compare: (1) genetic, environmental, and sampling variances of viabilities obtained with the two methods; (2) the accuracy of viability measurement using three replicate cultures versus that obtained by counting only one culture.

Matings were as before: Cy x Cy and Cy x Bl with 3% and 3-4% as parents in each line or combination. Parents were transferred every 24 hours for three days to give three replications. Counts were made three times during the emergence period. One chromosome, 2912, was discovered later to be semilethal by Cy x Cy tests, though of low quasinormal viability by Cy x Bl tests. Another chromosome, 2935, was sterile in Cy x Cy repeat tests. Hence, computations of average viability and variances for homozygous chromosomes have been made in two ways for the Cy x Bl tests: (a) including all thirteen chromosomes; (b) excluding 2912 and 2935.

The formulae given by Dobzhansky and Spassky (1953) and Wallace and Madden (1953) have been used to partition total variance (T) into its environmental (E), genetic (G), and sampling (S) components. Comparison of variances for the two methods are shown in Table 1. Total number of flies counted in each type of test cross is also included.

<table>
<thead>
<tr>
<th>Table 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>mean viability</td>
</tr>
<tr>
<td>----------</td>
</tr>
<tr>
<td>Heterozygotes</td>
</tr>
<tr>
<td>Cy x Bl</td>
</tr>
<tr>
<td>Cy x Cy</td>
</tr>
<tr>
<td>Homozygotes</td>
</tr>
<tr>
<td>(a) Cy x Bl</td>
</tr>
<tr>
<td>(b) Cy x Bl</td>
</tr>
<tr>
<td>Cy x Cy</td>
</tr>
</tbody>
</table>
In both heterozygotes and homozygotes total and sampling variances are lower using the Cy x Bl method. Environmental variances in heterozygotes are the same for either method, but lower in homozygotes with the Cy x Bl type of mating. In heterozygotes, genetic variances are likewise lower using these matings. In homozygotes genetic variances are approximately the same for either method.

For both methods heterozygotes show highest viability and least genetic variance—the typical picture when heterozygotes and homozygotes are compared. However, for this particular set of chromosomes, larger environmental variances are found among heterozygotes. This is sometimes observed in the data of others.

Data on the means of homozygous chromosomes tested are shown in Table 2. Data on 2912 are included in the table. Mean viability of 2935 was 24.1%, original viability measurement 25.0% in Cy x Bl tests. Correlations between average viability from the three replicates per chromosome line (X3) and viability previously computed from scoring the original test cross culture (Xo) are similar for both methods, 84.9% for Cy x Bl matings and 82.5% for Cy x Cy matings.

Table 2

<table>
<thead>
<tr>
<th>Line:</th>
<th>2905</th>
<th>2908</th>
<th>2912</th>
<th>2913</th>
<th>2915</th>
<th>2916</th>
<th>2920</th>
<th>2927</th>
<th>2928</th>
<th>2929</th>
<th>2936</th>
<th>2943</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cy x Bl</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X3</td>
<td>26.2</td>
<td>22.1</td>
<td>15.0</td>
<td>24.2</td>
<td>26.1</td>
<td>18.1</td>
<td>21.0</td>
<td>20.9</td>
<td>21.1</td>
<td>17.5</td>
<td>26.6</td>
<td>24.8</td>
</tr>
<tr>
<td>Xo</td>
<td>29.5</td>
<td>24.7</td>
<td>15.4</td>
<td>23.0</td>
<td>26.2</td>
<td>15.2</td>
<td>20.1</td>
<td>25.2</td>
<td>24.3</td>
<td>15.5</td>
<td>26.4</td>
<td>21.0</td>
</tr>
<tr>
<td>Cy x Cy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X3</td>
<td>32.5</td>
<td>29.6</td>
<td>10.0</td>
<td>32.3</td>
<td>29.3</td>
<td>29.0</td>
<td>34.2</td>
<td>33.8</td>
<td>29.2</td>
<td>28.1</td>
<td>19.3</td>
<td>33.3</td>
</tr>
<tr>
<td>Xo</td>
<td>28.8</td>
<td>29.1</td>
<td>9.1</td>
<td>29.8</td>
<td>28.1</td>
<td>20.7</td>
<td>31.0</td>
<td>27.3</td>
<td>25.9</td>
<td>22.9</td>
<td>18.3</td>
<td>24.7</td>
</tr>
</tbody>
</table>

Hence, for scoring viability for each chromosome derived homozygously from a natural population, counting only one culture appears satisfactory as an index of performance and enables the rapid computation of the frequency of drastic (le + sle) in the population. Either method gives comparable results.

However, this tells nothing about performance in heterozygous condition, nor enables comparisons between heterozygotes and homozygotes. For this, the Cy x Bl method, with replicated observations, is to be preferred, primarily because variances of viability measurements are generally less than, and certainly no more than equal to, those obtained with Cy x Cy scoring. In addition, a variety of ratio comparisons can be made. So the Cy x Bl cross for D. melanogaster, or the method by which four genotypes are scored among the progeny, is the more versatile. It is this method which has been employed in subsequent analyses of chromosome samples from this population.

(Supported by grants from the National Science Foundation and the National Research Council of Canada.)

Band, H. T. Preliminary evidence that variation in temperature affects viability of heterozygous wild type flies.

The correlations between drastic (le + sle) frequency and temperature variables of range and mean detected by Band and Ives (1961) suggest a dynamic relationship between environment and genetic structure of the S. Amherst D. melanogaster population. To determine if environment could influence viability of different kinds of heterozygotes, a small preliminary experiment was conducted using second chromosomes derived from the August 1960 collection. Crosses between chromosome lines were of two types: Cy/i x Cy/j and Cy/i x Bl/j such that the wild type (i/j) progeny were known to be genotypically drastic/drastic (d/d), drastic/nondrastic (d/nd) or nondrastic/nondrastic (nd/nd). Five crosses were constructed to yield d/d
progeny, 12 to give d/nd progeny and 6 to give nd/nd progeny. The same crosses were tested by both methods, and 2 Cy ffi and 2-3 Cy or Bl d' used for each cross. Parents were transferred every 24 hours for 4 days to give 4 replicate cultures. Oviposition was at room temperature. Replicates B and D were kept at 25°C; A and C were transferred between 17°C and 25°C constant temperature incubators. The F17/25 environment corresponds to a narrow range environment according to range data given in Band Ives (1961). Transfers were made to alternate temperatures every 24 hours. Progeny were counted 3 times during the emergence period. Wild type viabilities for the 3 kinds of heterozygotes are shown in Table 1. A = average viability computed from Cy x Cy matings; B = average viability computed from Cy x Bl matings; C = the ratio of wild types produced to 1/2 (Cy + Bl) + 1. The ratio is based on the total number of flies within observed genotypes in each of the 3 heterozygous combinations.

<table>
<thead>
<tr>
<th></th>
<th>C25</th>
<th>F17/25</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A(%)</td>
<td>B(%)</td>
</tr>
<tr>
<td>d/d</td>
<td>30.6</td>
<td>20.5</td>
</tr>
<tr>
<td>d/nd</td>
<td>35.4</td>
<td>24.0</td>
</tr>
<tr>
<td>nd/nd</td>
<td>33.8</td>
<td>26.1</td>
</tr>
</tbody>
</table>

Drastic/dramatic (d/d) heterozygotes have lowest viability at C25 but improve in the narrow range F17/25 environment. In this latter environment, however, nd/nd wild types have lowest viability. Otherwise, the two methods show slightly different results. By Cy x Cy crosses, d/nd heterozygotes appear decidedly superior at C25, but at F17/25 little difference is noted between different heterozygotes, though nd/nd is lowest in viability. By Cy x Bl methods, the different genotypes react differently in each environment; viability changes determined by % wild type viability are substantiated by ratio results.

Due to the small number of chromosomes involved and the small number of progeny realized, chi-square comparisons have been made using total number of flies and total number of wild types obtained in each method for the 3 kinds of heterozygotes. Even so, only 16,831 flies have been counted in the entire experiment. Between environments, three comparisons are of interest: change in viability of d/d heterozygotes, of d/nd heterozygotes, and of nd/nd heterozygotes. For Cy x Bl crosses, d/d viability has improved significantly in the F17/25 environment from the level shown at C25, likewise d/nd. For both P<.005. No difference is detected between nd/nd viabilities at the two temperatures. For Cy x Cy matings, only the viability of d/d shows significant changes between the two environments, with .025<P<.05.

Within environments both methods indicate that the different heterozygotes are significantly different in viability at C25; P<.005 for each method. At F17/25 only the results from the Cy x Bl crosses indicate significant differences between viabilities of the different heterozygotes. Again P<.005.

The outcome of this small experiment thus indicates: (1) that environment can influence heterozygous viabilities and so may affect the genetic structure of the population through selection at the heterozygote level; (2) heterozygotes carrying drastic chromosomes (either d/d or d/nd) can be favored in narrow range environments; (3) again indicates the Cy x Bl method to be more sensitive to genotypic differences than the Cy x Cy method.
In contrast to the viable mutant, dumpy, which is genotypically designated as dp0v1, or ov1, the double mutant dp02 dpv2 or ov2 (which was prepared from the heterozygote, ed dpv2 / dp02 cl) manifests a more extreme wing and thorax effect. Furthermore, the mutant ov1 is viable with all lethal members of the dumpy series, but the double mutant ov2 is inviable with some of the lethal alleles. For this reason a study of the dumpy lethals was made with ov2 at 220 C. and at 280 C. The results in Table 1 strongly suggest that these lethals can be classified in three ways. In one group the lethal acts as a dominant in the compound with ov2 at both temperatures. The absence of a change in activity with temperature for the lethals in this group is characteristic of that class of mutants designated as amorphs (lacking any detectable activity), in Muller’s classification of gene action (Muller, 1932, Proc. 6th Int. Congr. Genet. 1:213-255). The alleles in the first group include olw, olv54e, lv1, and lvx3. The second group of alleles shows a greater manifestation of lethality at 280 C. than at 220 C. These might be considered hypomorphic, with a partial normal activity of the lethal allele restored at lower temperature, just as is the case for the (o) and (v) effects. These mutants include O1m and olm. A third group of mutants is more difficult to classify. For example, olvbm shows little difference in temperature response with a suggestion of a reversal of sensitivity. The mutant olv, in contrast to olm, is much more viable with ov2 at 280 C.

When the lethals are tested in compound with the allele dpcm2 or cm2 the response is quite unexpected -- cm2 unquestionably reverses the manifestation of the lethal at these temperatures. This is not the case, however, for the (o) and (v) effects manifested by cm2 in its viable compounds, which show an exaggerated mutant expression at higher temperatures.

The developmental basis for this variation in expression of the lethal effect is not apparent from these data. However, the ov2 and cm2 alleles provide a useful classifying technique for phenotypically similar mutants and for determining the quantitative degree of expression of the lethal effect in the dumpy series.

Table 1. Lethal expression with temperature in the dumpy allelic system

<table>
<thead>
<tr>
<th>P1 Cross</th>
<th>280 C.</th>
<th></th>
<th>220 C.</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>non-Cy</td>
<td>Cy</td>
<td>% viable</td>
<td>non-Cy</td>
</tr>
<tr>
<td>olv1 x ov2</td>
<td>0</td>
<td>351</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>olvW x ov2</td>
<td>0</td>
<td>202</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>olv54e x ov2</td>
<td>0</td>
<td>249</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>lv1 x ov2</td>
<td>0</td>
<td>518</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>lvx3 x ov2</td>
<td>0</td>
<td>24</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>I1m x ov2</td>
<td>2</td>
<td>872</td>
<td>0.31</td>
<td>4</td>
</tr>
<tr>
<td>olm x ov2</td>
<td>1</td>
<td>259</td>
<td>0.36</td>
<td>26</td>
</tr>
<tr>
<td>olv x ov2</td>
<td>13</td>
<td>73</td>
<td>15.1</td>
<td>43</td>
</tr>
<tr>
<td>olvbm x ov2</td>
<td>3</td>
<td>236</td>
<td>1.25</td>
<td>1</td>
</tr>
<tr>
<td>olvW x cm2</td>
<td>87</td>
<td>304</td>
<td>22.2</td>
<td>5</td>
</tr>
</tbody>
</table>
(Carlson and Falk, Table 1—continued)

<table>
<thead>
<tr>
<th>P1 Cross</th>
<th>28°C</th>
<th>Cy</th>
<th>% viable</th>
<th>22°C</th>
<th>Cy</th>
<th>% viable</th>
</tr>
</thead>
<tbody>
<tr>
<td>olv5he x cm² Cy</td>
<td>27</td>
<td>211</td>
<td>11.3</td>
<td>13</td>
<td>594</td>
<td>2.14</td>
</tr>
<tr>
<td>olvbm x cm² Cy</td>
<td>76</td>
<td>409</td>
<td>15.7</td>
<td>2</td>
<td>518</td>
<td>0.39</td>
</tr>
<tr>
<td>olv¹ x cm² Cy</td>
<td>44</td>
<td>426</td>
<td>13.3</td>
<td>4</td>
<td>714</td>
<td>0.56</td>
</tr>
</tbody>
</table>

This work is supported by grant 14222 from the National Science Foundation.


The investigation has been carried on culturing cells derived from two organs of Drosophila melanogaster: the cephalic ganglia and the lymph gland, both in larval stage. For studying the ganglia the following wild stocks have been chosen: Barese, S. Maria and Aspra 52. For the lymph gland, stocks S. Maria, yw and Chieti, where the structure of the gland is very similar (lobes of the first pair of loose structure but fairly well developed).

Cultures were set in a drop of medium put on the coverslip, which is kept underneath, while the slide is above. The culture medium for Insects according to Kuroda and Tamura (1956) has been used.

The ganglia have been kept in culture for 28 days; the lymph gland for 21 days. In both cases, the culture medium has been replaced every 7th day. Each time both the organ and the cells released from it have been pipetted and transferred in a drop of fresh medium. Cells adhering to the coverslip have been washed, stained with May Grünwald-Giemsa, classified and counted.

Using this method it is possible to detect stable differences between the stocks, as far as the behaviour of the organ and of the cells is concerned. Since the culture conditions are strictly controlled, cultivation in vitro permits to recognize entirely new physiological properties at cellular level, which are apparently genetically controlled.


In our previous studies on the biochemical effects of lethal factors in D. melanogaster various abnormal patterns of free amino acids in the lethal homozygotes were reported. Since the amino acid pattern is characteristic for each mutant, it seems that the effect is locus-specific (for a general review, see Hadorn 1955). In order to obtain more insight into the intermediary metabolism of amino acids in Drosophila and as a basis for comparison between lethal and normal individuals, studies on the transaminase activities in the wild type have been carried out in our laboratory. Fat bodies from ten +/+ larvae aged about 96 hours (at 25°C) were dissected out in ice-cold insect Ringer's solution and homogenized. The homogenate was incubated in a sodium-potassium-phosphate buffer solution (0.067M; pH 7.56) at 38°C with one keto acid (ß-ketoglutaric acid, oxalacetic acid, pyruvic acid or glyoxylic acid) as the amino acceptor and one amino acid (aspartic acid, glutamic acid, ß-alanine, leucine, threonine, glycine, valine or arginine) as the amino donor. In addition, pyridoxal phosphate was added as
<table>
<thead>
<tr>
<th>Days after irradiation</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adults 12 h. control</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adults 6 d.</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adults 12 h.</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pupae 48 h. (500 r.)</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prepupeae 2 h.</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Larvae 72 h.</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Larvae 48 h.</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Larvae 24 h.</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.8</td>
<td>1.7</td>
<td>0.4</td>
<td>0.9</td>
<td>0.6</td>
<td>0.4</td>
<td>0.6</td>
<td>0.6</td>
<td>0.7</td>
<td>0.6</td>
<td>0.7</td>
<td>1.4</td>
</tr>
<tr>
<td>14.4</td>
<td>11.8</td>
<td>14.2</td>
<td>12.7</td>
<td>26.2</td>
<td>33.0</td>
<td>46.6</td>
<td>52.0</td>
<td>62.6</td>
<td>32.0</td>
<td>18.5</td>
<td>10.7</td>
</tr>
<tr>
<td>6.7</td>
<td>6.7</td>
<td>7.0</td>
<td>11.0</td>
<td>29.8</td>
<td>38.0</td>
<td>39.8</td>
<td>44.0</td>
<td>7.5</td>
<td>1.3</td>
<td>1.1</td>
<td>0.9</td>
</tr>
<tr>
<td>10.8</td>
<td>16.5</td>
<td>24.6</td>
<td>24.0</td>
<td>22.6</td>
<td>32.6</td>
<td>5.1</td>
<td>1.7</td>
<td>1.3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
From these data it appears quite likely that meiosis (beginning with pupation) has been sampled between the fourth and fifth days after irradiation. From the displacement of the peak value from day 7 to day 9 a. i. it is concluded that it may correspond to the most damaged cytes located at the end of the sampled cytes. It is also apparent that the value corresponding to the 8th day after irradiation includes a higher proportion of cytes as the number of germ cells increases during development. The subsequent drop in mutation rate coincides with the appearance of gonial mitosis (24 h. old larvae) and germ cell mortality. There is an apparent delay of the speed of the spermatogenesis probably due to the increase of the number of germ cells during development.

(Work supported by a grant to E. Ortiz and associates from the Junta de Energía Nuclear, Madrid.)

Gardner, E. J., and A. M. Hansen. Further studies on the transfer of the tumorous head maternal effect in D. melanogaster by injection. Gardner, Turner and Berseth (Genetics 45:905-913, 1960) injected extract prepared from tumorous head females and mated them with tu-h males. Eighteen per cent of the progeny expressed the tumorous head trait as compared to less than four per cent for the controls from un.injected females. These results were interpreted to indicate that the maternal effect had been transferred by the injection of tu-h female extract.

When this line of investigation was resumed several months later it was not possible to transfer the maternal effect by injections of tu-h female extract, tu-h female hemolymph, or tu-h oöplasm into Samarkand and Canton wild-type females. Experiments involving the original Lausanne stock were then repeated. Tumorous head flies appeared among the progeny of all the control crosses as well as among those of injected females. The Lausanne stock in our laboratory now carries the tu-l gene or an allele of tu-l that has presumably entered through mutation or contamination. A Lausanne S stock obtained from the California Institute of Technology was also found to carry tu-l or an allele of tu-l, but the tu-l gene was not present in a Lausanne S stock obtained from Johns Hopkins University. More than two thousand progeny were produced from 16 females of this stock that had been injected with tu-h female extract and mated with tu-h males. Less than one per cent of these expressed the tu-h phenotype. This result was comparable with that of the controls.

Another series of experiments was designed in which the extract was prepared in a more concentrated form and maintained in an ice bath during preparation and injection. Injections were made into either the thorax or abdomen of Samarkand and Cockaponsett females. Progeny from the injected flies did not express the tumorous head phenotype in greater proportions (less than one per cent) than the controls. The transfer of the maternal effect was not accomplished under the conditions of this experiment.

Glass, H. Bentley. The mutagenic effect of a 5-r dose of x-rays. The experiment in progress, reported in DIS-33 and DIS-34, has been completed with scoring of the mutations from 50 exactly balanced and coded control and irradiated series of tests. The total number of individuals scored is 1,360,948. As previously reported, each parent is exposed to 5r (dose rate about 40r/min), irradiated males are crossed with irradiated females, and dominant Minutes are scored in the F1. In 32 of the 50 replications of the experiment, the Minute mutations in the treated series have exceeded the number in the control series. The cumulative results are as follows:

<table>
<thead>
<tr>
<th></th>
<th>Minutes in unirradiated control series</th>
<th>Minutes in irradiated series</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>334/684,160</td>
<td>383/676,788</td>
</tr>
<tr>
<td>Minutes in unirradiated</td>
<td>0.049%</td>
<td>0.037%</td>
</tr>
<tr>
<td>Minutes in irradiated</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
The difference between the irradiated and control series is thus 0.008% in contrast to the prediction of 0.005% based on extrapolation downwards from dosages of 1000r and 2000r. A simple $X^2$ test yields a value, for one degree of freedom, of 3.92: $P = .047$ (when cultures with clusters of more than 3 Minutes are excluded from the data, inasmuch as clusters cannot be produced by the irradiation of mature spermatocytes and oocytes). When cultures with more than 2 Minutes are excluded, as an alternative statistical correction, the difference is statistically strengthened. $X^2 = 4.60$: $P = .032$

The viability and/or fecundity of the irradiated series is significantly lowered, by 7,540 flies, or 1.1%.

Glassman, E. Some observations on the prune-killer gene.

In DIS 33:136 there was reported a method for selecting back mutations at the prune locus using the following cross:

$$pn^- \times 1J1/sc^8\cdot Y; K-pn$$

The following observations of interest have been made.

1. The main progeny group following irradiation are the half-translocations, which suggests that this system might be used to test the effects of various agents on this group of chromosome aberrations.

2. A second, smaller group, consists of males derived by non-dysjunction in their fathers, which suggests that this system might have applications here, too.

3. It is found that the $pn^- = /sc^8\cdot Y; K-pn/+ female progeny die as pupae. This is unexpected since males which are $pn; K-pn/+ die in early second instar. This points to an interesting sex difference in the expression of these genes.


Gene dosage relations at the $ma-l$ and $ry$ loci.

Assays of xanthine dehydrogenase in flies heterozygous for $ry$ (ie $ry^+ / ry^1$ and $ry^+ / ry^2$) show that these heterozygotes have about 40-70% of the activity of normal. Thus, the $ry$ mutants appear to be similar to other genes in which the heterozygote has lower enzyme activity than the homozygous wild-type.

On the other hand, $ma-l^+ / ma-l$ heterozygotes show an activity of xanthine dehydrogenase which is greater than or equal to the wild-type. It is felt that the increase of activity in heterozygotes is associated with autosomal heterozygosis since males derived from $+ \times ma-l$ also show greater activity than $+$. The reason for this increase is not understood but it might be due to an increase in body mass or to an increase in vigor associated with heterozygosis.

That $ma-l^+ / ma-l$ flies are probably equivalent to $ma-l^+ / ma-l^+$ is substantiated by the insertion of $ma-l^+$-bearing chromosomal fragments derived either from the $T(1;4)BS$ translocation or the duplication $dp(1;3)BS^3$ into various diploid combinations of $ma-l^+$ and $ma-l$. The results demonstrate essentially no difference between the following genotypes:

$$ma-l^+ / ma-l / ma-l$$
$$ma-l^+ / ma-l +$$
$$ma-l^+ / ma-l^+ / ma-l^+$$

In other words, one dose of $ma-l^+$ produces an amount of xanthine dehydrogenase activity which is equal to three doses of $ma-l^+$.

If the analysis of Jacob and Monod on the lac region of $E. coli$ is applicable, then one might conclude from the above that $ry$ is a structural gene for xanthine dehydrogenase and $ma-l$ is a regulator gene. The regulation is expressed
not through a repressor substance, but through an internal inducer which is not the substrate. However, the fact that the ma-1 locus does have mutants which form a CRM and which show complementation, both of which are attributes of a structural gene, is not consistent with this hypothesis. Another possibility is that the ma-1 locus regulates the activation of the already formed enzyme molecule. Many models based on this assumption can be suggested, but none of them are subjectable to experimental analysis at the present time.

Goldberg, A., A. Schalet, and A. Chovnick.
On the lethality of double mutants of HnR-3 and various ry mutant alleles.

Taira has reported that the double mutant chromosome, HnR-3 ry behaves as a recessive lethal (DIS-34). If the synthetic lethality of HnR-3 and ry applied as well to combinations of HnR-3 and other rosy alleles, then the lethal effect might be used as the basis for a highly efficient system designed to select for pseudoallelic recombinants at the rosy locus. Moreover, selective systems could be developed for the study of reverse mutation of HnR-3 and rosy alleles, and for sex-linked suppressors and dominant suppressors of both HnR-3 and ry. Consequently, we synthesized five chromosomes bearing HnR-3 and each of five different rosy alleles (ry1, ry2, ry4, ry6, and ry9) in order to check for lethal effects of the mutant combinations (the three chromosomes with ry4, ry6, or ry9 also carried cu kar). Since both HnR-3 and ry affect pterine metabolism, the chemotypes of all genotypic combinations were examined by direct chromatography of heads and abdomens.

The mating scheme, run in parallel for all rosy mutants, was designed to provide an unambiguous answer to the question of the lethality of HnR-3 ry double mutants. Heterozygous females were produced, HnR-3 ry+sr/ry+, and back-crossed to HnR-3 ry+sr/HnR-3 ry+sr males. Single male offspring of the phenotype henna, non-stripe (crosses between HnR-3 and sr) were crossed to ry2 126 Sb Ubx/In(3)DoxF females to distinguish between the crossover classes Hn-ry, and ry-sr. Males of the former class would be HnR-3 ry+sr/HnR-3 ry+sr and would be distinguished from the latter class by producing rosy offspring. In those vials which did produce rosy offspring, there was no significant deviation from the expected frequency of rosy offspring. From vials which did produce rosy offspring, ry Sb Ubx males (HnR-3 ry/ry2 126 Sb Ubx) were crossed to M34 Dfd ry1/In(3)DoxF females, yielding HnR-3 ry/In(3)DoxF males and virgin females, which were mated. In the final cross, the expected ratio of HnR-3 ry/In(3)DoxF to HnR-3 ry/HnR-3 ry progeny is 2:1. The following table shows ratios obtained for all combinations of ry mutants grown at 26°C.

<table>
<thead>
<tr>
<th>HnR-3 ry1</th>
<th>HnR-3 ry2</th>
<th>HnR-3 ry4</th>
<th>HnR-3 ry6</th>
<th>HnR-3 ry9</th>
</tr>
</thead>
<tbody>
<tr>
<td>HnR-3 ry1</td>
<td>229/88</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HnR-3 ry2</td>
<td>357/156</td>
<td>439/152</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HnR-3 ry4</td>
<td>442/191</td>
<td>368/160</td>
<td>409/130</td>
<td></td>
</tr>
<tr>
<td>HnR-3 ry6</td>
<td>357/190</td>
<td>501/206</td>
<td>319/105</td>
<td>364/0</td>
</tr>
<tr>
<td>HnR-3 ry9</td>
<td>449/194</td>
<td>395/193</td>
<td>344/140</td>
<td>334/94</td>
</tr>
</tbody>
</table>

From the data presented, it can be seen that the double mutant HnR-3 ry6/ HnR-3 ry6 behaves as a lethal, but all other combinations are viable.
Of some interest are observations made on fluorescent pterines found in the various genotypes. For all rosy alleles tested, the double heterozygotes, $Hn^{-3}$ $+/+$ $ry$, appear phenotypically wild type, but exhibit a considerable increase over Oregon-R controls in content of sepiapterine, and the spot containing biopterin and 2-amino-4-hydroxypteridine. Examination of all viable double mutant homozygotes and all combinations of rosy mutant heterozygotes (homozygous for $Hn^{-3}$) revealed that they contained large amounts of sepiapterine, a considerable increase in the spot containing both biopterin and 2-amino-4-hydroxypteridine, and were lacking isoxanthopterin.

Mr. Alfred Goldberg is an undergraduate student at Harvard College, Cambridge, Massachusetts. His participation in this study was made possible by The National Science Foundation Undergraduate Research Participation Program conducted at The Biological Laboratory, Cold Spring Harbor, during the summer of 1961.

Rapaport (1939) first produced a phenocopy of the mutant yellow by adding soluble silver salts to the food of developing larvae. Yaffe (1956) ascribed this phenocopy to a blocking of tyrosinase activity by silver ions. He demonstrated the effect in vitro in a mixture of prepupal hemolymph with tyrosine solution. The blackening of this mixture is inhibited by the addition of silver nitrate.

In view of this specific influence of silver on melanin formation it appeared paradoxical that small amounts of silver nitrate were also reported to produce melanic inclusions in Drosophila (Sand & McDonald, 1954).

In order to test the effect of silver on the dark pigment of the 'pseudotumors' of Drosophila melanogaster, a stock with high tumor penetrance was grown on standard corn molasses agar medium containing silver nitrate. The emerging adults were scored for normal or 'yellow' body pigmentation and for presence or absence of tumors. Flies scored as non-tumorous on first inspection were cleared in 10% potash in order to reveal any tumors that had escaped attention. The results are summarized in the following table:

<table>
<thead>
<tr>
<th>Medium</th>
<th>No. bottles</th>
<th>Total flies</th>
<th>Non-melanotic Males</th>
<th>Non-melanotic Females</th>
<th>Melanotic Males</th>
<th>Melanotic Females</th>
<th>Total melanotic</th>
<th>With pale integument</th>
<th>With pigmented integument</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regular</td>
<td>6</td>
<td>3753</td>
<td>325 (8.7)</td>
<td>245 (6.5)</td>
<td>1439 (38.3)</td>
<td>46.5</td>
<td>84.8</td>
<td>0</td>
<td>100.0</td>
</tr>
<tr>
<td>+ 0.0125% AgNO$_3$</td>
<td>5</td>
<td>1307</td>
<td>2 (0.2)</td>
<td>5 (0.4)</td>
<td>655 (50.1)</td>
<td>49.3</td>
<td>99.5</td>
<td>99.0</td>
<td>1.0*</td>
</tr>
<tr>
<td>+ 0.0250% AgNO$_3$</td>
<td>6</td>
<td>860</td>
<td>2 (0.2)</td>
<td>4 (0.5)</td>
<td>446 (52.0)</td>
<td>47.6</td>
<td>99.3</td>
<td>99.0</td>
<td>0.1**</td>
</tr>
</tbody>
</table>

+ 5 males, 8 females all melanotic  
++ 1 male, melanotic

It is seen that silver nitrate concentrations which produce 100 per cent phenocopies give also rise to a drastic reduction in the number of emerging flies. Among the survivors the tumor incidence reaches almost 100%.
It cannot be decided whether the salt kills off most non-tumorous individuals along with a good many tumorous ones or whether it kills at random and raises tumor incidence among the survivors. The increase in tumor penetrance may well be due to an oxidizing effect of the silver ion becoming reduced to metallic silver. In the 'suppressor-erupt' stock, at all events, tumor incidence is much enhanced by various oxidizing agents (Plaine, 1955).

Especial interest attaches to the prevailing phenotype in the silver nitrate cultures. This is the fly exhibiting the pale integument, while harboring one or several deeply pigmented tumors.

This result may indicate that an enzyme which is essential for cuticular melanogenesis is not involved in the pigmentation of the cell aggregates forming the tumor sheath (Rizki & Rizki, 1959). Such an assumption gains support from the feeding experiments of Plaine and Glass (1955), who found tyrosine, the classical substrate of melanogenesis, to have little effect on tumor penetrance, while tryptophane feeding greatly enhanced the incidence of the pigmented aggregates.

Alternatively, the silver ions may fail to reach the aggregates at the critical stage of their blackening. The present experiment does not exclude this possibility. It should also be noted that the mouth hooks of larvae in silver nitrate cultures appear as dark as those of untreated individuals.


Gottschewski, G. H. M., and W. Querner. Earlier investigations demonstrated the spreading of injected fluorochromes in explanted cephalic complexes of different larval stages. Eventually the influence of the brain on the differentiation of the eye-imaginal-disc during development. This influence could be nervous or humural. In order to show the eventual spreading of substances from the brain to the eye disc, we injected smallest amounts (0.5 - 5 x 10^-10 ml) of fluorochrome-solutions in different parts of explanted cephalic complexes, taken from late second to late third larval stages. With a new arrangement of the Zeiss-photomicroscope for mixing normal light and UV-light in any proportion we could observe both, the structures of the tissue by phase contrast and the spreading of the fluorescent substances within the tissue. In a first series we injected in the late 3. larval stage either of 4 spots: posterior part of the hemisphere, anterior part, which becomes the medulla, eye-imaginal-disc, and antennal-disc. After injection in the posterior hemisphere, the substances spread in this part only, nothing reaches the anterior part of the neighboring hemisphere or the supraoesophageal ganglion, in spite of the connection between all parts. Likewise, the fluorochromes injected in the anterior part of the hemisphere do not spread into the posterior part, indicating a barrier inside the hemisphere, not allowing substances to pass. However, the fluorochromes pass from the anterior part of the hemisphere to the eye disc and from there to the antennal disc rather quickly. Vice versa, substances injected in
the eye-disc or in the antennal disc only spread in these two organ-Anlagen (then still connected). However, they do not pass in the brain or in the second eye-antennal disc of the explant. These results demonstrate, that the transport of the injected fluorochromes between brain and eye-disc is only allowed in one way (brain towards eye), the other one (eye towards brain) is blocked. In a second series we injected in earlier stages, where the frontal sac is in full action, its ends slide as a mucous layer on each hemisphere and the connection between brain and eye-disc by the nerve cord is not yet strong. Because of the difficulty to carry the injection-needle to the small eye-disc without stripping off the fluorochrome in the mucous frontal sac we only injected in the hemisphere. The same barrier in the hemisphere as in the late 3. stage was found. The substances do not pass from posterior to anterior and vice versa. And as in the first series the fluorochromes are spreading from the anterior brain part to the eye disc, and from there to the antennal disc. In contrast to the first series they spread from there to the second eye-antennal-Anlage, again ending nevertheless at the second hemisphere; thus demonstrating the same barrier from eye to brain as in the late 3. larval stage.* We assume, that other substances may pass from the medullar part of the brain to the eye as well as the fluorochromes, and that, consequently, there is a way to influence the differentiation of the eye by substances from the brain.

*Fixation of the explants by Formol or Alkohol after injection does not change the spreading effect of the fluorochromes, both barriers are visible before and after fixation.

Greenberg, Rayla. Two new cases of SD found in nature.

Natural populations of Drosophila melanogaster from two new localities have now been shown to exhibit segregation distortion. Among 503 males sent from Berea, Kentucky, by Dr. Frank Seto, 24, or a frequency of .048, carry the SD gene; in a group from DeKalb, Illinois, collected by Dr. Jack Bennett, 1 among 184 tested males, or .005, show distortion. The finding of such a low frequency in the latter locale indicates that other populations previously scored as negative (e.g. by Mange, 1961) may, on more extensive investigation, be found to contain SD. Indeed, the frequencies given here are minimal since a few additional SD's have appeared after successive backcrossing of the wild flies to a laboratory stock, suggesting that suppressors present in the males collected may now have been eliminated.

The Berea SD's examined conform in the following properties to the original SD discovered in Madison, Wisconsin (see Sandler, Hiraizumi and Sandler, 1959), and to the Baja California SD found by Mange (1961).

1) males heterozygous for a putative SD chromosome and a cn bw chromosome show mean segregation ratios ("k" values) of .99 in favor of the SD-bearing chromosome.

2) heterozygous females produce mean k values of .52.

3) one or more inversions are present in the right arm of the SD chromosome, as shown by suppression of crossing over between cn and bw.

4) SD action is inhibited when heterozygous with the Cy inversion.

5) a heterozygote between a Berea SD and a Madison SD exhibits no distortion.

These further findings of meiotic drive indicate its widespread occurrence and possible ubiquity in natural populations of Drosophila. Additional studies are being done, with particular interest in the question of sensitivity of wild non-SD chromosomes to the action of SD.
Bridges and Li (Morgan, Bridges and Schultz, 1936), Carnegie Year Book 35:293, and also quoted in Bridges and Brehme, 1944) describe the rearrangement associated with Glazed as a single pericentric inversion of the second chromosome with breakpoints at 27E and 51D. This inversion should permit crossing over between the tip of 2L and the breakpoint at 27E. Recent crossover tests failed to detect any recombination in this region among 12,727 flies. Salivary gland chromosome analysis reveals two additional breaks in the Glazed chromosome at 22D and 33F and which appear identical with those of In(2L)Cy. Furthermore, the heterozygote, Gla/In(2L)Cy shows a single pericentric inversion difference between the two chromosomes. It seems probable that the Glazed inversion was originally induced in an In(2L)Cy chromosome. The correct arrangement of the Glazed chromosome appears to be: 2L tip to 22D/33F to 27E/51D through the centromere to 33F/22D to 27E/51D to 2R tip. Ins(2LR)Gla is a good balancer for all of 2L and the proximal half of 2R.


inversion of the second chromosome with breakpoints at 27E and 51D. This inversion should permit crossing over between the tip of 2L and the breakpoint at 27E. Recent crossover tests failed to detect any recombination in this region among 12,727 flies. Salivary gland chromosome analysis reveals two additional breaks in the Glazed chromosome at 22D and 33F and which appear identical with those of In(2L)Cy. Furthermore, the heterozygote, Gla/In(2L)Cy shows a single pericentric inversion difference between the two chromosomes. It seems probable that the Glazed inversion was originally induced in an In(2L)Cy chromosome. The correct arrangement of the Glazed chromosome appears to be: 2L tip to 22D/33F to 27E/51D through the centromere to 33F/22D to 27E/51D to 2R tip. Ins(2LR)Gla is a good balancer for all of 2L and the proximal half of 2R.

Hadorn, E., and I. Faulhaber. Range of variability in cell number of larval salivaries. Earlier studies of our laboratory showed that some larval and pupal lethals affect the cell number of the salivary glands. This finding points to an early action of the mutant genotype during embryogenesis. Several genotypes have now been investigated by counting the nuclei in Gomori stained whole mounts of salivaries. Thereby a rather high variability was found which seems to depend on a polygenic basis. The following table shows a few examples from cultures kept at 25°C on standard food and under equal population density conditions.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Range of variability in single lobes</th>
<th>Mean for both lobes per individual</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild stock</td>
<td>107 - 144</td>
<td>256.7 ± 2.87</td>
<td>15</td>
</tr>
<tr>
<td>Sevelen</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1gl/Cy</td>
<td>90 - 153</td>
<td>247.3 ± 2.07</td>
<td>67</td>
</tr>
<tr>
<td>1gl/1gl not outcrossed</td>
<td>90 - 148</td>
<td>237.5 ± 2.26</td>
<td>73</td>
</tr>
<tr>
<td>1gl/1gl after outcrossing</td>
<td>97 - 160</td>
<td>257.3 ± 2.45</td>
<td>61</td>
</tr>
<tr>
<td>173 a larval lethal</td>
<td>109 - 174</td>
<td>293.0 ± 2.57</td>
<td>73</td>
</tr>
</tbody>
</table>

There is a distinct correlation between the cell number of the two lobes within an individual. The cell number increases when the time of embryonic development is prolonged by keeping the freshly laid eggs at 18°C instead of 25°C. Thus the high numbers found in some lethals (at 25°C) might result from the fact that these genotypes develop more slowly than normals.
An unusually low rate of recovery of females occurred when pair matings of \textit{vwm} attached X females to Canton \textit{S}\textsuperscript{+} males were made. One mating in particular gave 74\% males and only 26\% females. Cultures with the lowest recovery of females were selected when possible for the mating of the progeny with the following summary of results:

<table>
<thead>
<tr>
<th>Cross</th>
<th>No. of Cultures</th>
<th>\textit{Males}</th>
<th>\textit{Females}</th>
<th>%\text{ male}</th>
<th>% male range of individual cultures</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. \textit{vwm} \textit{F}\textit{1} \textit{♀} \textit{X} \textit{F}\textit{1} + \textit{♂}</td>
<td>16</td>
<td>1357</td>
<td>1132</td>
<td>55.3</td>
<td>49.3-74.0</td>
</tr>
<tr>
<td>2. \textit{vwm} \textit{F}\textit{2} \textit{♀} \textit{X} \textit{F}\textit{2} + \textit{♂}</td>
<td>6</td>
<td>481</td>
<td>325</td>
<td>59.6</td>
<td>41.1-75.4</td>
</tr>
<tr>
<td>3. \textit{vwm} \textit{F}\textit{3} \textit{♀} \textit{X} \textit{car, ru} \textit{♂}</td>
<td>7</td>
<td>453</td>
<td>139</td>
<td>76.5</td>
<td>60.4-96.9</td>
</tr>
<tr>
<td>4. \textit{vwm} \textit{F}\textit{4} \textit{♀} \textit{X} \textit{cår, ru} \textit{♂}</td>
<td>5</td>
<td>392</td>
<td>66</td>
<td>85.6</td>
<td>74.2-95.8</td>
</tr>
</tbody>
</table>

Controls:

<table>
<thead>
<tr>
<th>Males</th>
<th>Females</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. al, ru \textit{♀} \textit{X} \textit{F}\textit{2} + \textit{♂}</td>
<td>1060</td>
<td>1040</td>
</tr>
<tr>
<td>2. bw, st \textit{♀} \textit{X} \textit{F}\textit{2} + \textit{♂}</td>
<td>1248</td>
<td>1269</td>
</tr>
</tbody>
</table>

It is not known whether the low rate of recovery of females can be accounted for by their mortality, but apparently the effect is inherited in such a manner that only the female parent produces the effect. Thus the attached X itself or something which interacts with it is suggested to be the cause of the low recovery rate of females. Unfortunately the stock was lost when the furnace overheated at the University of Utah.

A new recessive mutant in \textit{D. melanogaster} has been found in a wild Cockaponsett stock. It was given the descriptive name scarp and the gene was symbolized \textit{scrp} (current DIS). Linkage studies placed the \textit{scrp} locus at 74\% in the second chromosome. Scarp overlaps wild-type completely at 25\°C, but the penetrance is approximately 80 per cent at 30\°C. A temperature effective period has been established that extends from the forty-second hour to the sixty-eighth hour after fertilization when development begins at 30\°C. The full 26 hours at 30\°C. are necessary for maximum penetrance.

A number of "wild" laboratory stocks and one sample from a natural population were tested for the presence of \textit{scrp}. Scarp was present in three of the laboratory stocks, but two of them were derived from the same stock and had been separated for about three years.

Examinations of histological sections of scarp eyes showed that certain ommatidia are shorter than others. The \textit{scrp} gene presumably controls this condition. There is a considerable time lapse between the temperature effective period and the time at which the ommatidia elongate. Further investigation is necessary to determine the intermediate steps.

The effects of high temperature upon the frequency of expression of \textit{welt}, \textit{lobe}, and \textit{lobe-recessive} have been examined. A temperature of thirty degrees centigrade significantly increased the frequency of expression in the \textit{F}\textsubscript{1} heterozygotes from crosses of \textit{lobe} or \textit{lobe-recessive} with wild-type. Homozygous \textit{welt} was found to be lethal at 30\°C.
Heed, W., J. Russell, and D. Harrington. Diversity and density of Drosophila in the immediate vicinity of Tucson with special reference to *D. pseudoobscura*.

The following list of species has been accumulated in 41 irregular collecting trips during a 3 1/2 year period (1958-61) within a 20 mile radius of Tucson, Arizona. The two main habitats collected (chiefly by lard cans containing old bananas) are the pine and fir in the Santa Catalina Mountains (6-9000') and the cactus and reperian in the desert (2-5000').

The three most abundant species (above 1000 individuals) in the total collection of 20,766 individuals and 32 species are *pseudoobscura* (50%), *hamatofila* (22%) and *simulans* (6%). The six next most common species (400 to 700 individuals) are each 2 or 3% of the total. They are *melanogaster*, *pseudoobscura-like*, *victoria*, *longicornis*, *rubrifrons* and *hydei*. The seven next most common species (100 to 300 individuals) are each .5 to 1% of the total. They are *nigrospiracula*, *macrospina*, *carbonaria*, *macroperta*, *nigrospiracula-like*, *innubila* and *azteca*. The remaining 16 species (one half of the total) are less than 70 individuals each. They are *tenebrosa* (65), *californica* (56), *Leucophenga varia* (35), *grisea* (28), *arizonensis* (15), *Clastopteromyia inversa* (10), *Gitonavisualis* (8), *Chymomyza sp.* (6), *busckii* (5), *Scaptomyza graminum* (5), *melanopalpa* (3), *montana* (2), *bifurcata* (2), *Giton america* (2), *nigrohydei* (1), and *Leucophenga pulcherrima* (1).

Of the 32 species collected, 1/3 of them (11) are restricted to the mountains, 1/3 of them (10) are restricted to the desert, and 1/3 of them (11) are found in both habitats. A total of 22 species were collected in the mountains and 19 in the desert. *D. pseudoobscura* represents 71% of the 13,038 individuals collected in the mountains. It represents 14% of the 7,728 individuals collected in the desert and is the third ranking species there. *D. hamatofila* makes up 12% of the mountain fauna and ranks second there and 38% of the desert fauna where it is the most abundant species. *D. simulans* is second in abundance in the desert (14%) but ranks 13th in the mountains (0.3%).

The main difference in species abundance at the two elevations is that the desert contains a higher frequency of common species (11 species are each 2% or more of the number of individuals in the desert). In the mountains only 4 species reach a frequency of at least 2%. The differential is due to the swamping effect of *pseudoobscura* in the mountains. Does *pseudoobscura* really affect the frequency distributions of other species' abundances in the mountains as compared to the desert? Reference to the table shows a surprising similarity in the two faunas in the ranking of species if *pseudoobscura* is deleted from the mountains. There are now one abundant species (42%), two common species (12 to 17%) and the remainder fall off gradually from about 6%.

### Species Ranked According to Density

<table>
<thead>
<tr>
<th>MOUNTAIN (without pseudoobscura)</th>
<th>DESERT</th>
</tr>
</thead>
<tbody>
<tr>
<td>N = 3,735</td>
<td>N = 7,728</td>
</tr>
<tr>
<td>(23 collections in four different months)</td>
<td>(18 collections in seven different months)</td>
</tr>
<tr>
<td>1. <em>hamatofila</em> 42.3%</td>
<td>37.6% <em>hamatofila</em></td>
</tr>
<tr>
<td>2. <em>pseudoobscura-like</em> 17.2</td>
<td>14.2% <em>simulans</em></td>
</tr>
<tr>
<td>3. <em>rubrifrons</em> 12.4</td>
<td>14.1% <em>pseudoobscura</em></td>
</tr>
<tr>
<td>4. <em>longicornis</em> 5.5</td>
<td>7.3% <em>victoria</em></td>
</tr>
<tr>
<td>5. <em>hydei</em> 4.9</td>
<td>7.1% <em>melanogaster</em></td>
</tr>
<tr>
<td>6. <em>macroperta</em> 4.8</td>
<td>4.4% <em>longicornis</em></td>
</tr>
<tr>
<td>7. <em>melanogaster</em> 3.4</td>
<td>3.9% <em>nigrospiracula</em></td>
</tr>
<tr>
<td>8. <em>innubila</em> 2.7</td>
<td>3.4% <em>macropina</em></td>
</tr>
<tr>
<td>9. <em>tenebrosa</em> 1.7</td>
<td>3.1% <em>hydei</em></td>
</tr>
<tr>
<td>10. <em>californica</em> 1.5</td>
<td>2.5% <em>carbonaria</em></td>
</tr>
<tr>
<td>11. <em>azteca</em> 1.4</td>
<td>1.9% <em>nigrospiracula-like</em></td>
</tr>
<tr>
<td>12. <em>simulans</em> 1.0</td>
<td>0.5% <em>azteca</em></td>
</tr>
<tr>
<td>13. <em>grisea</em> 0.7</td>
<td>0.5% <em>Leucophenga varia</em></td>
</tr>
<tr>
<td>0.1% or less, 8 species</td>
<td>0.1% or less, 6 species</td>
</tr>
</tbody>
</table>
The data indicate that the large pseudoobscura populations (larvae and/or adults) in the mountains possibly control over-all abundance within the other species, at least at the traps, but by the criterion of the desert fauna, they have little effect on the number of species or on their relative abundance. It appears that the only interaction here is that of random crowding at the site of collection.

The two new species, one similar to pseudoobscura, and the other, a morphological and ecological sibling of nigrospiracula, will soon be described.

Hess, Oswald. Scute\(^8\) as \(Y\) suppressed lethal factor.

In the course of cytological investigations of spermatogenesis in \(X/O\ \alpha\) of \(D.\ melangaster\) (Meyer, Hess and Beermann, 1961) we found that \(sc^8/0\ \alpha\) are lethal. Four different \(X\) chromosomes carrying the \(sc^8\) mutation were tested, namely Muller-5 \((sc^5_{1B} InS \ y^a-sc^8)\), FM 4 \((y^\text{31d} sc^8dm B)\), sc\(^8_{bb} w^a\), and sc\(^8_{bb} w\). These males were crossed with \(y^2-su-w^a xa bb/0\ \alpha\). From these crosses in \(F_1\ XX/Y\) daughters and \(sc^8/0\) sons are expected in the ratio 1:1. The actual ratio found, however, from three crosses apiece, is shown in the following table:

<table>
<thead>
<tr>
<th>Paternal X</th>
<th>XX/Y</th>
<th>X/O</th>
<th>(\alpha)</th>
<th>(\alpha)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild Berlin (control)</td>
<td>1015</td>
<td>1278</td>
<td>100</td>
<td>126</td>
</tr>
<tr>
<td>M-5</td>
<td>1005</td>
<td>111</td>
<td>100</td>
<td>11</td>
</tr>
<tr>
<td>FM 4</td>
<td>1070</td>
<td>21</td>
<td>100</td>
<td>6,6</td>
</tr>
<tr>
<td>sc(^8_{bb} w^a)</td>
<td>1047</td>
<td>2</td>
<td>100</td>
<td>0,19</td>
</tr>
<tr>
<td>sc(^8_{bb} w)</td>
<td>1028</td>
<td>24</td>
<td>100</td>
<td>2,3</td>
</tr>
</tbody>
</table>

The penetrance of the lethal effect of \(sc^8\) in \(X/O\ \alpha\) is below 100\%. It varies between 89 and 98\% in our experiments. We have not yet established whether the observed percentage of \(sc^8/0\) break throughs (Durchbrüner) are characteristic for the type of \(sc^8\) chromosome used. Since dead pupae have not been found in the crosses the critical period of the lethal effect must be earlier in development.

The lethality of \(sc^8\) is partially suppressed by fractional \(Y\) chromosomes, as is shown in the following table for \(Y^S\):

<table>
<thead>
<tr>
<th>Paternal X</th>
<th>XX/Y</th>
<th>X/Y(^S)</th>
<th>(\alpha)</th>
<th>(\alpha)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild Berlin (control)</td>
<td>975</td>
<td>1044</td>
<td>100</td>
<td>107</td>
</tr>
<tr>
<td>M-5</td>
<td>1316</td>
<td>943</td>
<td>100</td>
<td>72</td>
</tr>
<tr>
<td>FM 4</td>
<td>1136</td>
<td>661</td>
<td>100</td>
<td>58</td>
</tr>
<tr>
<td>sc(^8_{bb} w^a)</td>
<td>1241</td>
<td>989</td>
<td>100</td>
<td>79</td>
</tr>
<tr>
<td>sc(^8_{bb} w)</td>
<td>1087</td>
<td>791</td>
<td>100</td>
<td>73</td>
</tr>
</tbody>
</table>

Similar ratios were found with \(sc^8/Y^S, Y^S\) and \(sc^8/Y^{bb-}\ \alpha\).

The most likely hypothesis from these results is that \(sc^8\) is a lethal variegated position effect. Lindsley et al. have found that after X-raying of \(D.\ melangaster\ \alpha\ with 3-4 kr 20-25\% of the resulting lethals are suppressed by the \(Y\) chromosome and therefore overlooked in the usual tests. Another argument for this hypothesis is, that the \(sc^8\) mutation meets the basic requirement for a \(V\)-type position effect,
January 1962 Research Notes

The X chromosome has an inversion with one break in the euchromatin and another in the heterochromatin. We did not find any significant difference between cultures maintained at 22, 25, and 28°C, but we do not consider this to be a strong argument against the hypothesis. As many investigators have found, the amount of heterochromatin, especially in the Y chromosome, is a much stronger modifying factor than temperature differences.

Hildreth, P. Influence of different Y chromosomes on secondary nondisjunction in D. melanogaster. Females heterozygous for a wild type X chromosome from a Samarkand stock and an X chromosome of the composition \( y^{+}sc^{a}B inv^{49}v w^{a}sc^{a} \) were tested for the frequency of X-chromosomal nondisjunction and segregation of the X's when Y chromosomes of different types were present in the females. The Y chromosomes used were (1) a normal unmarked Y, (2) \( sc^{a}Y \), (3) \( sc^{B}Y.B^{a} \) and (4) \( y^{+}BY \) (a chromosome which arose in one of our experiments and has not yet been analyzed). Since this was only a preliminary test no attempt was made to isogenize the stocks. Larger-scale experiments are planned in which these and other Y's will be used and the genetic background will be strictly controlled.

Individual female inversion heterozygotes, each bearing one of these Y chromosomes, were mated with males carrying a normal Y chromosome and having the X chromosome marked with \( y \) and \( w \). The corrected nondisjunction rate was highest, 66.1%, in females carrying the normal Y chromosome. The frequencies decreased to 59.3% in the presence of \( y^{+}BY \), to 54.9% in the presence of \( sc^{a}Y \), and to 49.8% in the presence of \( sc^{B}Y.B^{a} \).

Table I indicates the percentage of recovery of the chromosomes singly and in combinations. Because of the markers used it was not always possible to ascertain whether or not the Y chromosome from the female was present. This was true entirely for the unmarked Y and partially for the \( sc^{a}Y \) chromosomes.

Table I

<table>
<thead>
<tr>
<th>Percentage recovery of Y and X chromosomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>P female</td>
</tr>
<tr>
<td>-----------</td>
</tr>
<tr>
<td>( y^{2}B v w^{a} )</td>
</tr>
<tr>
<td>( Y )</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>( y^{2}B v w^{a} )</td>
</tr>
<tr>
<td>( y^{+}BY )</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>( y^{2}B v w^{a} )</td>
</tr>
<tr>
<td>( + + + + sc^{a}Y )</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>( y^{2}B v w^{a} )</td>
</tr>
<tr>
<td>( + + + + sc^{a}Y.B^{a} )</td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

...
In each instance the $y^2B_v w^a$ chromosome is recovered nearly as frequently as the wild type chromosome, indicating in these cases that the viability effects of the two chromosomes are similar. The recovery of the $Y$ chromosome from the female approached or surpassed 50% in those cases in which the presence of the $Y$ could be ascertained, indicating relatively good viability of individuals carrying the $Y$ chromosome.

When apparent nondisjunction of the $X'$s took place the $Y$ chromosome was always recovered in higher frequency than the two $X'$s, as had been observed by Sturtevant and Beadle (1936) in their tests of several different inversion heterozygotes for secondary nondisjunction. The inversions used by them were not as complex as the one used here, however. The more complex inversion would lessen the chance for pairing between the $X'$s and therefore decrease the opportunity for crossing over. This in turn would mean a low frequency of anaphase bridges to account for death of eggs containing these, and therefore should not contribute greatly to the frequency of patroclinous males in our experiment. A test of inversion heterozygotes without $Y$ chromosomes failed to produce patroclinous males in higher frequency than matroclinous females.

Another interesting aspect is the frequency with which the $X$ chromosomes are recovered singly with and without the $Y$ chromosome. According to random expectation when one $X$ chromosome is retained in the egg nucleus, then 50% of the time it should be the inverted $X$ and 50% of the time it should be the wild type $X$. When the inverted $X$ is retained, then 50% of the time the $Y$ chromosome should be retained with it. The same is true for the wild type $X$, thus the four classes should occur with equal frequency. As is seen in Table I, the frequencies of wild type $X$ chromosomes with and without the $Y$ are nearly equal but there is great inequality in the frequencies of $y^2B_v w^a$ chromosomes recovered with the $Y$ and without it. Table II gives the expected and the observed ratios for the nondisjunctional and other classes.

<table>
<thead>
<tr>
<th>P female</th>
<th>Y : XX</th>
<th>$++ + + : + + + / Y : y^2B_v w^a : y^2B_v w^a / Y$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected</td>
<td>1</td>
<td>1 1 1 1</td>
</tr>
<tr>
<td>Observed</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td>$y^2B_v w^a$</td>
<td>1.39 1.00</td>
<td>-</td>
</tr>
<tr>
<td>$y^2B_v w^a$</td>
<td>1.21 1.00</td>
<td>2.09 2.01</td>
</tr>
<tr>
<td>$y^2B_v w^a$</td>
<td>1.28 1.00</td>
<td>-</td>
</tr>
<tr>
<td>$y^2B_v w^a$</td>
<td>1.66 1.00</td>
<td>1.50 1.58</td>
</tr>
</tbody>
</table>

It appears that the rate of secondary nondisjunction is influenced by the $Y$ chromosome and that the normal $Y$ is associated with the highest degree of nondisjunction, while the $sc^8,Y,BS$ is associated with the least degree. The wild type $X$ chromosome is recovered as frequently with the $Y$ as without it, but the inverted
X is recovered approximately twice as often without the Y as it is with the Y. From the total recovery of the Y and each of the two X chromosomes it seems unlikely that viability differences could account for this latter effect. It is possible that some mechanism causes the Y to be lost frequently from its association with the y^2B v w^a chromosome but not from its association with the wild type X chromosome.

(This work was done under the auspices of the U. S. Atomic Energy Commission.)

Hiraizumi, Y. Low viability induction by the segregation distorder (SD) locus; preliminary note.

In a heterozygous SD male, SD causes a breakage in its partner chromosome, perhaps at SD^+. This broken chromosome is eliminated in some stage before fertilization, thus more than 50% (usually 95% or more) SD-bearing chromosomes are transmitted to the next generation. Here a question arises whether the SD^+ bearing chromosomes found in the F_1 generation are 1) those which were not affected at all by SD action or 2) those which recovered from the break. If 2) is the case, then we may expect some changes, perhaps viability reduction, in the SD^+ bearing chromosomes from heterozygous SD males. Accordingly, SD/cn bw (and SD^+/cn bw as a control) males were crossed to cn bw/In(2L) Cy cn bw females. In each set of experiment the cn bw chromosomes in the heterozygous SD and SD^+ males in F-generation were derived from a single, lethal free chromosome and the remaining genetic background had been uniformed before the present experiments. From the F_1 of these matings cn bw/In(2L) Cy cn bw males were chosen to cross individually to cn bw/In(2LR) Cy females, and the F_2 cn bw/In(2LR) Cy sibs from each F_1 mating were mated to test the homozygote viabilities of cn bw chromosomes in comparison with their Cy heterozygotes. For the significance test the observed percentage of cn bw homozygotes (= r) in the F_2 in each culture vial was transformed according to the relation \( r = \sin^2 R \). The results are summarized in the table. Figure in parenthesis is the percentage of cn bw homozygotes corresponding to each R (= average of R) value.

The lethal-bearing cn bw chromosomes (indicated as +1 lethal etc.) were excluded from computing R. Each experimental set was made at a different time, but in each set the cn bw chromosomes from the SD/cn bw males showed, on the average, reduced viabilities (p < 0.01). It is interesting to note that original SD (= SD-72 and SD-5; strong SD) lines caused more viability reduction than recombinant SD (weak SD) lines, although the difference was not statistically significant. The detailed mechanism for this is not yet fully understood, but a small deletion accompanied by the breakage-reunion event could be responsible.

Table

<table>
<thead>
<tr>
<th>Exp. set No.</th>
<th>Original SD</th>
<th>No. of cultures</th>
<th>Recombinant SD</th>
<th>No. of cultures</th>
<th>SD^+</th>
<th>No. of cultures</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>31.16</td>
<td>8</td>
<td>31.49</td>
<td>20</td>
<td>34.34</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>(26.8)</td>
<td>(27.3)</td>
<td>(27.3)</td>
<td>(31.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>31.69</td>
<td>12</td>
<td>32.50</td>
<td>38</td>
<td>33.52</td>
<td>37</td>
</tr>
<tr>
<td></td>
<td>(27.6)</td>
<td>(28.9)</td>
<td>(28.9)</td>
<td>(30.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>31.80</td>
<td>16</td>
<td>32.42</td>
<td>52</td>
<td>33.38</td>
<td>31</td>
</tr>
<tr>
<td></td>
<td>(27.8)</td>
<td>(28.8)</td>
<td>(28.8)</td>
<td>(30.3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>31.62±0.34</td>
<td>32.30±0.28</td>
<td>33.63±0.27</td>
<td>30.7</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Hoenigsberg, H. F., Y. García Cortés, and D. Ortiz Rubio. The degree of sexual preference in D. melanogaster Cy BL and the fitness associated with it.


Hollander, W. F. Two mosaics.

(1) From our stock of y w/Basc triploid females and Basc males, a well-formed male was obtained having the left half of the body yellow, left eye white, right eye bar apricot (= Basc). This male lived a week, but failed to produce any progeny. The simplest explanation seems to be simultaneous fertilization, by normal Y-bearing sperms, of a binucleate egg with one diploid nucleus and one haploid nucleus (suggestion of Peter E. Thompson). The resulting intersex side might have been shifted toward normal maleness by coexistence with the male side, but not enough to permit fertility.

(2) In our attached-X stock #82 a male was obtained with the left eye reddish-colored, similar to "coral," the right eye white (typical color of males of stock). Mated to 5 virgin sibs he produced 15 sons with the new red eye color, and 100 sons white-eyed. Next he was mated to attached-X white-eyed females, and produced 32 sons, all white-eyed and yellow-bodied. It appears likely that this mosaic was somatically mutant from white to coral (1), including also a small portion of the germ cells. The new eye color is being maintained.

During recent studies of sexual behavior in Drosophila melanogaster mutants the authors found D. melanogaster Cy/BL to establish very definite preferences both in various elements of courtship which prompt response, and in copulations. The male choice method consisted in placing the male to choose between two females, one of his own type and the other a wild D. melanogaster female. Such preferences were also present in D. melanogaster Cy Pm but to a lesser extent, and not for all elements of courtship. Nevertheless, the phenomenon in Cy Pm resulted in courtship discrimination with subsequent nonrandomness in mating but extended to those elements of courtship which most elicited the lowering of the female threshold barrier. Moreover, the authors completed the studies by making the female choice as well. The results, which will be published elsewhere, show female choice preferences like those already apparent by the male choice method. In other cases we found discrimination in the other direction indicating a lesser importance of the female behavior as the deciding condition in sexual preferences.
Hollander, W. F., and Michael F. Festing. Matings of roughex males (rux60d - see DIS 34:50) with attached-X females have produced 17 homozygous roughex daughters in 9447 progeny examined. Any associated sex-linked markers also became homozygous. Secondary non-disjunction from these females has been below expectation; further tests are in progress.

Hunter, Alice S. Abnormal sex ratio in wild Drosophila pseudoobscura. During recent months, collections of Drosophila have been made in various natural localities in the vicinity of Bogotá, Colombia. In one of these collecting sites, a relatively high percentage of Drosophila pseudoobscura has been found in the collections made over a period of four months. The site, which we call "Pine Woods," is located at an altitude of 2,700 meters, with an average temperature of 15°C. Since Bogotá is located at a latitude of 4° North there is little or no variation in temperature throughout the year. There are, however, two rainy seasons, one March-April and the other October-November.

An inherited "sex ratio" condition is well known for D. pseudoobscura, but since this results in the production of all female offspring it could not be related to the high percentage of males collected in the Pine Woods. These collections are made by sweeping over a bait which consists of fruit skins and wastes of a wide variety of local fruits such as banana, pineapple, papaya, guava, mora, oranges and curuba. The data follow:

<table>
<thead>
<tr>
<th>Month</th>
<th>Total collections</th>
<th>Females</th>
<th>Males</th>
<th>Chi Square</th>
</tr>
</thead>
<tbody>
<tr>
<td>July</td>
<td>13</td>
<td>38</td>
<td>173</td>
<td>87</td>
</tr>
<tr>
<td>August</td>
<td>24</td>
<td>93</td>
<td>237</td>
<td>64</td>
</tr>
<tr>
<td>September</td>
<td>44</td>
<td>724</td>
<td>1,000</td>
<td>44</td>
</tr>
<tr>
<td>October</td>
<td>37</td>
<td>282</td>
<td>605</td>
<td>118</td>
</tr>
<tr>
<td>Total</td>
<td>118</td>
<td>1,137</td>
<td>2,015</td>
<td>122</td>
</tr>
</tbody>
</table>

It is obvious that the deviation from the expected 1:1 ratio is large each month and also for the total. It seems to us that such an abnormal ratio in a total of over 3,000 flies is worthy of investigation. Therefore both field and laboratory studies have been started in order to ascertain the basis of this abnormal sex ratio.

Collections are being made at another site roughly 400 meters from the Pine Woods. These collections are being made from traps which contain pure banana bait. Although there is a slight preponderance of males in these collections, the sex ratio is much closer to 1:1. This suggests that a comparison should be made of the different methods of collection, different types of bait and different ecological conditions of the collecting sites.

As a start in the laboratory investigations, isolated wild females collected in the Pine Woods were grown at outdoor temperature under optimal food conditions. Counts of the F1 from 54 different females showed that 47 produced a normal 1:1 ratio, while only 7 females produced offspring which varied significantly from the expected 1:1 ratio of males to females. Of these, 4 showed a higher percentage of females, and 3 a higher percentage of males. This suggests that environmental rather than hereditary factors are involved in the abnormal sex ratio found in the wild populations of D. pseudoobscura.
From genetical and cytological analyses, it becomes clear that females of the strain derived from a male of wild Miyazu strain irradiated by X-ray (previously reported) have two translocations in addition to the chromosomal constitution of XXY: one is a half-translocation between X and II, the other a mutual translocation between the rest of II and III. In the half-translocation, the distal 1/3 of the left arm of II including cn is located at the end of X; the rest of II is broken at the middle of the left arm and translocated with III. Thus, some genes on the left arm of the original II are linking with X, and some with III. The strain can be preserved by following crosses; Y/Basc/T(1;2) x Y/Basc or Y/y m/T(1;2) x Y/y m (the translocation between II and III is always contained in those females). In various crosses of this strain, the total mortality reaches 82-89%. The details will be reported in the "Cytologia."

In an experiment where regular F₁ males (expected type) carried vermilion on the X-chromosome and were homozygous for brown eyes, it was found that among several white-eyed exceptional males whose phenotype is due to the loss of the bw+ gene or the entire Y carrying it, one was fertile. On being mated to virgins from a stock homozygous for brown all his male offspring have a phenotype (bw+) which proves that they have the bw+ gene (by covering bw/bw) as well as in their being fertile showing they have the Y chromosome. It is apparent that the loss of the Y chromosome occurred from the primordial tissue forming the eyes but did not occur in his germ tissue on either side.

Collections were made with the use of traps for three days in the middle of August, 1961. A total of 831 specimens belonging to 24 species was obtained. In Utoro, a northern side of the Shiretoko Peninsula, the collection showed that dominant species were represented by D. lacertosa, D. auraria, D. nigromaculata and D. okadai. In Habomai, lying south to Utoro at a distance of 100 km, D. nigromaculata was the only dominant species showing the frequency of 66% (Table 1). The difference in distribution between the two localities is mostly attributed to the flora in their habitats.

<table>
<thead>
<tr>
<th>Species</th>
<th>No. of flies</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Utoro</td>
</tr>
<tr>
<td>D. nigromaculata</td>
<td>93</td>
</tr>
<tr>
<td>D. lacertosa</td>
<td>157</td>
</tr>
<tr>
<td>D. auraria</td>
<td>132</td>
</tr>
<tr>
<td>D. okadai</td>
<td>89</td>
</tr>
<tr>
<td>D. testacea</td>
<td>24</td>
</tr>
<tr>
<td>D. histrioides</td>
<td>1</td>
</tr>
<tr>
<td>D. suzuki</td>
<td>3</td>
</tr>
<tr>
<td>D. coracina</td>
<td>12</td>
</tr>
<tr>
<td>D. brachynephros</td>
<td>2</td>
</tr>
<tr>
<td>D. ezoana</td>
<td>5</td>
</tr>
<tr>
<td>D. funebris</td>
<td>2</td>
</tr>
<tr>
<td>D. moriwaki</td>
<td>1</td>
</tr>
<tr>
<td>D. nipponica</td>
<td>1</td>
</tr>
<tr>
<td>D. tenuicaua</td>
<td>1</td>
</tr>
<tr>
<td>D. trivittata</td>
<td>1</td>
</tr>
</tbody>
</table>
(Kaneko, Shima, and Momma, Table 1--continued)

<table>
<thead>
<tr>
<th>Species</th>
<th>Utoro</th>
<th>Habomai</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>D. sexvittata</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>D. sp. (fenestrarum group)</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>D. spp. (two different species)</td>
<td>8</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>Scaptomyza graminum</td>
<td>0</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>S. polygonia</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Parascaptomyza disticha</td>
<td>0</td>
<td>29</td>
<td>29</td>
</tr>
<tr>
<td>Amiota variegata</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>537</strong></td>
<td><strong>294</strong></td>
<td><strong>831</strong></td>
</tr>
</tbody>
</table>

External feature:
- Male and Female; **Body**: Dark brown, about 2.5 - 3 mm long, with remarkable black stripes on mesonotum.
- **Head**: Eyes dark red, with short piles.
- Ocellar triangle and perioral black and one prominent bristle.
- **Mesonotum (Fig. E)**: Yellowish brown, with 4 black longitudinal stripes, inner pair interrupted at posterior and outer pair interrupted at anterior.
- Broaden black spot below intrascutal suture.
- Scutellum black, with posterior portion brownish yellow. Thoracic pleura largely dark brown spots. Humerals two. Acrostical hairs 8 rows. Cross distance of 3r, about third the length distance. Anterior scut. slightly divergent. Stero-index about 0.4.
- **Legs**: Brownish yellow, ultimate femora and tarsal joints dark brown.
- Preapicals prominent on hind tibia. Apicals on middle. Wings fuscous.
- **Wings (Fig. A)**: Three rather large and distinct spots in the wing. One spot distributes around the anterior cross vein starting from the proximal end of the wing, another one in the central part of the wing extending from radius 213 to media 314 and surrounding the posterior cross vein, and the last one extends from the distal end up to nearly the middle part of the wing, covering the distal parts of marginal cell and submarginal cell, with a small puncture in the region of marginal cell. The distal end of the last spot draws a curve forming a concave connecting media 1 and radius 415 and the central end forms also a concave. C-index about 1.3; 4V-index about 1.4; 4C-index about 1.4; 5X-index about 1.0; C1-bristles 2; C3-bristles on basal 7/10. Halter white.
Abdominal tergites (Fig. F): Brownish yellow, with black patches. 1 tergite brownish yellow, black at lateral corners; 2T brownish yellow, with broad caudal black band, which is deeply incised at middle and with 4 brownish yellow spots at lateral sides; 3-5T with 4 brownish yellow spots on each lateral segment and one rod shaped spot on each middle side. 6-7T brownish yellow.

Periphallic organ (Fig. B): Genital arch yellow, elongate, somewhat triangular at lower tip, and with about 25 strong setae, upper portion densely hairy. Clasper yellow, broad, narrowing basally, and with about 10 teeth in a row, and about 6 secondary teeth arranged in 2 rows. Anal plate yellow, narrow and broad below, separated from genital arch, and with about 26 stout hairs including numerous short hairs.

Phallic organs (Fig. C): Aedeagus elongate, apically swollen in lateral view, flattened and elliptical. Anterior paramere brownish yellow, elongated, fused to novasternum. Novasternum brownish yellow, and nearly quadrangle, and with a spine on the inner edge. Posterior paramere seems to be absent. Ventral fragma brown, semielliptic, and rounded at tip.

Egg-guides (Fig. D): Lobe yellow, tapering at tip, and with about 14 marginal brown teeth. Basal isthmus brownish yellow, thick and short.

Holotype: Male, Kwang nung, Kyngki province, South Korea, 1 Male, June, 1961.
Allotype: Female, collected together with holotype.
Paratype: Kwang nung, Male 16 and female 12.
Distribution: South Korea.
Collecting method: Net sweeping on the decayed tees.

Kaplan, William D., V. E. Tinderholt, and D. H. Gugler. The number of sperm present in the reproductive tracts of Drosophila melanogaster females. In studying radioautographs of the female reproductive tracts for the presence or absence of labelled sperm, it was noted that the number of sperm present was not so great as was expected on the basis of earlier published reports. The number of Fuelgen-positive sperm heads is much less than the impression given by a fully-packed seminal receptacle or the paired spermathecae in which the sperm, with their extremely long tails, are contained. A count of sperm heads in these structures gave a maximum of about 650 in eight females examined. A mass of sperm cells is also present in the vagina and, two hours after a single copulation, this mass contained about 300 sperm cells. The total in this one female was, however, 750. We are now studying the way in which this vaginal sperm mass is utilized to replenish sperm in the seminal receptacle and the spermathecae.
Kikkawa, H. Strain differences in proteolytic enzyme activities in D. melanogaster and strains. Strain differences seem to be due to qualities and quantities of the enzymes. Of interest is that a strong inhibitor of trypsin is contained in the body fluid.

King, R. C. Vitellogenesis in Drosophila. Cytochemical studies have shown that the mature ovarian oocytes of Drosophila melanogaster and D. willistoni contain large yolk spheres between 1 and 3 microns in diameter which belong to two classes. The alpha yolk sphere contains proteins (extractable from sections by pepsin, trypsin, or papain) and acidic lipids (which appear to be relatively unsaturated). The second type of yolk, comprising the beta spheres, is devoid of protein and contains (1) periodic acid-Schiff-positive polyglucosans which are extracted from sections by alpha amylase, (2) alcian blue-positive, acidic polysaccharides, and (3) lipids (which are relatively saturated). Under the electron microscope the dense alpha spheres are seen to be covered by a double walled envelope; whereas the pale beta spheres appear to be devoid of an enclosing membrane. Beta yolk spheres often coalesce with one another.

The ooplasm contains myriads of mitochondria and lipid droplets which are just above the limit of resolution of the light microscope. In electron micrographs the mitochondria are seen to be ellipsoidal, and many are embedded in the cortex of the beta spheres. The lipoidal bodies have a stellate appearance. Stacks of annulate lamellae of the sort illustrated in Growth 22:323, Fig. 26, and isolated filaments of endoplasmic reticulum occur commonly also. The background ooplasm contains proteins, polysaccharides, lipids, glycoproteins, lipoproteins, and ribonucleoproteins. The so-called periplasm of the egg represents a region filled with layers of membranes arising from convolutions of the plasma membrane.

Cytological observations indicate that during vitellogenesis a stream of cytoplasmic material flows in a posterior direction through the cells of the nurse chamber and finally into the oocyte through pores which connect all these daughter cells. These pores are so large that particles the size of mitochondria and lipoidal droplets easily pass through them. Alpha yolk spheres are first seen in the oocyte during stage 8. During maturation of these spheres the lipids they contain appear to increase in concentration and/or to become less soluble in organic solvents (perhaps through formation of lipoprotein complexes). Immature alpha spheres contain a considerable amount of polysaccharide which is later lost. The alpha spheres are believed to arise by the growth while in the ooplasm of membrane enclosed droplets a few tenths of a micron in diameter. These precursor particles may arise in turn from tiny blebs which are pinched off the envelopes of the nurse cell nuclei and are subsequently carried into the oocyte.

The beta spheres arise during stage 13 (some 4 hours after the alpha spheres first make an appearance) from smaller particles of similar morphology. It appears that as the beta sphere grows the carbohydrates it contains become more resistant to extraction, since in osmium-fixed material they are lost from the small particles but retained in the larger masses. However, formalin-containing fixatives retain the polysaccharides of the precursor particles. The source of these beta sphere precursors is currently under study.

Koch, R., and H. Burla. Dispersal rates in Drosophila subobscura and Drosophila obscura in relation to factors of environment, sex and age. According to the method of Sakai et al. (Evolution 12, 1958, pp. 93-101), the two species, Drosophila subobscura and D. obscura, have been compared in reference to their dispersal capacities.

At 25°C the dispersal rates are higher for D. obscura than at 18°C, whereas it is the inverse for D. subobscura at the two temperatures. The maximum dispersal
Research Notes DIS: 36

rates for D. obscura have been shown to be at lower relative humidities. Presumably higher humidities are more optimal and the activity is reduced in the range of such preferred humidities. Furthermore, the case is the reverse for D. subobscura, their activity being increased at higher humidities. For both species the activity is greater on fresh food than it is on old food and is greater when the flies are starved than when well fed. Younger adults are more active than the older, as are the males when compared with the females. For both species the activity has been shown to be reduced in low air-pressure.

In general, D. subobscura reaches higher dispersal rates than D. obscura, while D. obscura is more sensitive to all factors except air-pressure. Consistently, D. obscura has been shown to be the less resistant species in reference to unfavorable conditions.

Generally the dispersal activity was increased by conditions of environment which were considered not to be optimal for the respective species.

Koref-Santibañez, Susi.
A comparative study of courtship behavior in some species of the mesophragmatica group of Drosophila.

Courtship behavior has been analyzed in the following five species of the mesophragmatica group: D. viracochi from Machu-Picchu (Peru); D. mesophragmatica, from Machu-Picchu (Peru); D. gasici, from Arica (Chile); D. pavani, from Bellavista (Chile) and D. gaucha from Rio Grande do Sul (Brazil).

The general courtship pattern in all five species is very similar, and follows the ritual described by Spieth (1952) for other species of the subgenus Drosophila. Nevertheless, there are differences which allow the individualization of each species. Thus, D. viracochi males have a slower wing vibration movement; D. gasici males circle and touch antennae of females much more profusely than do those of other species. D. mesophragmatica males and females display very little activity and all their movements are slow. Another characteristic for each species is the duration of copulations, which are significantly different in all.

When, by means of the "male choice method," individual males are allowed to discriminate between a female of their same species, and a female of any of the other four, the following facts are observed:

a) The general activity of both males and females increases significantly over that observed when only individuals of the same species are together.

b) All males court their own females for a longer period than they do foreign females.

c) Only some courtship elements are used significantly more towards their own females, while others do not seem to be discriminative. Thus, D. gasici males discriminate against D. gaucha, D. viracochi, and D. mesophragmatica females, using all courtship elements more towards D. gasici females; D. pavani males discriminate when confronted with D. viracochi and D. gasici; D. gaucha males discriminate against D. viracochi females while D. viracochi and D. mesophragmatica males use all elements equally towards their own or towards foreign females.

d) As regards mating, the males of almost all the species copulate only with the sister females. Only D. pavani and D. gaucha mate almost indiscriminately with one another.

The comparative analysis of the courtship behavior of the different species included in the mesophragmatica group agrees with the phylogenetic relationships previously determined by the morphologic and cytogenetic studies (Brncic and Koref-Santibañez, 1958, and Brncic, 1959). It may be postulated, that as regards divergence of courtship behavior, D. gasici is the most distant, as both males and females discriminate highly, and the ritual itself is the most diversified. D. pavani and D. gaucha, which are the most closely related (they are sibling species), discriminate very slightly against each other, but markedly against the other species. The apparent lack of preference shown by D. viracochi and D. mesophragmatica males may be due to their general low activity, or to the fact that in them, the females are the more discriminate.
The higher activity of males and females of each of the five species when confronted with individuals of any of the other may be tentatively interpreted as following: the males may receive repulsory stimuli from the foreign females which may increase their excitation and obliges them to a greater activity, raising also the stimuli threshold of the females, thus conditioning longer courtship time and greater utilization of all the elements which make up the courtship ritual.

(Partially supported by a grant from the U. S. Atomic Energy Commission: Contract AT (30-1) 2465.)

Lefevre, G., Jr., and Ulla-Britt Jonsson. D. melanogaster females that mate with two different males usually produce offspring by both males. However, as various investigators have noted, a considerable degree of individual variability is evident in the results. As a consequence, a number of differing conclusions have been offered in regard to questions of sperm displacement, sperm mixing, and the sequence of sperm utilization. Reinvestigating this problem, we have identified an important source of such variability, and we are now able to define some consistent features in the activities that follow double matings.

Individual virgin males were encouraged to mate in succession with several 2 to 3-day-old virgin females. Males will often proceed energetically with their task, fertilizing as many as 5 successive females in a period of 3 to 4 hours. Then, each female of the sequence (which we designate #1, #2, #3, etc.) was presented with a second, genetically different, virgin male. If a remating did not ensue during an observation period of about 4 hours, the female was left with the second male until the next day.

As a rule, #5 females, as well as the majority of #4 females, mated for the second time almost as readily as they did originally as virgins. The exceptions, particularly with #4 females, in which remating did not take place usually occurred when the first male had spent an unduly long time carrying out the sequence of matings. The #3 females, in contrast, remated much less readily than did #4 and #5 females; nonetheless, some were receptive to a second male soon after the completion of the first mating, others within 24 hours. The #1 and #2 females, however, rarely remated the same day, but a reasonable number of rematings occurred within 24 hours.

After an observed remating, or after having been with a second male for 24 hours, each female was removed to a new vial and then subcultured, generally at 2-day intervals, for 10 or 12 days. The resulting offspring were counted and classified as to paternity, and the hatches were compared with those obtained from females that had mated only once. Two striking conclusions arise from the comparisons: (1) once-mated females produce just as many offspring as do twice-mated females, and (2) after a remating, the number of progeny sired by the first male is always less than he would have produced had the female not remated with a second male.

Additional females were dissected at intervals after mating, ranging from a few minutes to a few days. Soon after mating, the ventral receptacle and the two spermathecae are completely filled with sperm in #1 and #2 females, most often noticeably less full in #3 females (with considerable variation), more sparsely filled in #4 females (indeed, frequently, they were empty), and almost always empty in #5 females (occasionally, a very few sperm were observed). Further, the number of sperm transferred by a male in his first or second mating, perhaps three or four thousand, far exceeds the capacity of the female sperm storage organs. Long after they are completely full, great quantities of motile sperm can be seen in the uterus, oviduct, and even in the ovary. Rough estimates indicate that only between 10 and 20% of the sperm deposited in a #1 female are actually stored, but perhaps as many as half of the stored sperm are subsequently used in fertilization. In #3 females, the on the other hand, relatively fewer excess sperm are present; yet, occasionally their storage organs are about as full as in #1 and #2 females. In #4 females, virtually all of the smaller number of sperm transferred are stored, and in #5
females the few sperm that are sometimes transferred find their way into the receptacle and spermathecae within a matter of minutes.

Dissections of males following the various matings showed that the decline in the number of sperm transferred with each successive mating was not so much a matter of exhausting the seminal vesicles of sperm, but rather was correlated with the condition of the accessory glands. Even after five matings, the vesicles have many motile sperm left, but the accessory glands are completely collapsed and devoid of secretion. When the sequence of matings proceeds more slowly than usual, as it sometimes does, or when the male is deliberately rested between matings, then even in a fifth mating an appreciable number of sperm may be transferred.

Stained whole mounts of dissected female sperm storage organs always showed sperm distributed throughout the receptacle. The sperm heads do not congregate at the distal blind end, but are dispersed more-or-less randomly throughout the length of the receptacle, and in fact may be identified "inbound" and "outbound" at all levels. Even during the period shortly after mating when sperm are being stored and many are still in the uterus, outBound sperm heads can be detected in the basal end of the receptacle near the entrance. Thus, a circulation of sperm appears to exist in the receptacle, so that even as some sperm are coming in, others are going out. Clearly, after sperm enter the receptacle, they do not simply proceed along as far as they can go and remain there until all the space in the receptacle is occupied; rather, throughout the period when sperm are in the uterus, we believe that sperm are continually entering the receptacle, reversing direction at any level, and even returning to the uterus, perhaps later to re-enter the receptacle, perhaps not. Eventually, the excess sperm thrash off in unrewarding directions (up the oviduct or out the vagina), leaving only the better oriented ones finally accommodated within the storage organs. It is abundantly evident, in any event, that the female sperm storage organs are normally filled to capacity by a single copulation.

Upon remating, the sperm circulation (which appears to continue between matings in the same manner) results in the emergence from the receptacle of previously stored sperm where, in the uterus, they are diluted by the vastly greater quantity of sperm newly deposited by the second male. The likelihood of re-entrance of the first sperm is thereby greatly reduced. Finally, when the excess sperm have vanished and all sperm possible have been stored, sperm from the first male still present in the receptacle are quite diminished in number.

Remating surely does not result in the storage of a double quantity of sperm, nor do the two kinds of sperm form any sort of layers in the receptacle; but rather remating results in the displacement of a greater or lesser proportion of the sperm originally present. It is not apparent from dissections that a similar activity occurs in the spermathecae, but judging from the offspring produced by remated females, which sometimes contain very few sired by the first male, we are inclined to suspect that to some degree it does.

Lefevre, G., Jr., and Ulla-Britt Jonsson. In 1949, Novitski and Rush (Biol. Bull. 97:150-157) reported that fertilized D. melanogaster females can be desemnated by an exposure to sub-zero temperatures. An effective treatment was -10° C. for 10 minutes. At the same time, they stated that males subjected to the same treatment were not affected in regard to their subsequent fertility, at least.

It seemed paradoxical that mature sperm stored in the female should be killed by cold treatment, while similar sperm stored in the male should be immune. Reinvestigating the effect on males, we have found that exposure to -10° C. for 10 minutes does in fact inactivate all of the fully mature, motile sperm stored in the seminal vesicles of the male, exactly paralleling the effect on mature, motile sperm stored in the ventral receptacle and spermathecae of females. For a period of 24 hours or so after treatment, no motile sperm can be detected in the male reproductive organs, nor are any sperm transferred to the female during
copulation. The male regains fertility, however, as the apparently more resistant, less mature, immotile sperm in the testis mature and enter the seminal vesicle, becoming available then for insemination. If the males are first irradiated with 4000r, then exposed to the cold treatment, a virtually complete sterility occurs from which there is little or no recovery.

Dissections indicate that the treated male is unable to expel from the vesicles the mass of dead sperm produced by the cold shock. This results in a greater or lesser obstruction to the passage of newly maturing sperm, and occasionally motile sperm can be seen in the testis itself, having been unable to descend into the vesicle. Thus, none of the treated males regain a normal degree of fertility, and in particular are unable to inseminate more than one or two females within a 24-hour period. There is no loss of sex drive, however, and treated males will copulate repeatedly without transfer of sperm. In fact, unless such males are removed from the presence of females and rested between matings, it is very rare to find a given male capable of successfully inseminating more than one female. It would appear that the dead sperm masses in the seminal vesicles so impede the passage of motile sperm into the ejaculatory duct that an appreciable period of time is required to build up the volume of sperm required for a normal insemination.

Cold-treated females, on the other hand, are able to expel the dead sperm from their sperm storage organs, so that following a reinsemination, there is no significant effect of subsequent fertility.

Lewis, E. B. Salivary gland chromosome analysis of segregation distorter lines.

Analysis of segregation distorter chromosomes SD5 and SD72 (of Hiraizumi, Crow and Sandler) indicates that each carries an inversion in the distal part of the right arm of the second chromosome apparently identical with In(2R)NS. The SD5 chromosome carries an additional inversion in 2R having one breakage point somewhere in the region extending from 45C to 45F and another in region 49A. The SD72 chromosome lacks the latter inversion but has, in addition to In(2R)NS, a small pericentric inversion with one breakage point in the euchromatic region of sections 39 or 40 of 2L and the other breakage point close to or within euchromatic section 42A of 2R.

Lovellette, E., and F. Ratty. Comparisons of inbred and random bred larval survival to 1200r.

The following experiments were conducted to determine whether first instar larvae of various genotypes express a differential survival to acute X-irradiation. This work compares the survival of four lines including a random bred (line 1), a hybrid formed from random females and inbred males (line 2), a hybrid formed from inbred females and random males (line 3), and an inbred (line 4).

Five hundred first instar larvae were placed on a one inch plaque of standard Drosophila medium, and then exposed to 1200r (220 kv, 20 ma) or used as controls. Survival is herein defined as the proportion of larvae developing into the adult stage. The results presented in the following table are based upon the average number of larvae which hatched from an original plaque of 500.

<table>
<thead>
<tr>
<th>Line</th>
<th>Total sample</th>
<th>CONTROL (avg./500) Total</th>
<th>Total Sample</th>
<th>IRRADIATED (avg./500) Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>7,000</td>
<td>214 211 425</td>
<td>11,000</td>
<td>159 195 354</td>
</tr>
<tr>
<td>2</td>
<td>7,000</td>
<td>205 217 422</td>
<td>11,500</td>
<td>107 125 232</td>
</tr>
<tr>
<td>3</td>
<td>7,000</td>
<td>220 206 426</td>
<td>9,500</td>
<td>72 120 192</td>
</tr>
<tr>
<td>4</td>
<td>7,000</td>
<td>214 219 433</td>
<td>14,500</td>
<td>77 113 190</td>
</tr>
</tbody>
</table>

line 1(random x), line 2(random x inbred), line 3(inbred x random), line 4(inbred x).
These results indicate that the average total survival among the four controls is quite uniform. In the irradiated samples survival is highest in line 1, while the groups derived from the inbred strain do not survive as well.

Possible interpretations of these observations might be:

1. Differences in the rates of development which would result in lines derived from the inbred group being in a different stage of development at the time of irradiation and thereby having a lower survival. Critically timed studies on first instar larvae of ages 2, 4, and 6 hours indicate this effect is probably not relevant.

2. If survival is dependent upon a maternal effect the survival of line 1 would be expected to approximate that of line 2, since they both have random bred mothers, while that of line 3 should approximate that of line 4, both having had inbred mothers. This relationship is indicated for the combined survival of lines 3 and 4; however, the survival of line 1 is significantly higher than line 2 which does not support a strict maternal effect hypothesis. The survival of irradiated females from line 2 does not differ significantly from those of lines 3 and 4 which also would not be indicative of a maternal effect.

3. Possibly survival is related to the genotype in a particular line. Differential survival of lines 2 and 3 seems to indicate that the males of line 3 do not survive as well as those of line 2. These males differ only with respect to their X chromosome. This suggests that lower survival is partially a function of the X chromosome from the inbred line—possibly related to the fixation of deleterious recessives therein. However, this explanation would not account for the high survival of line 4 females in relation to those of lines 2 and 3.

4. In addition, survival is probably also related to epistatic effects between the autosomes and the X chromosome of the inbred line. When these chromosomes are homozygous they tend to increase survival, as indicated by the comparative values of lines 3 and 4.

Lüönd-Luchsinger, S.
The riboflavin content in Malpighian tubules of D. hydei.
Chromatographic and fluorometric methods were used in determining the riboflavin quantities in the Malpighian tubules of larvae, pupae and imagos. Two maxima were found, one at pupation time and the other at the time of eclosion. A sex difference becomes apparent only in imagos, where females contain about twice the quantity of riboflavin as males. Adding riboflavin to the standard food results in a strong increase of the substance in Malpighian tubules of larvae and pupae. On the other hand, the feeding of riboflavin to imagos leads to almost no increase of this substance in their Malpighian tubules.

Malogolowkin, Ch. A new sibling species of the D. willistoni group.
A new species of the subgenus Sophophora, morphologically very similar to D. willistoni and D. paulistorum has been found in the states of Guanabara, Rio de Janeiro, Bahia, Salvador and in Pernambuco, Recife. This species crosses to D. willistoni and to strains of D. paulistorum from the Andean-South-Brazilian group of species. This species is being studied at the Department of Zoology of Columbia University and a formal description, together with genetic and cytological data, will be published elsewhere.

(This investigation is being supported by a fellowship from the Pan American Union.)

Malogolowkin, Ch. A new transitional race in Drosophila paulistorum.
The species Drosophila paulistorum is known to be a complex of six races or incipient species sharing varying degrees of reproductive isolation. Five of the races produce completely sterile F1 hybrids, if they can be crossed at all. The sixth race, termed Transitional by Dobzhansky and Spassky, produces fertile hybrids with
at least some strains of the other races. Now, the Amazonian race, which lives in the northern part of South America, from Belen to Panama, showed very strong reproductive isolation from the Andean-South-Brazilian race, which occurs from Colombia and Peru, to southern Brazil. Now collections made by myself in Central and Northeastern Brazil have disclosed the existence of a new Transitional race, which crosses and yields fertile hybrids of both sexes with the Amazonian as well as with the South-Brazilian strains. Strains of the new Transitional race have been isolated from populations of Ceara (Maranguape), and Bahia (Salvador), and may occur in other regions as well.

(This work has been assisted by grants from the Conselho Nacional de Pesquisas de Brazil and from the Rockefeller Foundation.)

Mettler, Lawrence E. Fertility relationships of recombination-hybrid males from the cross of D. mojavensis and D. arizonensis. Baker (1957) demonstrated that hybrid males from the cross arizonensis-mojavensis were sterile and that the reciprocal mating (mojavensis-arizonensis) produced partially sterile males. Population studies have indicated that introgressive (recombination) hybrid males may be partially fertile when the initial cross is arizonensis-mojavensis. The present investigation is to determine if the sterility is due to a simple X-Y chromosome unbalance, or if autosomal recombination and/or the cytoplasm influences fertility. The acquisition of a spontaneous white-eyed (X chromosome) mutant, which is apparently closely linked to the region in the X chromosome which differs in the two species by a paracentric inversion, makes such a study possible.

Cross A was arizonensis (white eye)-mojavensis and cross B was the reciprocal mojavensis-arizonensis (white eye). The F1 females from these two crosses were backcrossed to mojavensis males and to arizonensis (white eye) males. The 4 backcrosses produced 8 classes of male progeny in respect to the cytoplasm and the X and Y chromosomes:

| Cross A backcrossed to arizonensis | ariz | ariz | ariz | 4.0 |
| Cross B backcrossed to arizonensis | ariz | ariz | moja | 4.0 |
| Cross A backcrossed to arizonensis | ariz | moja | ariz | 4.0 |
| Cross B backcrossed to arizonensis | ariz | moja | moja | 6.0 |
| Cross A backcrossed to mojavensis | moja | ariz | ariz | 24.0 |
| Cross B backcrossed to mojavensis | moja | ariz | moja | 20.0 |
| Cross A backcrossed to mojavensis | moja | moja | ariz | 42.0 |
| Cross B backcrossed to mojavensis | moja | moja | moja | 77.0 |

Fertility tests were made by placing 100 males of each class individually with 3 virgin mojavensis females. The number (per cent) of those cultures producing offspring are listed above.

Those males carrying mojavensis Y chromosomes and X chromosomes are more fertile. The cytoplasm appears to have some effect especially when the X and Y chromosomes are both mojavensis. The fact that the combination moja-ariz-ariz (which is similar to the F1 sterile males from the cross arizonensis-mojavensis) shows 24 per cent fertility indicates an influence of autosomal recombination. Crossing over cannot be ruled out. The arizonensis arrangement is a simple inversion and it is being identified by a mutant locus. Certain backcross progeny with white eyes may actually carry the mojavensis X chromosome arrangement. If this is true the problem becomes more interesting. It would mean that the sterility loci are closely related to those loci differing by an inversion in the two species. Experiments are now in progress to verify these results and to ascertain the amount of crossing over.
Mettler, Lawrence E. Locating mutants by crossing species with chromosome differences.

An autosomal recessive mutant was recently found in D. arizonensis. In order to ascertain the autosome in which the mutant resides, the homozygous mutant strain was crossed to D. mojavensis (which differs from arizonensis by paracentric inversions in two of the four major autosomes). The F₁ of this cross was wild type. The F₁ hybrid females were backcrossed to homozygous mutant arizonensis males. Half of the progeny of this cross showed the mutant phenotype. Those flies showing the mutant phenotype were intercrossed and the salivary chromosomes of 20 of the progeny larvae were examined to determine which chromosome (2nd, 3rd or neither) was always homozygous for the arizonensis arrangement. All third chromosomes were such. Thus, the mutant was found to reside in this chromosome.

Mettler, Lawrence E. Drosophila pachea.

A single record of 2 males and 2 females collected about 3 miles north of Hermosillo, Sonora, Mexico (August, 1941) led to the description of Drosophila pachea Patterson & Wheeler 1942. The species was placed in the hydei subgroup of the repleta group. These flies did not breed in the laboratory. Recently (October, 1961) 4 males were collected near Hermosillo and 1 female near San Felipe, B. C., Mexico. These were classified as pachea by Dr. W. B. Heed. The female produced a few eggs on banana media but they did not hatch. The egg is rather small (1.76 mm. including filaments) with 2 relatively large, flat blade-like filaments which are as long as the body of the egg. Since all described eggs of the repleta group are characterized by 4 thread-like filaments, and since pachea lacks the general feature of mesonotum spotting, it is suggested that this species might belong in the subgenus Sophophora instead of Drosophila. Further collections and possible culturing will be attempted to help ascertain its true position.

Milkman, R. D. cv ultral relaxants in the progenies of wild inseminated females.

An experiment similar to a previous one in Ann Arbor (Science 131:225-226) has been conducted on the progenies of 29 wild inseminated Drosophila melanogaster females collected at Syracuse. Five hundred flies in the F₃ of each line were examined for posterior crossvein defects. The distribution of frequencies of such defects implies a polygenic basis for this deviant phenotype. The number of crossveinless flies of a total of 500 examined are presented, ranked as follows: 30, 17, 13, 12, 10, 6, 4, 3, 3. In addition, 5 strains contained 2 cve flies, 7 contained 1, and 8 contained none. These results are consistent with previous findings. Attempts to select true-breeding cve strains from the top 5 strains are in progress.

The analysis of a polygenic cve strain selected from the progeny of a wild inseminated female in the Ann Arbor experiment shows that all autosomes are involved. The second chromosome is more important, its homozygous state being a necessary and sufficient condition for the appearance of crossvein defects. Two genes between
the Star locus (1.3) and the sternopleural locus (22.0) seem to be involved. The X chromosome, on the other hand, favors the production of normal crossveins more than the X chromosome in Oregon R. In contrast to a cve strain previously analyzed, not all the alleles have increased expression at 18°, although some of them do.

Milkman, R. D. Protection against phenocopying by pre-treatment at high temperatures.

Resistance, both to the production of crossvein defects and to death at 40.5° C., has been described for pre-treatment in the 31°-38° range. Although temperatures throughout this range are comparably effective in conferring resistance to death at the higher temperature, a high temperature coefficient has been found for conferring resistance to transfiguration. The Q1 seems to be around 1.3 based on comparison of the effectiveness of treatment at various durations at 38.5°, 36.5°, 34.5°, and 32.0°. It is necessary to return pupae to room temperature for a short time, such as 2 minutes, in order for the protection to take effect. Pre-treatments lasting as little as 10 seconds at 36.5° have a measurable effect. Short pre-treatments at 40.5° also have a protective effect in terms of subsequent treatments at the same temperature, if the interval is of the order of 2 hours. Longer pre-treatments at 40.5° do not protect in the same way. It is possible to think of several states of a protein, whose interconversion is akin to denaturation, as the basis for these phenomena, and experiments are in progress to test this hypothesis further.

Momma, E., A. Kaneko, and T. Shima. The sensitivity for X-rays to spermatogenesis was analysed preliminarily. A total of 663 pupae was irradiated with 2000r in various developing stages, and the rate of their emergence was examined. The results are shown in Table 1. All the pupae irradiated within 20 hours after pupation died before the emergence. With the pupae irradiated 20 hours or more after pupation, emerged flies tended to increase gradually in number. Normal ratio of emergence was observed in pupae irradiated 40 hours or more after pupation. A few decrease of the ratio was observed in pupae irradiated 58 and 122 hours after pupation.

Table 1

<table>
<thead>
<tr>
<th>Age of pupae at irradiation after pupation (hours)</th>
<th>No. of irradiated pupae</th>
<th>No. of emerged flies</th>
<th>Rate of emergence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-15</td>
<td>127</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>16-25</td>
<td>77</td>
<td>0</td>
<td>0.1</td>
</tr>
<tr>
<td>26-35</td>
<td>41</td>
<td>12</td>
<td>18</td>
</tr>
<tr>
<td>36-45</td>
<td>29</td>
<td>18</td>
<td>27</td>
</tr>
<tr>
<td>46-55</td>
<td>30</td>
<td>14</td>
<td>29</td>
</tr>
<tr>
<td>56-65</td>
<td>89</td>
<td>32</td>
<td>67</td>
</tr>
<tr>
<td>66-75</td>
<td>26</td>
<td>12</td>
<td>22</td>
</tr>
<tr>
<td>76-85</td>
<td>60</td>
<td>31</td>
<td>60</td>
</tr>
<tr>
<td>86-95</td>
<td>11</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td>96-105</td>
<td>72</td>
<td>25</td>
<td>61</td>
</tr>
<tr>
<td>106-115</td>
<td>8</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>116-125</td>
<td>4</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>126-135</td>
<td>37</td>
<td>13</td>
<td>32</td>
</tr>
<tr>
<td>136-145</td>
<td>3</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>146-155</td>
<td>44</td>
<td>31</td>
<td>37</td>
</tr>
<tr>
<td>156-160</td>
<td>5</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Total</td>
<td>663</td>
<td>203</td>
<td>382</td>
</tr>
<tr>
<td>Control</td>
<td>558</td>
<td>243</td>
<td>496</td>
</tr>
</tbody>
</table>
Moree, Ray. Relative fecundity involving the e locus in *D. melanogaster.*

Male and female carriers of +/+ and e11/e11 were crossed in all possible combinations, each combination being made in a separate culture bottle. The 9 combinations were distributed over 160 bottles in a (1:2:1)² ratio. To minimize larval inviability each culture contained but 5 parental flies of each sex. The results in terms of total number of progeny per combination were as shown in Table 1. Dividing twice the number of homozygotes by the number of heterozygotes gave the relative fecundity coefficients shown in Table 2. Corrections for a small amount of larval inviability changed the fecundity coefficients only very slightly. As a matter of interest the frequencies of the progeny genotypes +/+ and e11/e11 were, in that order, 0.25, 0.52 and 0.23. The requisite statistical tests (made on the numbers and on the logarithms of numbers when that was necessary) support the obvious conclusion that the heterozygotes are heterotic and that the relative fecundities for the genotype sequence +/e11 > +/+ > e11/e11. This sequence is similar to that for relative viabilities (Moree and King, Genetics, in press) and helps to explain the long known fact that the e alleles are maintained at low frequency for long periods in cage populations.


It has been reported that effect of irradiation on the mutation rates changes depending on the different stages of germ cells. Paying attention to this point, we investigated the shift of the sex-ratio in the progeny of males irradiated at different stages of the male germ cells in *D. melanogaster.* Using the isogenic Oregon-R wild strain, male flies of 4½ hour-old were irradiated with 1000 r, 2000 r and 3000 r and crossed with the same wild type females of about 3 day-old. The crossings were made immediately after irradiation, and every other day thereafter with new virgin females, continuing up to 16 days after irradiation to obtain 8 different classes. Every class was divided into four groups, one control and three treated; thus 32 groups in total were made for 8 classes. In the next generation, the number of males and females were noted and the sex-ratio in each group was calculated. The counted flies amount to about 700,000, and the sex-ratio in each group changes dependent on the dose as well as the germ cell stage. The sex-ratio shifts to the lowest level in the group where progeny come from males of 6 - 8 days after irradiation (class IV). In table, regression coefficients are given, showing relationship between sex-ratio and dose with respect to every class. It can be said that in each class sex-ratio depresses in proportion to dose, and the coefficient is - 0.0237 per 1000 r in the average.

### Table 1

<table>
<thead>
<tr>
<th>d'</th>
<th>++</th>
<th>+/e11</th>
<th>e11/e11</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>++</td>
<td>17795</td>
<td>19795</td>
<td>9011</td>
<td>37452</td>
</tr>
<tr>
<td>+/e11</td>
<td>8646</td>
<td>83476</td>
<td>30222</td>
<td>151150</td>
</tr>
<tr>
<td>e11/e11</td>
<td>7504</td>
<td>15386</td>
<td>7332</td>
<td>37409</td>
</tr>
</tbody>
</table>

### Table 2

<table>
<thead>
<tr>
<th>d'</th>
<th>++</th>
<th>+/e11</th>
<th>e11/e11</th>
</tr>
</thead>
<tbody>
<tr>
<td>++</td>
<td>0.897</td>
<td>1</td>
<td>0.724</td>
</tr>
<tr>
<td>+/e11</td>
<td>0.894</td>
<td>1</td>
<td>0.952</td>
</tr>
<tr>
<td>e11/e11</td>
<td>0.896</td>
<td>1</td>
<td>0.835</td>
</tr>
</tbody>
</table>

### Regression coefficients (b)

<table>
<thead>
<tr>
<th>Class</th>
<th>b/1000 r</th>
<th>Class</th>
<th>b/1000 r</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>-0.00597</td>
<td>V</td>
<td>-0.02506</td>
</tr>
<tr>
<td>II</td>
<td>-0.01473</td>
<td>VI</td>
<td>-0.02702</td>
</tr>
<tr>
<td>III</td>
<td>-0.03495</td>
<td>VII</td>
<td>-0.01969</td>
</tr>
<tr>
<td>IV</td>
<td>-0.06013</td>
<td>VIII</td>
<td>+0.00277</td>
</tr>
<tr>
<td>Average</td>
<td>-0.02371</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Moriwaki, D., and H. Ikeda.
Disturbance of "sex-ratio" condition by X-ray irradiation.

In several species of Drosophila, "sex-ratio" condition has been analysed in various ways. "Sex-ratio" flies of D. bifasciata in Japan have been kept more than about sixty generations in our laboratory, producing almost only female progeny. The note reported here concerns whether X-ray irradiation can disturb "sex-ratio" condition in D. bifasciata or not. Normal males irradiated by X-rays (2000r, 4000r, 6000r) in each generation were mated with "sex-ratio" females successively for several generations. Their progenies were tested in each generation, but no male offspring could be detected. On the other hand, "sex-ratio" females which had been exposed to X-rays (2000r, 3000r) were mated with normal untreated males over a series of generations. Although no male appeared in the F1 generation, in the F2, for the first time, twelve per cent of the treated females raised progenies comprising some degree of males. Using the sister flies of the exceptional males without treatment, sex ratio in the next generation was examined, where some were realized as quasi-normal ratio and others behaved in the manner typical to the original "sex-ratio" strain. The result indicates that X-ray irradiation can inactivate the "sex-ratio" factors in egg cytoplasm originated from "sex-ratio" females. Further it is almost similar to results obtained in the previous investigations on temperature cure of the "sex-ratio" condition in D. bifasciata (Magni, 1953; Moriwaki and Kitagawa, 1957).

Mukai, T., and S. Chigusa.
Radiation-induced mutation rates of polygenes controlling the number of sternopleural bristles in D. melanogaster.

The males of an isogenic line (Burick's W160) and an inbred Oregon R (Hiraizumi's M-Oregon) were acutely irradiated with X-rays and Y-rays at 500r. Immediately after irradiation, the irradiated males were mated to the females of the same lines. The numbers of sternopleural bristles in females and males which hatched on or before the 13th day after the mating were scored. Therefore, the heterozygous effects of radiation-induced mutations were tested. The experiments are still in progress, but the results at hand are reported here.

The data of females only were analysed by using a technique in which the means, variances, and the third moments about the means of the distribution patterns of sternopleural bristle numbers are used. The summarized results are presented in the table.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Isogenic W160</th>
<th>Inbred Oregon R</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>X-rays</td>
<td>Y-rays</td>
</tr>
<tr>
<td>No. of genomes tested</td>
<td>1088(975)</td>
<td>1653(1392)</td>
</tr>
<tr>
<td>No. of mutations per individual</td>
<td>0.284</td>
<td>0.112</td>
</tr>
<tr>
<td>*Variance increase rate</td>
<td>4.87x10^{-4}/r</td>
<td>1.98x10^{-4}/r</td>
</tr>
<tr>
<td>**Mutation rate</td>
<td>1.14x10^{-6}/locus/r</td>
<td>0.41x10^{-6}/locus/r</td>
</tr>
</tbody>
</table>

* heterozygote basis
** assuming 500 loci
( ) no. of tested genomes in the control
To our surprise, the relative biological effectiveness of X-rays to \( \gamma \)-rays turned out to be about three. The detailed studies of this problem are now under way. The polygenic mutation rate is higher than that of major genes as previously described by Burdick and Mukai (1958), and the variance increase rate is also higher than the estimates obtained previously by several investigators. This is supposed to have been caused by the different response of the males and females to the irradiation, i.e., the variance of females was increased by irradiation while that of males was not increased but decreased. In spite of this phenomenon, they used pooled data of the males and females.

Mukherjee, A. S., and R. C. Strohman. A preliminary study on the chromatographic behavior of the heterozygous and homozygous conditions of a mutant and that of wild type Prospilina melanogaster.

In an attempt to make a comparative study of the chromatographic pattern of mutants in heterozygous and homozygous conditions we selected vestigial due to the following reason. In the description of vestigial (vg) wing mutation it has been reported that this mutant is temperature sensitive.1 The wings are completely vestigial at room temperature or below, but at 30°C and above the wings tend to be stretched to take a normal phenotype. This is an example of the fact that the phenotypic expression of the gene might be governed by the metabolic rate. With this in mind and encouraged by the work of Buzzati-Traverso, we attempted, first, to distinguish between the flies homozygous for vestigial and those heterozygous for the same gene, and between these and normal flies (OregonR+), with respect to their amino acid constituents and then tried to correlate the differences with the assumed phenotypic change caused by the temperature.

The temperatures selected for this experiment were as follows: 18°C, 22°C ± 1°C (as room temperature), 30°C, and 33°C. Unfortunately, the flies rearing at 33°C and above either became sterile or did not give rise to any living adult. However, they were reared at 33°C to be used for chromatographic purpose. As regards the methods for resolution of the difference between the various types of flies, we adopted the ascending chromatographic techniques, as given by Buzzati-Traverso, with the following modifications. We used Whatman filter paper No. 1 and No. 42. Six decapitated flies, at a time, were washed in 95% ethyl alcohol, boiled in distilled water for a minute or two, and then squashed at a spot on the paper. Males and females were separately squashed and recorded. The chromatogram was developed in a mixture of n-propanol and 1% ammonia in 2:1 ratio for a period of 18 to 24 hours. The \( R_f \) values are being presented in the table and discussed later. The flies obtained from the vestigial stock were, however, not tested for their isogenicity; but, rather, they were collected every time from the F2 with wild type inbred in the laboratory for a long period of time. This leaves an open question of other genes affecting the chromatographic pattern. However, an over-all picture can be obtained considering the relative qualitative and quantitative differences.

Results and conclusion:
The homozygous vestigial males and females differ from both heterozygous and normal flies. The difference is mainly based on the number and color of the spots and on their \( R_f \) values. The spots were of two types. In all the cases both fluorescent and ultra-violet-absorption spots were obtained. These results have been summarized in the table. There is a great difference in the \( R_f \) values of the fluorescent spots of males and females, within a given genotype.

Quantitative aspect:
There is a reduction in the \( R_f \) values for fluorescent spots obtained in the cases of flies raised at 33°C as compared to those raised at room temperature, except for \(+/-\) male and \(vg/+\) female, which show an increase. The significance of the increase in \(+/-\) male is, however, very poor. Distinction is possible between \(+/-, vg/vg\) and \(vg/+\) from their \( R_f \) values both at room temperature and at 33°C. It is interesting to note that \(vg/vg\) and \(+/-\) can be more easily distinguished from...
those heterozygous for vg, than between each other. There is a considerable change for both UV-absorption as well as fluorescence from room temperature to 33°C.

Qualitative aspect:

There is a great difference in the kinds of fluorescent spots in the different genotypic conditions of the flies used. These are presented in the table. One point needs to be mentioned here, that while at room temperature the qualitative differences are very pronounced, at 33°C these differences seem to be minimized. Another experiment, done with vgNo (vestigial-notched) mutant shows similar qualitative differences between the male and the female and also between this and other genotypic conditions (see the table).

Our results agree, in general, with those of Buzzati-Traverso. In our case we can further distinguish, by UV-absorption spots, the qualitative and quantitative nature of patterns in the different genotypes of the flies. The fluorescent spots might be either due to tyrosine or tryptophan or both. Both of these show fluorescence in conditions similar to those of the experiment.

In our experiment we could not get the variable phenotypic expression of the mutant vestigial in different temperatures (stretching of wings at high temperature). However, these preliminary results indicate that the study on the relationship between the temperature sensitivity of the gene and the chromatographic analysis of its products would probably be a very interesting topic for careful research.

---


Table: Summary of the results of the chromatographic behavior of vestigial and wild type flies of Drosophila melanogaster in different genotypic and sexual conditions. F = fluorescent spots, UVA = ultraviolet-absorption spots, Rm° C. = room temperature, ++ = color (bluish or yellowish in the bright spots) and separate, -- = no color but separate spots, -~ = no color, no separate spots but fused and continuous, ..... = not done, * = in separate expt.

<table>
<thead>
<tr>
<th>Conditions</th>
<th>Rf values at Rm° C. for</th>
<th>Rf values at 33° C. for</th>
<th>Number of spots</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>F</td>
<td>UVA</td>
<td>F</td>
</tr>
<tr>
<td>MALE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+/+(Oregon R+)</td>
<td>0.447</td>
<td>0.259</td>
<td>0.47</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>vg/vg</td>
<td>0.427</td>
<td>0.269</td>
<td></td>
</tr>
<tr>
<td>vg/+</td>
<td>0.5</td>
<td>0.3</td>
<td>0.456</td>
</tr>
<tr>
<td>vgNo/vgNo</td>
<td>0.447</td>
<td>0.194</td>
<td></td>
</tr>
<tr>
<td>FEMALE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+/+(Oregon R+)</td>
<td>0.638</td>
<td>0.212</td>
<td>0.33</td>
</tr>
<tr>
<td>vg/vg</td>
<td>0.627</td>
<td>0.221</td>
<td>0.356</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>vg/+</td>
<td>0.241</td>
<td>0.308</td>
<td>0.493</td>
</tr>
<tr>
<td>vgNo/vgNo</td>
<td>0.52</td>
<td>0.163</td>
<td></td>
</tr>
</tbody>
</table>
Munz, P. Xanthindehydrogenase activity in D. melanogaster (Oregon-R).

The enzyme activity in homogenates from different stages and sexes was determined by measuring the amount of newly-formed isoxanthopterin after 90 minutes incubation with 2-amino-4-hydroxypteridine as substrate. This reaction product was found in the following amounts (relative units of fluorescence): Pupae with faintly yellowish eyes $\phi = 11$; $\sigma = 12$. Imagos immediately after eclosion $\phi = 13$; $\sigma = 8$. Imagos three days old $\phi = 49$; $\sigma = 28$. These figures are relative to fresh weight. It is remarkable that females exhibit a higher enzyme activity than males though males contain much more isoxanthopterin than female imagos.

Narain, P. Effect of age of female on the rate of egg production in D. melanogaster.

(India) were studied for their daily egg production for life-time (about 40-50 days) starting from the first day of egg laying. The flies were raised on a standard medium evolved by Burdick (1954) which was also used for testing the females' level of egg production. The eggs were collected on food (coloured green) placed on card-board chips treated with paraffin. A little live yeast was put on the food-chip which was inserted in test tubes of size 6" x 0.7" where a female and her partner were kept. The food-chips were changed daily (usually after 24 hours) and the number of eggs laid therein were counted on a stereoscopic binocular microscope with 10 x magnification. The flies were kept at 25°C ± 1°C.

It was observed that the daily egg production showed rapid increase in the first few days reaching the peak on the 4th day of egg laying. Thereafter the daily egg production gradually decreased till 40th day of egg laying. The declining pattern of egg production after the attainment of peak was found to follow the following exponential:

$Y = 66.56 e^{-0.0269 t}$

where $Y$ represents the number of eggs corresponding to a particular day ($t$) of egg laying. It is apparent from the equation that after attaining the peak, the daily egg production fell, on an average, at the rate of 2.7% per day.

The results of comparing the mean egg production over life-time (average 52.5 eggs/day) with mean egg production taken over various 3- and 10-day periods for a set of 37 females are shown in the table below:

<table>
<thead>
<tr>
<th>Days of egg laying</th>
<th>Average egg production per day</th>
<th>Rank correlation coefficient between the two means</th>
</tr>
</thead>
<tbody>
<tr>
<td>2nd to 4th</td>
<td>68.1</td>
<td>0.54**</td>
</tr>
<tr>
<td>3rd to 5th</td>
<td>74.0</td>
<td>0.67**</td>
</tr>
<tr>
<td>4th to 6th</td>
<td>71.5</td>
<td>0.69**</td>
</tr>
<tr>
<td>5th to 7th</td>
<td>70.0</td>
<td>0.75**</td>
</tr>
<tr>
<td>6th to 8th</td>
<td>66.0</td>
<td>0.66**</td>
</tr>
<tr>
<td>7th to 9th</td>
<td>64.3</td>
<td>0.76**</td>
</tr>
<tr>
<td>8th to 10th</td>
<td>63.6</td>
<td>0.72**</td>
</tr>
<tr>
<td>1st to 10th</td>
<td>64.0</td>
<td>0.79**</td>
</tr>
</tbody>
</table>

**Significant at 1% level
Each of the rank correlation coefficient was found to be significantly different from zero. When tested for homogeneity, the various rank correlation coefficients were found to differ insignificantly among themselves. An estimate of the weighted values of rank correlation coefficients based on Fisher's z-transformation was found to be 0.71.

Thus, these findings revealed that the age of females has a significant bearing on their rate of egg production. There is a peak (4th day of egg laying) in their life-time when the females lay eggs with maximum rate. It is also evident that to take advantage of the Drosophila flies in breeding at short intervals of 8-10 days and the special feature of 3-day egg laying periods, the females should be tested for egg production when they are 6 to 8 days of age which is equivalent to 4th to 6th day of egg laying. Females of this age group will also be having sufficient vigour for producing their progenies. These findings varied but little with those reported by Gowen and Johnson (1946) and adopted by Bell et al. (1955), Rasmuson (1956), and Robertson (1957) in regard to the best period for testing females for their egg production levels.

Similar results have recently been obtained by Prabhu and Bhat (unpublished 1961) in D. ananassae also.


Narain, P., C. Joshi, and S. S. Prabhu. Response to selection for fecundity in D. melanogaster. In a quantitative character like fecundity, which is closely connected with fitness and which is less heritable, heritability being estimated at 5 to 15% by Bell et al. (1955) and at 18% by Robertson (1957), the response to a selection pressure is expected little and still less if the selection is applied to a laboratory population of Drosophila flies which has been under mass culture for 20 to 25 generations. The selection would also cease to be effective after few generations, the number of generations depending upon the level of heritable variation present in the initial population to which the selection is applied. In such a case to maintain the degree of heterozygosity and hence to keep the selection effective, a slight modification in the 'mass' selection method may be helpful. This was tried in the present investigation.

Selection was practised in two laboratory populations, initially derived from different localities in India, viz., Nai Basti and Matunga. Five cultures were set up each generation and selection was made in either directions with total of 100 females each generation, 4 females being selected out of 20 females in each culture. A control was also run simultaneously. While making pair matings from each culture, the mates of the females were taken in a random fashion from the other remaining cultures instead of the same. Further, after selection, the eggs
laid by the selected females were mixed and distributed at random in almost equal numbers in the cultures set up for the next generation. This method ensured minimum chances of inbreeding in the population, though inbreeding, as such, cannot be completely dispensed with in a closed population. The females were tested for their egg production level on 4th to 6th days of egg laying and the method of testing and other details were as reported by Narain (unpublished, 1961).

The results of selection experiments conducted for 10 and 7 generations respectively in Nai Basti and Matunga stocks of D. melanogaster expressed in terms of average egg production per day per female in each generation are shown in the table below. The level of average egg production in the initial foundation stocks of Nai Basti and Matunga were $85.3 \pm 1.50$ and $37.5 \pm 1.36$ eggs per day respectively. The standard errors are based on 'within' culture variations.

Table 1

<table>
<thead>
<tr>
<th>Generation No.</th>
<th>Nai Basti</th>
<th>Matunga</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>High</td>
<td>Control</td>
</tr>
<tr>
<td>1</td>
<td>71.8±1.35</td>
<td>64.9±2.06</td>
</tr>
<tr>
<td>2</td>
<td>82.7±1.50</td>
<td>77.4±1.69</td>
</tr>
<tr>
<td>3</td>
<td>76.1±0.88</td>
<td>69.6±1.42</td>
</tr>
<tr>
<td>4</td>
<td>67.6±0.80</td>
<td>65.9±1.71</td>
</tr>
<tr>
<td>5</td>
<td>88.9±1.22</td>
<td>80.5±2.12</td>
</tr>
<tr>
<td>6</td>
<td>79.3±0.94</td>
<td>70.9±2.61</td>
</tr>
<tr>
<td>7</td>
<td>67.2±0.97</td>
<td>65.6±0.96</td>
</tr>
<tr>
<td>8</td>
<td>40.3±0.68</td>
<td>34.9±0.72</td>
</tr>
<tr>
<td>9</td>
<td>46.8±1.13</td>
<td>38.1±1.58</td>
</tr>
<tr>
<td>10</td>
<td>52.6±1.40</td>
<td>51.6±1.43</td>
</tr>
</tbody>
</table>

The data in Table 1 indicate that there are wide fluctuations from generation to generation even in the control series where no selection was practised. The difference in "high" and "low" lines in Nai Basti stock, is quite high in second generation being $12.6 \pm 2.55$ whereas it falls down to quite a low figure of $4.3 \pm 1.38$ in fourth generation. Thereafter it fluctuates and does not go below $8.5 \pm 1.92$ of 10th generation. In Matunga stock, the situation is different, the maximum divergence being $22.1 \pm 3.99$ in the 5th generation whereas minimum being $3.4 \pm 2.95$ in the 7th generation. To get an idea of how the response to selection behaved as a fraction of selective force applied (which was measured by selection differential), the ratio of divergence (between high and low) with the cumulative selection differential which is frequently referred to as realized heritability (Falconer, 1960) was worked out after each round of selection. The results after each round of selection are shown in Table 2.
Table 2

Realized heritability for egg production after each round of selection

<table>
<thead>
<tr>
<th>Round of selection</th>
<th>Nai Basti</th>
<th>Matunga</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cumulative</td>
<td>Divergence</td>
</tr>
<tr>
<td></td>
<td>differential</td>
<td>h^2</td>
</tr>
<tr>
<td>One</td>
<td>36.7</td>
<td>10.7</td>
</tr>
<tr>
<td>Two</td>
<td>56.5</td>
<td>12.6</td>
</tr>
<tr>
<td>Three</td>
<td>59.9</td>
<td>10.2</td>
</tr>
<tr>
<td>Four</td>
<td>110.2</td>
<td>4.3</td>
</tr>
<tr>
<td>Five</td>
<td>129.4</td>
<td>8.6</td>
</tr>
<tr>
<td>Six</td>
<td>157.3</td>
<td>9.7</td>
</tr>
<tr>
<td>Seven</td>
<td>180.8</td>
<td>8.7</td>
</tr>
<tr>
<td>Eight</td>
<td>202.2</td>
<td>9.9</td>
</tr>
<tr>
<td>Nine</td>
<td>216.5</td>
<td>10.5</td>
</tr>
<tr>
<td>Ten</td>
<td>238.3</td>
<td>8.5</td>
</tr>
</tbody>
</table>

It is apparent from the above results that the response to selection was substantial in the initial few generations as judged by the values of realized heritability. After four rounds of selection in Nai Basti stock and after six rounds of selection in Matunga the response ceased to be of appreciable magnitude.

These results indicate that in a character like fecundity which is largely determined by environment, the effect of the modified 'mass' selection is to deviate the 'high' and 'low' lines initially to a significant extent and then preserve the heterozygosity at a level characteristic of the character, i.e., exhibiting only 5 to 7% of variation as genetic. Such a situation is likely to arise in characters largely determined by non-additive actions of genes, viz., dominance and epistasis.

References:

Narise, T. Genetic studies on migrating activity in D. melanogaster. A number of inbred strains which differed from each other with regard to random- and mass-migrating activities were used for the present genetic study (for details of material and methods refer to DIS 30, p. 149, 32, p. 153, and 34, p. 94). The findings obtained in the present experiment are: 1) Selection for high migrating activity was quite effective and the selected lines were found to be quite active in either mass-migration or random-migration. 2) Both the migrating activities were genetic characters and showed a dominance effect in some cases but not others. 3) By the method of substitution of II and III chromosomes, both migrating activities were found to be highly controlled by genes included in these chromosomes.
Nash, D. Selection for changes in the manifestation of the Hairless mutant.

Using the Hairless mutant of D. melanogaster balanced against LVM, the distribution of bristle absence and bristle vestige ("Socket") presence between the fourteen dorsal head macrochaetae has been studied. Selection has altered the susceptibility to loss of all macrochaetae except the median orbitals. Directional selection using four sub-culture, rotationally mated, populations has increased the mean numbers of bristles lost from four to ten, and reduced it to two. Breeding from the flies possessing sockets at all sites at which bristles are lost, and selecting for increased loss amongst this class of flies, has resulted in a 95% level of socket presence (base stock 65%) and a mean loss per fly of 7.5 bristles; the loss is concentrated at the eight sites where sockets were usually or commonly present in the base population.

Selection for the presence of one geographical group of bristles, the anterior and posterior orbitals, simultaneously with selection for absence of a second group, the anterior and posterior verticals, and for the converse, has met with limited success. It is considered that the limitation is a function of the developmental system; the verticals are developmentally associated and hence can react similarly to selection, but the posterior orbital is more closely associated with the verticals than with the anterior orbital.

The series of associations deduced from reactions to selection coincides with the series of bristles placed in order of percentage socket presence at sites where bristles are missing. Since there is no simple spacial relationship between the associated bristles in either of these series, and since the difference between presence and absence of sockets is probably a function of the time during bristle development at which the mutant is effective, it is suggested that there exists a developmentally significant temporal inter-relationship between bristles.

The effect of selection upon the manifestation of Hairless

<table>
<thead>
<tr>
<th>Stock</th>
<th>Base stock</th>
<th>Increased loss</th>
<th>Decreased loss</th>
<th>Orbital loss</th>
<th>Vertical loss</th>
<th>High socket presence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bristle</td>
<td>(gen 0)</td>
<td>(gen 19)</td>
<td>(gen 19)</td>
<td>(gen 19)</td>
<td>(gen 19)</td>
<td>(gen 19)</td>
</tr>
<tr>
<td>ant. vertical</td>
<td>(a) (b)</td>
<td>(a) (b)</td>
<td>(a) (b)</td>
<td>(a) (b)</td>
<td>(a) (b)</td>
<td>(a) (b)</td>
</tr>
<tr>
<td>post. vertical</td>
<td>25 98</td>
<td>83 100</td>
<td>1 100</td>
<td>14 100</td>
<td>72 100</td>
<td>95 100</td>
</tr>
<tr>
<td>post. orbital</td>
<td>21 98</td>
<td>84 99</td>
<td>3 100</td>
<td>28 85</td>
<td>48 98</td>
<td>70 100</td>
</tr>
<tr>
<td>post-vertical</td>
<td>39 88</td>
<td>97 49</td>
<td>4 92</td>
<td>52 64</td>
<td>40 97</td>
<td>86 100</td>
</tr>
<tr>
<td>ant. vertical</td>
<td>99 61</td>
<td>100 35</td>
<td>91 96</td>
<td>100 27</td>
<td>100 54</td>
<td>100 89</td>
</tr>
<tr>
<td>ocellar</td>
<td>14 3</td>
<td>92 5</td>
<td>2 0</td>
<td>80 1</td>
<td>16 12</td>
<td>19 85</td>
</tr>
<tr>
<td>(med. orbital)</td>
<td>.9 1</td>
<td>.3 100</td>
<td>0 -</td>
<td>0 -</td>
<td>1 50</td>
<td>1 33</td>
</tr>
</tbody>
</table>

Column (a) shows the percentage loss of bristles at a particular site; (b) shows the percentage of the sites at which the bristle vestiges are present.

Each percentage based upon data collected from 80 flies of each sex; i.e., 320 bristles.

For a description of the selection criteria, see text.
The definitive description of miniature-dominant \( (m^D) \) by Slatis (Genetics, 1954) revealed that viability was about 20% to 50% in hemizygotes and 5% in homozygotes and that there was very low fertility in homozygous females. The present note includes results of further investigations involving this mutant. The foundation stocks for these experiments were developed from stock No. 65 from Pasadena (DIS-34). The stock is maintained by breeding \( m^D/FM3 \) females to \( m^D \) males. In matings of \( m^D/m^D \) by \( m^D \) flies from the stock cultures approximately 35% of single-pair matings were fertile. Mass selection from the fertile matings has yielded stocks in which the per cent fertile matings is between 60% and 70%. It is possible to get direct comparisons of the relative viabilities of the \( m^D \) homozygotes compared to the hemizygotes as these are the only two classes of progeny segregating in these matings. A total of 3456 progeny were counted, comprising 817 females and 2639 males indicating \( m^D/m^D \) females are only 31% as viable as \( m^D \) males.

A preliminary study indicates that fertility is not impaired in \( m^D/+ \) females.

The occurrence of riboflavine and sepiapteridine in the heads, testis and Malpighian tubules of Drosophila melanogaster (wild type "Sevelen" and homozygous sepia-mutant) has been qualitatively investigated. By purifying the extracts with several solvents in column-chromatograms small quantities of practically pure riboflavine and sepiapteridine were obtained. The results are given in the table in a semiquantitative form.

<table>
<thead>
<tr>
<th>genotype</th>
<th>organs</th>
<th>riboflavine</th>
<th>sepiapteridine</th>
</tr>
</thead>
<tbody>
<tr>
<td>wild type</td>
<td>heads</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>testis</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>Malpigh.</td>
<td>++++</td>
<td>+</td>
</tr>
<tr>
<td>sepia-mutant</td>
<td>heads</td>
<td>-</td>
<td>+++++</td>
</tr>
<tr>
<td></td>
<td>testis</td>
<td>-</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>Malpigh.</td>
<td>++++</td>
<td>+</td>
</tr>
</tbody>
</table>

Sepiapteridine was found in all of the tested organs of the two genotypes, while riboflavine was not detectable in the heads and testis of the sepia-mutant. Therefore it appears that the sepia-locus not only affects the pteridines but also the flavines (although probably only quantitatively), since riboflavine was easily found in the heads and testis of the wild type.

In both genotypes the Malphigian tubules contain much riboflavine. The UV-spectrum of this substance was found to be identical with that of synthetic riboflavine.

Terzaghi and Knapp (Evolution, 1960) have shown that if D. pseudoobscura is heterozygous for inversions in only one pair of chromosomes, there is only a very small amount of zygote mortality but that if there are heterozygous inversions in two or more independent chromosomes, mortality is high, running up to fifteen per cent. This is similar to the result reported earlier for D. melanogaster (Cooper, Krivshenko and Zimmering) but is of particular interest for this species because it provides a simple explanation for the accumulation of inversions in the third...
chromosome only of pseudoobscura: if a certain chromosome, as the third, by chance becomes variable in sequence, there will be no great selective disadvantage (indeed, it may provide an advantage through heterosis), but if subsequently any other chromosome (as the second) suffered an inversion, the new sequence would immediately be at an extreme disadvantage because of the great zygote mortality elicited when this new sequence found itself in a heterozygote in combination with an inversion heterozygote in the other chromosome and, similar to the argument applicable to the gradual decrease of the least frequent allele when a heterozygote has reduced adaptive value, inversions in any chromosomes other than the one with the fortuitous headstart will be eliminated.

While this is an attractive scheme to account for the situation in D. pseudoobscura, and a few other species in which variability is almost completely limited to one chromosome, it does not appear to hold for other species (D. robusta, for instance) where inversions appear to be distributed generally throughout the complement. It is then a natural, and important, question as to whether this difference has its basis in a different behavior of the chromosomes in the two types, or whether other (possibly non-genetic) factors (such as an initial geographic isolation of inversions in non-homologous chromosomes) played some role.

It is possible of course to answer this question for each species by a direct experimental attack; however, the writer would like to make a simple observation that suggests strongly that the chromosomes in all reasonably well studied species of Drosophila will behave the same and that the answer to this question must rest in the second alternative.

The zygote mortality described above is undoubtedly caused by non-homologous pairing, followed by irregular disjunction of non-homologous chromosomes. As has been emphasized recently by Sandler and Novitski (Genetics, 1956), such pairing manifests itself quite obviously in 3N meioses, where the frequency of balanced gametes is considerably lower than one would expect if the distribution of the single member of each trivalent towards a given pole were simply \((1/2)^N\) as elementary texts tell us it should be. If the mortality is high in a species like melanogaster with three major autosomes, then it should be very much greater (certainly more than doubled) for a species with four major chromosomes because of the greater possibilities for non-homologous pairing in the first place, and the reduced fraction of balanced gametes among all possible combinations in the second. It is now clear why the writer was unsuccessful in producing a permanent 3N stock of D. pseudoobscura (with four major chromosomes) although the initial 3N female did produce one 3N offspring (DIS 20). Similarly, it undoubtedly is not fortuitous that 3N lines of simulans can be maintained without difficulty, but that a 3N line in D. virilis is unknown despite the great amount of work done on this form, especially in Japan, by the most acute observers. The one apparent exception to this trend (to my knowledge) is the case of polyploidy reported by Stalker (Genetics, 1942), but this strengthens rather than weakens the argument, since it occurred in americana which has two pairs of its elements joined, thereby shifting the form from the five chromosome group to the three chromosome group like melanogaster.

The conclusion, then, is that the absence of fertile triploid stocks in species other than those with three major autosomes suggests that in most if not all species of Drosophila we would expect to find pairing potentialities conducive to the accumulation of inversions in some one chromosome (as in the third of pseudoobscura) at the expense of the others.

Novitski, E. A note on Sturtevant and Beadle's 1936 inversion paper. For years I have puzzled over the explanation given on pp. 584-586, Genetics, 1936, concerning the behavior of tandem metacentric chromosomes. Questions directed to the two authors have been of no help, because of the long time that has elapsed since that paper was written, and the loss of the notes on which that section was based. Others also have been mystified by their calculations, so I should like to report partial success, at least, in unravelling their puzzle.
The question is: how is the figure of 90.8%, given as the best estimate of the frequency of single crossing over in their tandem metacentric, arrived at? We know now, of course, that their experiment was confounded by non-random disjunction, which was unrecognized at the time. For this reason, the number of recovered crossovers was excessive, in fact, was greater than the number of non-crossovers and by their method of calculation should have amounted to 114% single exchanges (or four times the number of recovered rings, 313, divided by the number of patroclinous males, 1098). A second method of computation for the singles is to take the excess of males over females, 445, multiplied by four because each lost egg comes presumably from that one fourth of the exchanges producing dicentrics, the product then to be divided by twice the number of patroclinous males, 1098. This gives a value of 81.06%. (At this juncture an arithmetical slip seems to have been made, because to get the right answer we must use the value of 81.6%.)

Now we are confronted with the nasty problem of having two distinctly different estimates from the same data, 114% and 81.6%. This is resolved by, first, reducing the 114% to 100% since the first figure is clearly impossible, and then averaging 81.6% and 100% to get 90.8%!

It might also be mentioned that Table 15 on which the logic of the arithmetic is based is in error by having several figures reversed (undoubtedly a typesetter's error) and that for a better approach to this problem of estimating exchange in tandem metacentrics, one should refer to the author's paper on non-random disjunction (Genetics, 1951).

Ogaki, M. Inheritance of heat tolerance in D. melanogaster.

Two strains of Hikone-H and Mino-H have been reared at higher temperature than 30°C through more than three years. These stocks are able to breed at 31°C successively, but heat susceptible stocks, for instance B;e11, se ss and others, reproduce no fly at the same temperature in the next generation. Genetical analysis of heat tolerance indicated that this character was polygenic, but the major gene was recognized as dominant, and located around the spindle fiber attachment on the third chromosome. It is notable that this locus is almost the same place as that of the nicotine sulfate resistant gene in D. melanogaster. It is also suggested that the selection pressure to nicotine sulfate increases the heat resistability without contact to heat. This seems to imply that the main gene of heat tolerance is very closely related to the nicotine resistant gene. Otherwise it is assumed that the heat tolerance might be a kind of vigor tolerance manifested by the same gene as that of the nicotine resistance.


Methylbutyrate was scarcely hydrolyzed by homogenates of two mutant strains (bw;st ss and bw;st;svn), while homogenates of wild type strains have high methylbutyrate-splitting capacity. From the results of genetical analyses, it became clear that the low ali-esterase activity was controlled by a recessive factor on the 3rd chromosome. Although it is still premature to conclude that the low activity is controlled by only one gene, it may be called ali for convenience sake. It is very interesting that the activity of the hybrid (ali/ali+) obtained from the crosses of low activity flies (ali/ali) and high activity flies (ali+/ali+), reveals an intermediate level of the parent strains.

The cholinesterase activity, however, showed no difference among these strains. In fact, the distribution of ali-esterase in parts of the body was distinctly different from that of cholinesterase. The results suggest that these esterase activities are controlled by different genes.
Okada, T. "Speed index" shown by the apodemes of drosophilid flies. There is known to occur a phenomenon among systematic groups of Drosophilidae that the length of the apodeme of aedeagus shows gradual decrease in accordance with the gradual development of the aedeagus itself. In other words, the phallosomal index (ratio in length of the aedeagus and its apodeme) tends to increase in the more advanced forms. A similar pattern of decrease is seen in the ratio in length of the basal plate and the stalk of the ejaculatory apodeme. The increase of these ratios should be of high biological significance, since it turns out to bring the more speedy contraction of muscles attached to the apodemes by means of shortening distance of muscular action. Therefore, this phenomenon can be interpreted by that of "speed index" proposed by Lull for the ratio in length of the metacarpus III and the femur, which becomes higher in the more speedy mammals as shown by N. American fossil horses.

Okada, T. "Compensatory adaptation" of the ejaculatory apodeme of drosophilid flies. Although the general process of differentiation of the ejaculatory apodeme of the drosophilid flies seems to be the gradual shortening of the stalk in accordance with the development of the plate, as expressible by the phenomenon of "speed index" (see above), in some forms, e.g., the members of the subgenera Sophophora and Lordiphosa, the stalk remains elongate and the effectiveness of muscle contraction seems to be compensatorily attained by the centripetal shifting of the junction of stalk on the plate. This kind of structural differentiation concerning a certain functional adaptation may be called "compensatory adaptation."

Okasa, T. A. The effect of autosomal inversion heterozygosity on crossing-over frequency in the X chromosome of D. melanogaster. It is well known that inversion heterozygosity produces an increase in crossing-over frequencies in nonhomologous chromosomes. This increase has been found to be most striking around the centromere and at the tips of the chromosome arms, being less pronounced in the central regions of the arms. When the X chromosome has been the affected chromosome and the relative increases in crossing-over frequencies have been determined in different regions along its whole length an essentially U-shaped curve has been obtained (e.g. Schultz and Redfield, Cold Spring Harbor Symp. Quant. Biol. 16, p. 184, fig. 5, 1951). However, in papers dealing with this phenomenon the long central part of the X has not been closely marked. Therefore, the present author has carried out a more detailed analysis concerning the region from crossveinless to forked. This region was divided into five subregions (see below) and each of them was tested separately in four parallel experiments: the standard autosomal homozygote as a control, the In(2L+2R)Cy heterozygote, the In(3L+3R)P heterozygote, and the combination of these two autosomal inversion heterozygotes. The following relative increases (in per cent) from the control were found:

<table>
<thead>
<tr>
<th>Region</th>
<th>Curly</th>
<th>Payne</th>
<th>Curly: Payne</th>
</tr>
</thead>
<tbody>
<tr>
<td>cv - sn</td>
<td>18.3</td>
<td>19.5</td>
<td>56.6</td>
</tr>
<tr>
<td>sn - lz</td>
<td>3.1</td>
<td>4.8</td>
<td>33.5</td>
</tr>
<tr>
<td>lz - m</td>
<td>37.7</td>
<td>44.7</td>
<td>87.1</td>
</tr>
<tr>
<td>m - g</td>
<td>9.8</td>
<td>12.1</td>
<td>55.2</td>
</tr>
<tr>
<td>g - f</td>
<td>14.8</td>
<td>37.4</td>
<td>50.4</td>
</tr>
</tbody>
</table>
The curve computed from these data together with the data for the ends published by Schultz and Redfield is not U- but rather W-shaped with a fairly conspicuous peak in the very middle of the one armed chromosome (more or less around vermilion). This result is very much what one should expect on the basis of the tetrad analysis carried out by Redfield (Genetics 42, p. 723, Table 4, 1957). This analysis showed that in the presence of autosomal inversions the singles in the X tend to be replaced by triples (and to smaller extent by quadruples). When there are three crossovers in the same tetrad it is but natural that they are, due to interference, situated as far from each other as possible at even intervals, i.e. at both ends and in the middle. This state of affairs is reflected in the three-peaked curve obtained.

Further interpretation of the above findings is possible on the basis of the hypothesis put forward by the present author in an earlier connection (Oksala, Cold Spring Harbor Symp. Quant. Biol. 23, pp. 197-210, 1958).

Oshima, C. The persistence of deleterious genes in natural populations of D. melanogaster. The second chromosomes were isolated from several Japanese wild populations by using the method of completely marked inversion. The relative frequencies of chromosomes carrying lethal, semi-lethal, subvital and normal genes were estimated. The results obtained in 1959 had been reported in DIS 34 (p. 99). A similar sampling of second chromosomes from the same populations was carried out also in 1960 and the different classes of deleterious chromosomes were similar in relative frequencies. There was apparently no fluctuation between samples collected in 1959 and 1960. The lethal chromosomes have been maintained in the Cy balanced system in the successive generations. Diadial crosses were performed with all lethal strains to determine the allelic rate within and between populations. After maintaining the lethal chromosomes during the year 1959, they were subjected to cross-testing with new lethal chromosomes isolated in 1960 from the same populations. The allelic rates underwent scarcely any change during one year, but they seemed to have increased slightly in 1960. Most interesting was that the two lethals isolated in 1959 were found again in 1960 (allelic rate: 0.87 per cent). This finding shows that the two lethals have been maintained in the same population at least for a year.

Oshima, C. Dieldrin resistance in D. pseudoobscura. About ten strains of each five kind of chromosomes for homozygous ST, AR, CH, TL and PP, which had been established by Prof. Th. Dobzhansky from flies collected in California in 1959, were transferred to us in 1960. Several strains having the same chromosomal arrangement were mated, and female and male flies in the offspring were tested separately with test papers, containing 0.8, 0.4, 0.2, 0.1 and 0.05 per cent Dieldrin. Ten flies were exposed to test paper for one hour in a small vial and then transferred into another vial containing wet filter paper. After 24 hours, the number of dead flies was scored. Such a test was repeated ten times. On the other hand, two different chromosomal strains were crossed and the hybrid offspring were tested by the same method described above. The mortalities obtained in the experiments were transformed into arc-sine units and analyzed statistically. From the results of analysis of variance, the difference between chromosomal strains was highly significant and the order in resistance was observed as follows: TL > PP > CH > ST > AR. Flies having the relatively rare chromosomes such as TL and PP in California would be more resistant than flies having common chromosomes. The significant differences between resistances of both sexes and mortalities in doses were observed expectedly, but the difference between resistances of monochromosomal strain and hybrid strain could not be detected significantly.
Parsons, P. A. A biochemical polymorphism in Drosophila melanogaster. Ebony (e"e") flies have less tyrosinase than wild type flies. Flies can be rapidly tested for tyrosinase content by growing larvae on the tyrosinase inhibitor phenyl-thio-carbamide (P. T. C.), and ascertaining the lethal concentration of P. T. C. Recently, in an Oregon-R stock of flies an allele at the ebony locus, reacting to P. T. C. in a similar way to e"e" files, has been found. This "ebony" allele is wild type in body colour. The Oregon-R stock is polymorphic for this allele with a gene frequency of 13%. Flies collected in the wild in September, 1961, at Eugene, Oregon, have also turned out to be polymorphic. Hence the allele in the Oregon stock probably came from the wild population from which the stock was derived. Other polymorphic stocks found so far are Oregon-K, Kaduna (from Africa) and Bikini with gene frequencies of 43%, 40%, and 67% respectively, while a Canton-S stock from the California Institute of Technology is 100% "ebony." Flies collected in the wild near Cambridge are also polymorphic. Hence this polymorphism is probably very widespread in the wild. It probably explains, in part, the variability of tyrosinase estimations found in different stocks.

The selective basis of the polymorphism is unknown, but two observations deserve mention, namely (1) the gene frequencies in the wild Cambridge flies has not varied much during the summer of 1961 and (2) there is a degree of male sterility of the "ebony" homozygotes in the Cambridge flies. It is, however, remarkable that the "ebony" allele has probably persisted in the Oregon Laboratory population for at least 35 years, since the Oregon-R stock was collected at Roseburg, Oregon, in 1925 or before by D. E. Lancefield (Bridges & Brehme, 1944).

Pelecanos, M. Induced oögonial lethals in Drosophila. A simple method for the induction of high frequencies of oögonial sex-linked recessive lethal mutations by larval feeding with diethylsulphate is reported.

Tests for the mutagenicity of chemicals are usually carried out on males. By the larval feeding method, only one chemical (chloroethyl methanesulphonate) has so far been shown to have a mutagenic effect on females (Auerbach and Sonbati, 1957). Larvae were cultured on an aseptic medium with the following composition:

<table>
<thead>
<tr>
<th>Substance</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>10%</td>
</tr>
<tr>
<td>D. C. L. Yeast</td>
<td>10%</td>
</tr>
<tr>
<td>Agar</td>
<td>3%</td>
</tr>
</tbody>
</table>

The medium was autoclaved and 0.4 per cent propionic acid was added at 65°C, and diethylsulphate at 60°C. The medium was dispensed as 25 ml portions into sterile bottles. Oregon-K eggs were sterilised using Sang's (1956) method, and one hundred newly hatched larvae were transferred to each culture. Hatching males and females were collected as virgins and tested for sex-linked recessive lethal mutations by the Muller-5 mating method. Four three-day broods were studied; each male was mated with two females and vice versa. Table 1 shows the results obtained over four broods for treated males and females. A high rate of sex-linked recessive lethals was maintained for both sexes over the four broods tested.

Results so far available do not allow comparisons between the effects of larval treatment in males and females, since both spermatogonia and primary spermatocytes are present during the male larval period, whereas only oögonia occur in the female larvae. However, the significant heterogeneity $X^2$ in females suggests that there might be stages of different sensitivity among the larval oögonia.

Table 1.
Concentration of diethylsulphate 0.5% (Molarity 3.2 x 10^-4). Treatment throughout the larval life. Temp. 25°C.

| Time of Development | 9 days |  
|---------------------|--|---
| % Survival         | % lethal | % Survival | % lethal |
| **d♂ Treated**     | **♀ Treated** |
| Broods | No. chrom. | No. lethal | % lethal | No. chrom. | No. lethal | % lethal |
| 1st    | 672       | 83        | 12.35    | 856       | 135        | 15.77    |
| 2nd    | 626       | 63        | 10.06    | 864       | 136        | 15.81    |
| 3rd    | 514       | 43        | 8.36     | 514       | 99         | 19.26    |
| 4th    | 422       | 52        | 12.32    | 672       | 143        | 21.28    |
| Total  | 2234      | 241       | 10.79    | 2906      | 513        | 17.65    |

Heterogeneity $X^2$ 3 D.f.
for $d♂ = 6.01 (0.20 > P > 0.10)$
for $♀ X^2 = 11.91 (0.001 > P > 0.01)$

Time of action of the lethal effect in the dumpy series.

Various alleles of the dumpy series manifest a lethality as homozygotes or as compounds derived from interallelic crosses. Several such lethal-bearing alleles were mated to Ore R wild type flies and their heterozygous $F_1$ progeny were used for examination of the time of action of the lethal effect. In most of these tests, the progeny were mated in vials and transferred to petri dishes containing a sugar and bactoagar medium treated with streptomycin. This system permitted a relatively germ free development on a transparent food medium.

In Table 1 the flies were examined for their egg hatchability after fertilization. These results suggest that the homozygous alleles $ol^6$, $lv^1$, and $olv^1$ all die in the embryonic stage (before the egg collapses with the emergence of a first instar larva). The mutant $l^m$, as a homozygote, survives this stage and dies at a later stage (the third larval instar). In heterallelic crosses, a partial complementation for the lethality exists for most of the crosses. Thus $l^m/ol^6$, $ol^6/olv^1$, and $l^m/olv^1$ show little mortality in the embryo. In $lv^1/ol^6$, and $l^m/lv^1$ there is less lethality in the embryo than for homozygous $lv^1$. Only $lv^1/olv^1$ maintains a high mortality in the embryonic stage.

In Table 2, the mortality is shown to exist prior to the pupal stage, based on the survivors from pupae to adults in these crosses. This is important because both the oblique wing ($o$) and the thoracic vortex ($v$) effects of the dumpy series are manifested at the pupal stage at slightly different times (see Carlson and Falk, this issue). In Table 3, the lethality not manifested at the embryonic stage is shown to exist primarily during the third instar larval stage.

The mechanism for the lethality may differ, not only in those instances where their time of action differs, but in those cases where the same apparent stage is affected. Thus uncollapsed eggs of $ol^6/ol^6$ and $olv^1/olv^1$ genotypes are yellow, but eggs of $lv^1/lv^1$ genotype are white. The color of uncollapsed $l^m/l^m$ embryos (which account for only a small portion of the total lethality of this compound) is also white. No attempt was made in this study to examine first and
second instar larval mortality, but they must be slight, judging by the results of Tables 1 and 3. Because of the small numbers used in each test, no attempt was made to subtract the control values from the experimental cultures. This provides a maximal lethality which is, of course, higher than the corrected values which should have the frequency of the control lethality subtracted.

Table 1

<table>
<thead>
<tr>
<th>Cross</th>
<th>Total eggs laid</th>
<th>Uncollapsed eggs</th>
<th>Maximal % lethality</th>
</tr>
</thead>
<tbody>
<tr>
<td>im/+ x im/+</td>
<td>274</td>
<td>9</td>
<td>3.3</td>
</tr>
<tr>
<td>olS/+ x olS/+</td>
<td>133</td>
<td>31</td>
<td>23.3</td>
</tr>
<tr>
<td>lv1/+ x lv1/+</td>
<td>130</td>
<td>28</td>
<td>21.5</td>
</tr>
<tr>
<td>olv1/+ x olv1/+</td>
<td>177</td>
<td>42</td>
<td>23.7</td>
</tr>
<tr>
<td>lv1/+ x olS/+</td>
<td>132</td>
<td>8</td>
<td>6.1</td>
</tr>
<tr>
<td>1m/+ x lv1/+</td>
<td>127</td>
<td>13</td>
<td>10.2</td>
</tr>
<tr>
<td>lv1/+ x olv1/+</td>
<td>149</td>
<td>23</td>
<td>15.4</td>
</tr>
<tr>
<td>1m/+ x olS/+</td>
<td>124</td>
<td>2</td>
<td>1.6</td>
</tr>
<tr>
<td>olS/+ x olv1/+</td>
<td>126</td>
<td>3</td>
<td>2.4</td>
</tr>
<tr>
<td>1m/+ x olv1/+</td>
<td>169</td>
<td>9</td>
<td>5.3</td>
</tr>
</tbody>
</table>

Table 2

<table>
<thead>
<tr>
<th>Cross</th>
<th>Eggs laid</th>
<th>Pupae</th>
<th>Adults</th>
</tr>
</thead>
<tbody>
<tr>
<td>1m/+ x olv1/+</td>
<td>183</td>
<td>124</td>
<td>122</td>
</tr>
<tr>
<td>olS/+ x olS/+</td>
<td>109</td>
<td>68</td>
<td>66</td>
</tr>
<tr>
<td>1m/+ x 1m/+</td>
<td>227</td>
<td>122</td>
<td>119</td>
</tr>
<tr>
<td>olv1/+ x olv1/+</td>
<td>192</td>
<td>137</td>
<td>120</td>
</tr>
<tr>
<td>lv1/+ x lv1/+</td>
<td>104</td>
<td>73</td>
<td>70</td>
</tr>
<tr>
<td>+/- x +/-</td>
<td>64</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>lv1/+ x olS/+</td>
<td>187</td>
<td>131</td>
<td>129</td>
</tr>
<tr>
<td>1m/+ x lv1/+</td>
<td>145</td>
<td>97</td>
<td>93</td>
</tr>
<tr>
<td>olS/+ x 1m/+</td>
<td>99</td>
<td>59</td>
<td>59</td>
</tr>
<tr>
<td>olv1/+ x lv1/+</td>
<td>129</td>
<td>99</td>
<td>93</td>
</tr>
<tr>
<td>olS/+ x olv1/+</td>
<td>100</td>
<td>58</td>
<td>56</td>
</tr>
</tbody>
</table>
Table 3  
Maximal frequencies of lethality at different developmental stages for dumpy alleles

<table>
<thead>
<tr>
<th>Cross</th>
<th>Eggs laid</th>
<th>Lethality stage A</th>
<th>Lethality stage B</th>
<th>Lethality stage C</th>
<th>Pupae</th>
<th>Lethality stage C</th>
<th>Adults</th>
<th>Maximal per cent lethality</th>
</tr>
</thead>
<tbody>
<tr>
<td>+/+ x +/+</td>
<td>55</td>
<td>9.1</td>
<td>50</td>
<td>3.64</td>
<td>48</td>
<td>0</td>
<td>48</td>
<td>12.7</td>
</tr>
<tr>
<td>olv¹/+ x olv¹/+</td>
<td>92</td>
<td>25.0</td>
<td>69</td>
<td>16.4</td>
<td>54</td>
<td>5.4</td>
<td>49</td>
<td>46.8</td>
</tr>
<tr>
<td>l⁻¹m/+ x l⁻¹m/+</td>
<td>105</td>
<td>4.8</td>
<td>100</td>
<td>23.8</td>
<td>75</td>
<td>0</td>
<td>75</td>
<td>28.6</td>
</tr>
<tr>
<td>l⁻¹m/+ x olv¹/+</td>
<td>122</td>
<td>4.1</td>
<td>117</td>
<td>38.7</td>
<td>82</td>
<td>0</td>
<td>82</td>
<td>32.8</td>
</tr>
<tr>
<td>ol⁺¹⁸/+ x olv¹/+</td>
<td>185</td>
<td>10.8</td>
<td>165</td>
<td>31.4</td>
<td>107</td>
<td>0.5</td>
<td>106</td>
<td>42.6</td>
</tr>
</tbody>
</table>

Stage A = lethals occurring from embryo to second instar larvae  
Stage B = lethals from third instar larvae to beginning of pupation  
Stage C = lethals during pupation  

(This work was supported by grant 14222 from The National Science Foundation.)
Several spontaneously occurring rare mosaic types have been observed in the progeny of triploid females in the course of experimental work. These verify the previously known facts of sex determination and fail to give indication of hormonal effects of tissues of diverse origins upon one another except in the last case. Certain mosaics demonstrate the simultaneous loss of X, II, and III. Descriptions of the mosaics follow: (1) Triploid-diploid female. This individual, coming from a cross of ru, T(3,4) 30/ca, T(3,4) 28 males with y²; ru ca triploids, had a gray body and displayed the mutants ru and ca on both sides. Both eyes possessed the large eye facets typical of 3A tissue. The left wing showed the coarse wing texture characteristic of 3A tissue and was slightly longer than the right wing which exhibited the fine texture of diploid wings. All primary and secondary sexual characters were female. It is presumed that the genetic composition of the left side was +/-y²/y²; II/II/II; ru, 30L+ca, 28R/ru ca/ru ca (hyper-triploid for the short section between T(3,4) 30 and T(3,4) 28.) The right side was 2X2A, having lost one y² X chromosome, one II, and either an intact III or both the 30L and the 28R fragments of III. (2) Triploid-diploid female. This individual arose from a cross of ca, T(3,4) 85C/ru, T(3,4) 89E males with y²; ru ca triploids. The mosaic possessed gray body color throughout, and both eyes were red (not claret) and not-roughoid, but they exhibited diploid facet size. Both wings, on the other hand, showed coarse 3A texture. All primary and secondary sexual characters were female. The fly presumably started out with +/-y²/y²; II/II/II; 85CL + 89ER/ru ca/ru ca (a hyper-triploid for the short section between T(3,4) 85C and T(3,4) 89E). Subsequent loss of one y² X, one II, and one intact ru ca III chromosome produced the diploid eyes. (3) Triploid-intersex. This individual was found in the homozygous y²; ru ca triploid stock and was therefore homozygous for these three mutants. Both eyes were the same size and shape and appeared like those of a triploid female. The right wing was distinctly shorter than the left but both wings showed coarse 3A texture. The right foreleg bore a sex comb of 8 prongs, the usual size of intersex sex combs in this stock. No sex comb was on the left foreleg. The abdomen was typically female, not bent or misshapen, and genitalia were female. Presumably the anterior portion of the right or intersex side arose from tissue which had lost 1X from the 3X3A intersex complement. (4) Triploid-diploid female-intersex triplet mosaic. This individual occurred in the progeny of T(3,4) 28/ru ca males with y²; ru ca triploid females. It is thought to be the result both or double fertilization and chromosome loss. The left wing was a mere stub. The head, forelegs, left half the thorax, left half plus the right distal half the abdomen were gray (not yellow²). The left eye was triploid (3A) in facet size and showed the mutants ru and ca. The top one fifth of the right eye was likewise ru ca and triploid in facet size. The lower four fifths of the right eye was not-ru, red (not-ca) and diploid in facet size. The right half the thorax, right anterior half the abdomen were yellow. The right wing was clearly of 3A texture. All primary and secondary sexual characters were female. Presumably the left gray, ru ca triploid part of the body arose from a union of sperm bearing a gray X and intact ru ca chromosome III and one chromosome II with a diploid egg possessing y²/y²; II,II; ru ca/ru ca. The yellow tissue including the 3A right wing, right half of thorax, and right anterior half of abdomen was apparently intersex, arising from the same zygote as the gray half except for loss of the gray X chromosome. The red, not-ru (not-roughoid) part of the right eye must have arisen by fertilization of a haploid second oöcyte (one y² X/one II/one intact ru ca III) by a sperm carrying a gray X, one II/ and the T(3,4) 28 chromosome fragments bearing the normal alleles of ru and ca. An alternate explanation of the non-roughoid, red (not-ca) lower 4/5 of the right eye is that this tissue is haploid, derived from the development of above sperm alone. (5) Intersex-hypointersex mosaic. This individual arose from a cross of ru, T(3,4) 12/ca, T(3,4) H3 males and y²; ru ca triploids. It was typically intersexual throughout. The eyes showed 3A texture on both sides, but the left eye was...
not-roughoid, claret; the right eye, roughoid and claret. The mosaicism of the eyes was unmistakable since the mutant ru narrows the eye and disturbs facets in the intersex eye considerably more than in a diploid ru eye. Body color was y² throughout. Wings were slightly outstretched and had coarse 3A texture. Genitalia were of the fragmentary male type; anal plates, of the female type. Sex combs of 8 and 9 prongs were present on the right and left forelegs respectively. The fly is presumed to be the result of union of a sperm bearing a Y chromosome/one II/ and the ca, T(3, 4) H3 chromosome III with a diploid egg containing y²/ y²; II/II; and ru ca III/ru ca III followed by the loss of the H3L fragment in the tissue giving rise at least to the right eye. This mosaic is interesting because the H3L fragment represents half of chromosome III, and a hypointersex lacking such a long fragment does not ordinarily live. Even if the right eye were derived from a different syngamy; i.e., from the union of sperm bearing a Y chromosome; one II; and ru, 12L + H3R with a diploid second oöcyte carrying y²/ y²; II/II; ru ca/ru ca, then hypointersex tissue would result. This hypointersex tissue would lack in triplicate one dose of the section of the III chromosome between T(3, 4) 12 and T(3, 4) H3 and would not be expected to survive except as mosaic tissue. In this mosaic alone we may observe evidence of a "hormone effect" since the right eye hypointersex tissue survived in the mosaic fly.

(This work was supported by Research Grant 3453, Public Health Service, Bethesda, Md.)

Pozzi, L. V., S. Giavelli, G. P. Sironi, and E. Gallucci. Frequency of recessive sex-linked lethals in D. melanogaster spermatogenesis, in O₂, N₂ and air, with 600 r and 1200 r. Using a mating system of one irradiated male to three Muller 5 females, with renewed matings every 24 hours, a sensitivity spectrum of different sperm cells stages is scored, which, in air, shows a peak corresponding to spermatids. Gas treatment is given before, during and after irradiation. N₂ treatment removes the sensitivity peak, that is different developing stages do not show significantly different mutation frequencies. Maximum O₂ effect is found in meiotic stages. The frequency pattern of the two doses in regard to gas atmosphere is quite constant.

Reitan, P. J., and M. E. Aman. The effects of dehydration on the frequency of irradiation induced embryonic abnormalities in Drosophila. It has been demonstrated by Herskowitz that the dehydration of Drosophila females prior to their exposure to irradiation increased the frequency of mortality of eggs laid during the first eight days following treatment and increased the number of cross-over like exchanges in the X chromosome and gross chromosomal rearrangements. It was suggested that the increases noted were probably due to induced lethal mutations. The work reported here has demonstrated an increase in the number of induced embryonic abnormalities in the eggs of females that had been desiccated prior to irradiation. Abnormalities noted include only those in which a substantial number of cells had been produced and in which the developmental sequence was interrupted prior to sixteen hours of development.

Virgin Drosophila melanogaster females were desiccated, desiccated and irradiated or irradiated only. Desiccation was carried out by exposure to 25% humidity for six hours. Irradiation consisted of exposure to 3,000 r or 4,000 r from a cobalt-60 source at 40 r per minute. Eggs were collected at eight hour intervals during the first five days following treatment. They were divided so that some were used for embryological examination and the remainder for the determination of hatchability. Eggs for embryological examination were allowed to develop for 16 hours, then fixed in Carnoy's and stained with iron haemaotoxylin.
The table shows the results from three experimental groups. Hatchability data was highly variable in all treatment groups; however, the number of gross developmental abnormalities noted among the eggs which had clearly undergone development was consistently greater in the embryos from desiccated and irradiated flies. Desiccation alone did not have any effect on the embryonic development. The data support the hypothesis that dehydration increases the number of irradiation induced lethal mutations.

<table>
<thead>
<tr>
<th>Experimental Group</th>
<th>No. of embryos</th>
<th>No. unfertilized eggs</th>
<th>No. grossly abnormal</th>
<th>% of developing embryos showing gross abnormalities</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ax 45</td>
<td>8</td>
<td>6</td>
<td>16.2</td>
<td></td>
</tr>
<tr>
<td>3,000 r bx 82</td>
<td>27</td>
<td>3</td>
<td>5.4</td>
<td></td>
</tr>
<tr>
<td>2 ax 197</td>
<td>74</td>
<td>22</td>
<td>17.3</td>
<td></td>
</tr>
<tr>
<td>4,000 r bx 263</td>
<td>100</td>
<td>14</td>
<td>8.5</td>
<td></td>
</tr>
<tr>
<td>3 ax 525</td>
<td>124</td>
<td>37</td>
<td>9.2</td>
<td></td>
</tr>
<tr>
<td>4,000 r bx 338</td>
<td>68</td>
<td>22</td>
<td>8.1</td>
<td></td>
</tr>
<tr>
<td>Combined ax 767</td>
<td>206</td>
<td>65</td>
<td>11.6</td>
<td></td>
</tr>
<tr>
<td>bx 683</td>
<td>195</td>
<td>39</td>
<td>7.9</td>
<td></td>
</tr>
</tbody>
</table>

ax - denotes desiccated and irradiated
bx - denotes irradiated only

(This work was supported by grant A-2162 from the National Institute of Health.)


Ovaries from third instar larvae of the genotype y/y; cand/cand were transplanted into third instar Oregon-R female larvae. Each survivor bearing an ovarian implant was mated, after eclosion, to a y B male. All but one of the females so mated produced no yellow or exceptional offspring due to failure of the transplanted ovary to become connected to an oviduct. However, from a female giving both y and y' progeny of both sexes, all of the 59 y' progeny were regular. Of the 20 y progeny obtained, three were exceptional males and one was a gynandromorph. The percentage of exceptional progeny (20%) from a homozygous cand ovary developing in a wild type female is not significantly different from the percentage of exceptions obtained in this laboratory from mating homozygous cand females.

Ronen, Amiram. Induced somatic recombination in the third chromosome of Drosophila melanogaster.

An attempt was made to study spontaneous and induced somatic recombination in the third chromosome of Drosophila melanogaster in flies of various genotypes. All individuals investigated were heterozygous for the same marker gene, Sb, carried on a structurally normal chromosome. Their X- and second chromosomes were either homozygous for the standard arrangement or heterozygous for various inversions. Third instar larvae of each genotype were given an X-ray dose of 1170 r (190 r per minute) at 80-90 hours after hatching and were then allowed to pupate. Individuals of each genotype were kept
as unirradiated controls. The adult flies were searched for normal bristles, assumed to be due to somatic crossing-over between the locus Sb and the centromere.

The bristles were scored according to a schedule fixed in advance. This schedule included 40 specified bristles on the head and thorax in the first series of the experiment, while 34 bristles of each individual were examined in the second experimental series (some of the bristles, numbers 8, 13 and 14 in Table 1, having been found to give only a small frequency of normals).

The frequency of normal bristles on the head and thorax was as high as 0.08 per irradiated fly among 4970 flies, but not a single normal bristle was found in 1590 unirradiated controls. The frequency of flies exhibiting normal bristles did not differ significantly between the various genotypes (the over-all \( X^2 \) test did not indicate any significant deviation from homogeneity), but different bristles showed different frequencies of normals (Table 1). It should be remembered that most of the bristles examined in these experiments (except for the humerals) are derived from the dorsal meso-thoracal imaginal disc.

The interpretation of the results on the basis of induced somatic crossing-over is complicated by several factors. The most important among these is the complete and unexpected absence of spontaneous recombination in the unirradiated controls.

However, it should be stressed that the frequency of normal bristles observed in the irradiated individuals appears too high to be accounted for by induced somatic back mutations. Even on the assumption that normal bristles may be due to induced mutations to suppressor genes of Sb at up to 10 different loci, in addition to back mutation at the Sb locus itself, the average mutation rate per r per locus would still have to be as high as \( 1.4 \times 10^{-7} \) in order to account for the observed effect.

### Table 1

<table>
<thead>
<tr>
<th>No.</th>
<th>Bristles</th>
<th>Experiment I</th>
<th>Experiment II</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>No. Normal</td>
<td>No. Normal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Mean</td>
<td>Mean</td>
</tr>
<tr>
<td>1-4</td>
<td>ocellars, orbitals, verticals, post verticals</td>
<td>16</td>
<td>56</td>
</tr>
<tr>
<td>5</td>
<td>ant. dorsocentrals</td>
<td>2</td>
<td>12</td>
</tr>
<tr>
<td>6</td>
<td>post. dorsocentrals</td>
<td>2</td>
<td>15</td>
</tr>
<tr>
<td>7</td>
<td>ant. postalars</td>
<td>2</td>
<td>19</td>
</tr>
<tr>
<td>8</td>
<td>post. postalars</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td>9</td>
<td>ant. scutellars</td>
<td>2</td>
<td>18</td>
</tr>
<tr>
<td>10</td>
<td>post. scutellars</td>
<td>2</td>
<td>10</td>
</tr>
<tr>
<td>11</td>
<td>supralars</td>
<td>2</td>
<td>}</td>
</tr>
<tr>
<td>12</td>
<td>ant. notopleurals</td>
<td>2</td>
<td>}</td>
</tr>
<tr>
<td>13</td>
<td>post. notopleurals</td>
<td>2</td>
<td>33</td>
</tr>
<tr>
<td>14</td>
<td>prescutellars</td>
<td>2</td>
<td>}</td>
</tr>
<tr>
<td>15</td>
<td>humerals</td>
<td>4</td>
<td>}</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>40</td>
<td>170</td>
</tr>
</tbody>
</table>
Schepers, A. M. An interaction in Pteridine metabolism between garnet and brown genes in D. melanogaster.

Two-dimensional paper chromatograms using various solvents were made of extracts from 25 heads. All flies used were three days old. A fluoroscopic comparison was made of wild type, a garnet allele resembling g2, a brown allele, and the double mutant g ; bw. Both mutants were found to differ from wild type only quantitatively in respect to their pteridine patterns. In particular, 2-amino-4-hydroxypteridine is present in both of them although in lower concentration than in wild type. The double mutant, on the other hand, contains no detectable quantities of this substance. The activity of xanthine dehydrogenase which has been implicated in formation of pteridines, was determined following the method described by Mitchell, and measured fluorometrically. Enzyme activity in the double mutant was found to be present though reduced with respect to wild type.

Schulten, G. M. A case of aberrant sex-ratio in D. melanogaster.

A Kr/Cy stock which had been selected for a high penetrance of Kr gave in contrast to the original stock a sex-ratio of about 1♀ : 4♂ at 25°C. At lower temperatures the ratio shifts towards normal, with a 1♀ : 1.5♂ ratio at 17°C.

When the males of this stock were crossed to unrelated stocks, the aberrant sex-ratio did not reappear in F1 or subsequent generations. Also, crosses of "sex-ratio" females to unrelated males gave normal sex-ratios, save in the cross to Cy-Oster/Fm ; Ubx130/Sb males in which case there was a deficiency in females of the genotype Cy-Oster/Kr resulting in a ratio of 1♀ : 2.5♂.

Substitution, in the "sex-ratio" stock, of a Fm-chromosome for the Cy-chromosome, resulted in a ratio of about 1♀ : 1.5♂. After substitution of a Fm-chromosome for the Kr-chromosome, a ratio of about 1♀ : 3♂ was obtained. The sex-ratio became normal in all cases where the X-chromosomes of the "sex-ratio" stock had been replaced by X-chromosomes from other sources.

It is tentatively assumed that this temperature-sensitive deviation from a normal sex-ratio is caused by anomalous behaviour of a mutant X-chromosome depending on the presence of the Cy and Kr second chromosomes.

Schwinck, I. Drosopterin formation and semi-lethality of the mutant rosy in temperature experiments.

The pleiotropic pattern of the mutant rosy (ry) includes the following characters:

1. reduced amount of drosopterins (red eye pigments),
2. nonautonomous formation of drosopterins as demonstrated in transplantation experiments, increased drosopterin formation at low breeding temperature,
3. semi-lethality in the late pupal stage and during emergence of the fly,
4. aberrant morphology and function of the Malpighian tubes,
5. no xanthine dehydrogenase activity and no accumulation of isoxanthropterin and uric acid.

In course of the study of cause and relation of these characters and their dependence on temperature, first the critical time for the manifestation of drosopterin quantity was determined. For the strain v ; ry2 this was found to be the very late pupal stage, as revealed by changing the breeding temperature from 18°C to 26°C, and vice versa, at the following stages: (a) 1. larval stage, (b) early 3. larval stage, (c) prepupa, (d) pupa 36 hours after pupation, (e) pupa with beginning red pigment formation, (f) imago 0-2 hours after emergence. It seems rather interesting that the critical time determining the drosopterin quantity is the period when actually the drosopterin pigments are deposited in the eyes. Furthermore, the temperature effect on the drosopterin quantity and on the semi-lethality of the late pupa and emerging flies was studied in various rosy alleles and compounds, as well as in isogenic cn ; ry2 strains. With decreasing temperature the drosopterin formation increases parallel to increasing viability of pupae and young flies. For the radiation induced rosy alleles ry4, ry6, ry8 and ry9 (Chovnick, A., A. Schalet, and R. P. Kernaghan, Rec.
Genetics Soc. America 30, p. 68, and Genetics 46, p. 858, 1961) the temperature effect on both, the drosoperin formation and the semi-lethality, is similar to ry⁴ and ry⁵; the death rate in the late pupal stage being about 40-60% for a breeding temperature of 26°C, as compared with below 5% for ry⁺ strains. However, for the compounds of these rosy alleles the semi-lethality at 26°C, decreases in certain crosses below 5%, although the eye color is typical for 26°C rosy-breed and the aberrant morphological structure of the Malpighian tubes could not be distinguished from those of rosy strains with low emergence rate. A rather strong influence of the genetic background on the temperature dependence of the drosoperin quantity and the pupal semi-lethality was found in experiments with various isogenic on; ry² strains. Crossing flies from certain isogenic strains with flies from the original ry² stock resulted in an improved viability at 26°C, the pupal death rate being about 10% in the F₁ as compared with about 50% in the parental stocks raised in the same incubator.

(Supported by a grant from the USPHS RG-7464.)

Seki, T. Absence of beta-alanine in hydrolyzate of the pupal sheaths of ebony mutant of D. virilis.

After washing in water and drying in the air, 50 mg of pupal sheaths were homogenized with 80% ethanol and filtered and washed with 80% ethanol, followed by 99% ethanol. The residue was hydrolyzed in 6 N HCl at 110°C for 24 hours in a sealed tube.

The fluid was evaporated rapidly by using rotary evaporator and dissolved in 2 ml of distilled water. One ml of the solution was added on a column of Amberlite CG-120 (H form, 0.8 X 140 cm) and eluted with 1.2 N HCl. The effluent was collected in fractions of 2.9 ml. Each fraction was neutralized with sodium hydroxide solution and assayed according to the method of Yemm and Shen.

No beta-alanine was detected in the hydrolyzate of pupal sheaths of ebony mutant strain, in contrast with that of wild strain, in which a considerable amount of beta-alanine was present. Similar results were obtained with black pupa mutant strain of Musca domestica and with sooty mutant of Bombyx mori.

Sherwood, Eva R. All-male offspring from heatshocked cultures.

Accidental exposure of a few hours' duration to heatshock of 31°C of four bottles with crosses of y² sc wa ec, sc⁸,Y ¿¿ x ywB ¿¿ yielded only male progeny (a total of approximately 250-300) of ywB, sc⁸,Y constitution, except for 2 females of the expected genotype. At 25°C, approximately equal numbers of male and females were produced. Repeated single pair crosses resulted in the same sex ratio effect, when the shock was given on the 4th, 5th or 7th day after the start of the cultures. Those shocked before the third day after mating had no offspring.

Heatshock to crosses involving different attached and unattached X chromosomes, as well as the sc⁸,Y carried in male or female showed that the particular attached XXs of y² sc wa ec constitution were responsible for the lack of female progeny in the next generation.

Shima, T., A. Kaneko, and E. Momma. Hatchability of eggs during varying lapses from the time of mating in D. virilis.

Preliminary examinations were made in order to analyse the sensitivity for X-rays to spermatogenesis. Fifteen young virgin females were mated singly in vials each with a single young male for three days, and then the females were kept without males. Eggs were laid and their hatchabilities were observed every day during 15 days after the separation from males. As shown in Table 1, the largest number of eggs (29.5 per single female) were observed on the 5th day of single culture, the smallest one (1.4) being on the 8th day. Hatchability
of the eggs laid on every day was about 90 per cent within the first ten days. From the 11th day on, however, it showed a remarkable decrease (6%). No egg hatched out from those laid after the 12th day. Dissected seminal receptacles showed a rapid decrease of sperm after the 11th day.

Table 1

<table>
<thead>
<tr>
<th>Days after mating</th>
<th>No. of eggs laid</th>
<th>No. of hatched eggs</th>
<th>Hatchability (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>86</td>
<td>81</td>
<td>94</td>
</tr>
<tr>
<td>2</td>
<td>120</td>
<td>107</td>
<td>89</td>
</tr>
<tr>
<td>3</td>
<td>93</td>
<td>86</td>
<td>93</td>
</tr>
<tr>
<td>4</td>
<td>90</td>
<td>86</td>
<td>93</td>
</tr>
<tr>
<td>5</td>
<td>443</td>
<td>434</td>
<td>98</td>
</tr>
<tr>
<td>6</td>
<td>167</td>
<td>158</td>
<td>95</td>
</tr>
<tr>
<td>7</td>
<td>357</td>
<td>346</td>
<td>97</td>
</tr>
<tr>
<td>8</td>
<td>20</td>
<td>15</td>
<td>75</td>
</tr>
<tr>
<td>9</td>
<td>50</td>
<td>42</td>
<td>84</td>
</tr>
<tr>
<td>10</td>
<td>266</td>
<td>244</td>
<td>92</td>
</tr>
<tr>
<td>11</td>
<td>235</td>
<td>14</td>
<td>6</td>
</tr>
<tr>
<td>12</td>
<td>93</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>13</td>
<td>28</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>14</td>
<td>173</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>15</td>
<td>51</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>2272</td>
<td>1613</td>
<td>71</td>
</tr>
</tbody>
</table>

Snyder, L. A. The effect on TEM-induced mutations and translocations of storing treated spermatozoa in the female.

Studies by Schalet (1955: Genetics 40, 594), and Herskowitz (1956: Genetics 41, 605) on the effects of storing nitrogen mustard- or triethylene melamine-treated spermatozoa in females, revealed sharp increases in translocation frequencies, with no sensible change in frequencies of sex-linked recessive lethals or chromosome losses. The post-copulatory vaginal douche treatments of females admit to uncertainty in interpretation since the chemicals used are highly penetrating and would be expected to reach the ovaries of the females in which treatment was carried out. Similar experiments were repeated, using inter-abdominal injection of day-old males, and the results were in agreement with those reported earlier. Using $2 \times 10^{-4}$ triethylene melamine in 0.7% saline resulted in a 4-fold increase in translocations after 3 days storage in the female of treated spermatozoa, with no increase in sex-linked recessive lethals or chromosome losses.


The present experiments describe the effect of selection on the expression of the ocelli-less mutant, a sex-linked recessive, variable for the number of bristles and ocelli on top of the head in Drosophila subobscura. Earlier experiments had shown that if selection for higher numbers of structures were practised on a population homozygous for the mutant, it was possible to obtain a population in which a high
proportion of flies had a wild-type phenotype. There was found to be a partial barrier at the wild-type phenotype, preventing progress beyond the "score" of six bristles and three ocelli (Fig. 1 A). The frequency of repeated bristles (two or more bristles lying close together at the site normally occupied by one) was found to be much greater than in the foundation population. In a few flies more than three ocelli were seen. In the present experiments an attempt was made to observe the effect of continued selection, in the hope that an increase in the frequency of genes for higher score might bring forth the expression of new structures.

The results of the present experiment show the appearance of a novel pair of bristles. These bristles (Fig. 1 B) always appear at a specific site and have a definite orientation. The "neomorphs" are normally absent in Drosophilidae but they are present (Fig. 1 C) in a family closely related to it, the Aulacigastriidae.

The appearance of these structures as a consequence of continued selection on the mutant population, after the wild-type phenotype had been reached, suggests that a particular frequency of genes for higher score is required for their expression. This is also suggested by the higher frequency of neomorphs in females, which have a higher mean number of bristles and ocelli than males.

A hypothesis is postulated, similar to that suggested by Stern (1954, Proc. 9th Intern. Congr. Genet., 6:355-369), to explain the origin of neomorphs in terms of an unvarying "prepattern" which determines the positions of these structures and
a common "precursor" of bristles and ocelli which must be present in the required amount if structures are to be formed. It is suggested that if the amount of the precursor is increased to a certain threshold, a new bristle is formed in response to the peak of the prepattern which is present in the wild-type flies, but to which wild-type cells are not competent to respond.

Sperlich, D. Hybrids between D. melanogaster and D. simulans in nature.

In Drosophila samples caught in May/June at the island of Lipari (South Italy) the most frequent species observed were D. melanogaster and D. simulans. Whereas the males of the two species can be easily detected and separated, the females are practically identical in appearance. In order to determine the frequency of females in nature, cultures from individual females, mated already in nature, were reared. Out of a total of 141 cultures the examination of the male offspring showed that 81 (57.5%) of the females belonged to D. melanogaster and 53 (37.5%) to D. simulans. The remaining 7 cultures (5%) gave only sterile female offspring of a typical appearance, most probably resulting from the cross D. melanogaster ♀ x D. simulans ♂ in nature. This frequency of cross-matings is surprisingly high and cannot be explained just as a chance happening. It may be noted that crosses D. melanogaster ♀ x D. simulans ♂ with Lipari-strains are almost always successful also in the laboratory. It seems that the sexual isolation between these two species at the island is not so strong as on other places, perhaps on a genetical basis. But we could not establish the existence of the reciprocal matings in nature.

Stern, Curt, and Eva R. Sherwood.

Can primordial germcells of the genotype XXY produce functional sperm?

Spermatogonia with two X chromosomes in transformed phenotypic tra male individuals are unable to develop into functional sperm (Seidel, 1960, Naturw.) but the recovery of clusters of sperm with attached X chromosomes from 3XY males (Neuhaus, 1937, Genetics) suggests that spermatogonia with two X chromosomes in males of non-tra genotype can develop normally. An attempt was made to discover gonosomal mosaics in which the soma is normally XY male and part or all of the testes contain primordial germcells with two X chromosomes. Would they form functional sperm? Frost (1961, DIS 35:81) has shown that somatic double nucleus mosaics occur among the offspring of 3N females with a low frequency. Therefore, sons of 3N females y² sc w² ec, FM4 mated to FM4, sc²,Y were progeny tested. These sons were somatically FM4, sc²,Y. Would an occasional individual contain functional germcells derived from an attached-X female pronucleus fertilized by a sc²,Y sperm? If so, y² sc w² ec daughters should be observed among the offspring of the FM4, sc²,Y males mated to attached-X females of the yf genotype. None of 3,744 FM4 males, mated in groups of 6, produced y² sc w² ec daughters. In a similar experiment, none of 4,740 wvB sons from the cross yf x wvB when mated to y² sc w² ec, sc²,Y females produced any yf, sc²,Y daughters. Thus, either none of the total of nearly 8,500 males were gonosomal XXY-XY mosaics, or no functional sperm is formed by cells derived from XXY primordial germcells. Only experiments on a much larger scale than those reported here can furnish a decision.


During the summers of 1950, 1951, 1952, and 1954, collections of D. robusta were made from several sites on Unaka Mountain in northeastern Tennessee. Salivary gland preparations were made, and, at the time, the samples were considered too small to be statistically significant. The data, however, may be of some interest to other workers.
The accompanying table shows the frequencies of inversion heterozygotes at different elevations. These data are in sharp contrast to those of Stalker and Carson (1948), who collected in the Smoky Mountains.

It is suggested that the Unaka collections were, in the main, from marginal populations (in the sense of Carson, 1955). During the 1920's a fire destroyed much of the forest land, and most of the collecting sites were in or very near areas that had not yet restocked. Selection pressure at these sites was evidently quite severe. The collecting sites have been described elsewhere (Stevenson, 1952).

Large collections could not be made because of the inaccessibility of some of the stations, and, later, several of the stations were disturbed by logging operations.


Table to show the frequencies of inversions in D. robusta at different altitudes on Unaka Mountain, Tenn.

<table>
<thead>
<tr>
<th>Altitude</th>
<th>n</th>
<th>XL</th>
<th>XL-1</th>
<th>XL-2</th>
<th>XL-3</th>
<th>XR</th>
<th>XR-1</th>
<th>XR-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>2200 ft.</td>
<td>36</td>
<td>.92</td>
<td>.052</td>
<td>.026</td>
<td>.89</td>
<td>.056</td>
<td>.056</td>
<td></td>
</tr>
<tr>
<td>2500</td>
<td>48</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3000</td>
<td>44</td>
<td>.98</td>
<td>.021</td>
<td>1.00</td>
<td>.96</td>
<td>.042</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3500</td>
<td>9</td>
<td>1.00</td>
<td></td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3760</td>
<td>2</td>
<td>1.00</td>
<td></td>
<td></td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4080</td>
<td>63</td>
<td>.92</td>
<td>.091</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4200</td>
<td>24</td>
<td>.83</td>
<td>.007</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4400</td>
<td>26</td>
<td>1.00</td>
<td></td>
<td></td>
<td>.81</td>
<td>.038</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4800</td>
<td>10</td>
<td>1.00</td>
<td></td>
<td></td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>2L</th>
<th>2L-1</th>
<th>2L-2</th>
<th>2L-3</th>
<th>2R</th>
<th>2R-1</th>
<th>3R</th>
<th>3R-1</th>
</tr>
</thead>
<tbody>
<tr>
<td>2200</td>
<td>36</td>
<td>.81</td>
<td>.083</td>
<td>.083</td>
<td>.028</td>
<td>.97</td>
<td>.028</td>
<td>.86</td>
</tr>
<tr>
<td>2500</td>
<td>48</td>
<td>.79</td>
<td>.145</td>
<td>.016</td>
<td>.049</td>
<td>.98</td>
<td>.016</td>
<td>.90</td>
</tr>
<tr>
<td>3000</td>
<td>44</td>
<td>.95</td>
<td>.023</td>
<td>.023</td>
<td>.91</td>
<td>.091</td>
<td>.98</td>
<td>.023</td>
</tr>
<tr>
<td>3500</td>
<td>9</td>
<td>.67</td>
<td></td>
<td>.333</td>
<td>1.00</td>
<td>.89</td>
<td>.112</td>
<td></td>
</tr>
<tr>
<td>3760</td>
<td>2</td>
<td>1.00</td>
<td></td>
<td>1.00</td>
<td></td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4080</td>
<td>63</td>
<td>.92</td>
<td>.042</td>
<td>.042</td>
<td>1.00</td>
<td>.80</td>
<td>.200</td>
<td></td>
</tr>
<tr>
<td>4200</td>
<td>24</td>
<td>.88</td>
<td></td>
<td></td>
<td>1.00</td>
<td>.73</td>
<td>.270</td>
<td></td>
</tr>
<tr>
<td>4400</td>
<td>26</td>
<td>1.00</td>
<td></td>
<td></td>
<td>1.00</td>
<td>.80</td>
<td>.200</td>
<td></td>
</tr>
<tr>
<td>4800</td>
<td>10</td>
<td>1.00</td>
<td></td>
<td></td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Stone, L. E. Structure and variation of the salivary gland chromosomes in Drosophila affinis. Standard salivary gland chromosome maps have been prepared for D. affinis using a homozygous strain from central Nebraska. The chromosomes of this species have seven large euchromatic arms and a dot-like element. These represent a V-shaped X-chromosome (LX and SX) and four pairs of autosomes, which includes a J (3L and 3S), a near-V (2L and 2S), a rod (4) and a dot-like element (5).
Chromosomal variation is being investigated by mating males from various laboratory stocks to females carrying the Standard sequence. A number of slides have been prepared with lactic-acetic orcein, but the best success has been obtained with a modification of Cohen's Sudan Black B reagent (Cohen, 1949, Stain Technol., 24:177-184). Preliminary data on chromosomal variation has been gathered from a number of strains representing 35 localities in 18 states (Table 1).

Table 1

<table>
<thead>
<tr>
<th>State</th>
<th>Number of Strains</th>
<th>Sequences besides Standard*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alabama</td>
<td>3</td>
<td>SX/SX-1</td>
</tr>
<tr>
<td>Florida</td>
<td>4</td>
<td>LX/LX-1; 4/4-3</td>
</tr>
<tr>
<td>Georgia</td>
<td>2</td>
<td>SX/SX-1</td>
</tr>
<tr>
<td>Iowa</td>
<td>2</td>
<td>4/4-2</td>
</tr>
<tr>
<td>Kentucky</td>
<td>1</td>
<td>?*</td>
</tr>
<tr>
<td>Louisiana</td>
<td>10</td>
<td>LX/LX-1; SX/SX-1</td>
</tr>
<tr>
<td>Massachusetts</td>
<td>1</td>
<td>?*</td>
</tr>
<tr>
<td>Minnesota</td>
<td>3</td>
<td>LX/LX-1; SX/SX-1</td>
</tr>
<tr>
<td>Mississippi</td>
<td>2</td>
<td>LX/LX-1; 4/4-2</td>
</tr>
<tr>
<td>Missouri</td>
<td>2</td>
<td>LX/LX-1; SX/SX-1</td>
</tr>
<tr>
<td>Nebraska</td>
<td>53</td>
<td>LX/LX-1; LX/LX-2; LX/LX-4; 4/4-2</td>
</tr>
<tr>
<td>New York</td>
<td>1</td>
<td>SX/SX-1</td>
</tr>
<tr>
<td>North Carolina</td>
<td>1</td>
<td>LX/LX-3; 4/4-2</td>
</tr>
<tr>
<td>Ohio</td>
<td>1</td>
<td>4/4-2</td>
</tr>
<tr>
<td>Oklahoma</td>
<td>1</td>
<td>LX/LX-1; SX/SX-1</td>
</tr>
<tr>
<td>South Carolina</td>
<td>2</td>
<td>LX/LX-1; SX/SX-1</td>
</tr>
<tr>
<td>Tennessee</td>
<td>2</td>
<td>?*</td>
</tr>
<tr>
<td>Texas</td>
<td>2</td>
<td>LX/LX-1</td>
</tr>
</tbody>
</table>

*In addition to the non-Standard sequences listed in Table 1 a complex configuration has been found in the distal part of chromosome four. It has been found in at least one strain from each of the states listed in the table. Since the pairing in this configuration is so variable, it has not yet been determined how many different inversions may be involved or what differences there may be between strains.

Excluding the dot-like element there are four rather long arms that have been found to contain only the Standard sequence in all of the strains tested to date.

**Strangio, V. A.** Recessive lethals, sex chromosome loss, and nondisjunction followed simultaneously. The incidence of sex-linked recessive lethals induced at various stages in the spermatogenesis of *D. melanogaster* has been investigated simultaneously with the induction of sex-chromosome loss and nondisjunction of X and Y chromosomes. This has been achieved by the use of a modified Muller-5 (Basc) stock in which the Bar marker had reverted to wild type. The daily brood technique involved the mating of fresh virgin females of this stock to irradiated males (800r) carrying a normal X and the doubly-marked y+w Y BS. In the F1, sex chromosome loss and marker deletion from the Y were scored as exceptional non-Bar males, primary nondisjunction of X and Y as Bar females. Impregnated F1 females were placed individually in minilinks to provide an F2 from which the sex-linked lethal frequency was obtained. If the first
appearance of induced nondisjunction is accepted as a valid reflection of the irradiation of early meiotic stages, then the condensed results presented in the following table indicate two radio-sensitive peaks during spermatogenesis, one in spermatids (5th day) for recessive lethals; the other in spermatocytes (7th day) for breakage-loss aberrations. Detailed discussion will be published elsewhere.

<table>
<thead>
<tr>
<th>Daily Brood</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>% lethals</td>
<td>4.67</td>
<td>5.43</td>
<td>3.61</td>
<td>2.92</td>
<td>3.30</td>
</tr>
<tr>
<td>% non Bar dd</td>
<td>0.23</td>
<td>0.23</td>
<td>0.84</td>
<td>1.70</td>
<td>1.47</td>
</tr>
<tr>
<td>% Bar ??</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.43</td>
<td>0.17</td>
</tr>
</tbody>
</table>

The investigation of possible pseudo-allelism at the spineless-aristapedia locus is in progress by means of a crossover selector technique. The work is at present being hampered by the low fertility of the selector cross males.

Previously the authors (1959, 1960) reported that in several strains of *D. melanogaster* spontaneously there occurs definite deficiency of the compound eye. The occurrence is quite seldom on standard medium, but may be considerably increased on experimental media containing soybean powder or monosodium glutamate. Further it has been shown that these deficient eyes, once appeared, are inherited if inbred by means of sib-mating, while crossing of them by the wild type generally fails to be reproduced. Especially it was striking that the eye-deficiency is inherited not only by the same degree but also by higher ones. Therefore, if inbreedings were properly continued, gradual diminution and, in extreme cases, complete disappearance of the compound eye eventually results. On this account, these are designated as 'erosion eye.' Since these facts suggest that the erosion eye is a hereditary character which represents peculiar mode of inheritance, further evidences were collected in the following experiments.

In an attempt to prepare a pure strain of fly which produces no erosion progeny, a selection experiment was carried out with wild type flies of Oregon. In the pair matings carried out randomly with flies of this stock, frequency of the erosion eye was about 0.3%. Later it was recognized that among flies of this stock there are ones the compound eyes of which represent slight irregularities in outline. Mating of these was liable to produce erosion offspring more often. In the present selection, therefore, special care was taken not to choose these aberrant flies. Even in this way erosion offsprings were found to be produced in the first 3 generations of selection. But in later generations erosion flies were ceased to be produced, and in none of the offsprings successively examined during 50 generations the eye deficiency was met with (Table 1). However, when offspring flies produced in every generation of this selection were reared on the medium containing 10% of soybean powder, occurrence of the erosion progeny was not only increased in rate but also continued in still later generations. In this medium the rates obtained were 2 or 3 times higher than those in the standard medium, and the occurrence was continued till the F6 generation. Among the offsprings produced in later generations than this, there was none which presented erosion eyes. Therefore, it may be safe to assume that the offspring flies produced in relatively later generations of the selection can be regarded as a pure strain which are free from the erosion-eye producing factors.
In a second experiment crossing was tested between erosion flies and wild types. The erosion flies were chosen from the strains kept in our laboratory and the grade of erosion was very low, the facets of the compound eye counting about 600. The wild types were of 2 sources: one from unselected strain and the other from offsprings of the F50 generation of the pure strain. When the crossing was done by using wild types of the unselected strain, erosion offsprings were produced, being found in 4 out of 13 matings examined. Rates of the occurrence considerably varied in individual matings, but the rate averaged throughout the cases was 0.44%. The same crossing tested on the soybean medium produced erosion progeny far more frequently than on the standard one. The average rate amounted 1.42% (Table 2A). When wild type flies of the pure strain were used in the crossing, all the offsprings produced failed to represent the eye deficiency being normal so far as they were reared on the standard medium. But the same crossing tested on the soybean medium brought about erosion flies invariably in all the matings examined. Rates of the occurrence were nearly equal in individual cases, and, moreover, they were very high, the average being 2.29% (Table 2B).

From the results above mentioned it can be surmised that spontaneous occurrence of the erosion eye is due to certain hereditary factors which are widely distributed among population of Oregon stock. By means of the inbreeding properly carried out for several successive generations, these factors can be swept off so that the flies are quite unable to produce erosion progeny. These flies again acquire the ability to produce erosion offsprings if they were crossed with the flies possessed with the factors. Further it was striking that the phenotypic representation of the erosion eye is shown to be influenced by the medium on which flies were reared.

Table 1

Frequencies of erosion flies produced in selection experiment carried out by means of pair-matings of wild type flies

<table>
<thead>
<tr>
<th>Generation</th>
<th>Number of total flies</th>
<th>Number of erosion flies</th>
<th>Per cent of erosion flies</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>634 (508)</td>
<td>2 (11)</td>
<td>0.29 (2.17)</td>
</tr>
<tr>
<td>F2</td>
<td>410 (573)</td>
<td>0 (7)</td>
<td>-</td>
</tr>
<tr>
<td>F3</td>
<td>402 (425)</td>
<td>1 (2)</td>
<td>0.25 (0.47)</td>
</tr>
<tr>
<td>F4</td>
<td>480 (492)</td>
<td>0 (3)</td>
<td>-</td>
</tr>
<tr>
<td>F5</td>
<td>573 (260)</td>
<td>0 (1)</td>
<td>-</td>
</tr>
<tr>
<td>F6</td>
<td>649 (487)</td>
<td>0 (1)</td>
<td>0.25 (0.25)</td>
</tr>
<tr>
<td>F7</td>
<td>214 (522)</td>
<td>0 (0)</td>
<td>-</td>
</tr>
<tr>
<td>F8</td>
<td>526 (613)</td>
<td>0 (0)</td>
<td>-</td>
</tr>
<tr>
<td>F9</td>
<td>139 (439)</td>
<td>0 (0)</td>
<td>-</td>
</tr>
<tr>
<td>F10</td>
<td>225 (341)</td>
<td>0 (0)</td>
<td>-</td>
</tr>
<tr>
<td>F50</td>
<td>607 (735)</td>
<td>0 (0)</td>
<td>-</td>
</tr>
</tbody>
</table>

Numerals without parentheses represent results on standard medium and those within parentheses results on soybean medium.
Table 2
Results of crossing erosion flies by wild types
of different strains

<table>
<thead>
<tr>
<th></th>
<th>Standard medium</th>
<th>Soybean medium (10%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total flies</td>
<td>Erosion flies</td>
<td>%</td>
</tr>
<tr>
<td>A. Wild type flies of unselected strain</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1597</td>
<td>7</td>
<td>0.44</td>
</tr>
<tr>
<td>B. Wild type flies of selected strain</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1474</td>
<td>0</td>
<td>-</td>
</tr>
</tbody>
</table>

Thompson, Peter E. The basis
for high "nondisjunction" from
maroon-like females.

Females homozygous for the mutant maroon-like
(ma-l) produce unusually high frequencies
of apparent primary nondisjunction. The
exceptional offspring which result are
predominantly male; the frequency of exceptional females is not appreciably greater
than the normal rate (see Spieler, Rec. Gen. Soc. Am., 1961). In this study crosses
of y² ma-l/y² ma-l; T(3;4)89E/+; y² ma-l/y²; T(3;4)89E/+; and y² ma-l/y² ma-l females
to w males were made to test whether the X's tend to interact with unpaired auto-
somal elements when the translocation is present. The disruption of homologies by
heterozygous T(3;4)89E results in appreciable frequencies of haplo-4 and triplo-4
offspring. If failure of pairing is the basis of nondisjunction in ma-l lines, the
availability of nonhomologous elements for pairing should enhance the effect (Grell
and Grell, 1960). The progeny of the above crosses were:

Series A: y² ma-l / y² ma-l; T(3;4)89E/+♀ x w³♂

<table>
<thead>
<tr>
<th></th>
<th>y² ma-l♂♂</th>
<th>y² ma-l♀♀</th>
<th>w³♂♂</th>
<th>w³♀♀</th>
</tr>
</thead>
<tbody>
<tr>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td></td>
</tr>
<tr>
<td>2058</td>
<td>91</td>
<td>1691</td>
<td>62</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>excep.♀♀ = .09%</td>
<td>excep.♂♂ = 1.35%</td>
<td>haplo-IV = 3.9%</td>
<td></td>
</tr>
</tbody>
</table>

Series B: y² ma-l/y²; T(3;4)89E/+♀ x w³♂

<table>
<thead>
<tr>
<th></th>
<th>y²♂♂</th>
<th>y²♀♀</th>
<th>w³♂♂</th>
<th>w³♀♀</th>
</tr>
</thead>
<tbody>
<tr>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td></td>
</tr>
<tr>
<td>2208</td>
<td>95</td>
<td>1966</td>
<td>79</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>excep.♀♀ = .43%</td>
<td>excep.♂♂ = .68%</td>
<td>haplo-IV = 4.1%</td>
<td></td>
</tr>
</tbody>
</table>

Series C. y² ma-l/y² ma-l♀♀ x w³♂♂

<table>
<thead>
<tr>
<th></th>
<th>y² ma-l♂♂</th>
<th>y² ma-l♀♀</th>
<th>w³♂♂</th>
<th>w³♀♀</th>
</tr>
</thead>
<tbody>
<tr>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td>diplo-4 haplo-4</td>
<td></td>
</tr>
<tr>
<td>2501</td>
<td>0</td>
<td>2082</td>
<td>3</td>
<td>31</td>
</tr>
<tr>
<td></td>
<td>excep.♀♀ = .12%</td>
<td>excep.♂♂ = 1.4%</td>
<td>haplo-IV = 0.0%</td>
<td></td>
</tr>
</tbody>
</table>
The presence of homozygous ma-l and the translocation in Series A did not result in any appreciable increase in X-chromosome exceptions over crosses lacking the translocation (Series C), or in chromosome-4 nondisjunction over crosses where ma-l was not homozygous (Series B). Furthermore, no coincidence of X and 4 nondisjunction was observed in Series A; this absence of coincidence even falls below an expectation based on the presence of the translocation alone (see Series B).

It appears that the basis for the occurrence of exceptional types from maroon-like females is not a looseness or failure of pairing. The great predominance of exceptional males over exceptional females suggests chromosome loss or elimination as the underlying mechanism.

Tsukamoto, M. Comparative studies on the oxidation of DDT in D. melanogaster. Analyses of DDT, which have a replaceable hydrogen atom in the alkyl moiety between two p-chlorophenyl groups, have been metabolized to their corresponding alcohols. On the other hand, para-substituted analogues of DDT were rapidly metabolized but the corresponding alcoholic metabolites could not be detected among the recovered fractions.

Toyofuku, Y. Non-random association of inversions in D. nigromaculata. In DIS 36, the author reported that in D. nigromaculata in natural population there occur twenty-two different kinds of chromosomal aberrations, all being represented by heterozygous inversions. Cytological analyses of these chromosomal aberrations have been carried out in more detail. This species showed a variety of inversions in each arm of two or more chromosomes. Frequency distributions of inversions in each chromosome were as follows: C-chromosome showed 34.6% in frequency, D-chromosome ranked next, showing 25.7%, A-chromosome occurred at 24.76% of a total inversion, B-chromosome showed 8.91%, and the X-chromosome was found at the lowest frequency of 5.94%. The results are referred to in table. It is interesting to note that in C-chromosome, for example, the inversion d includes sections 65C - 67B often appeared in combination with the inversion b and inversion e involves sections 67C - 72A, containing new inversion c. This suggests that the distribution of inversions is not random.

<table>
<thead>
<tr>
<th>Chromosome type</th>
<th>Part of inversion</th>
<th>Inversion observed</th>
<th>Frequencies (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2</td>
</tr>
<tr>
<td>X a</td>
<td>3B - 9A</td>
<td>1</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5,94</td>
</tr>
<tr>
<td>X b</td>
<td>5A - 9C</td>
<td>4</td>
<td>67</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>82.5</td>
</tr>
<tr>
<td>X c</td>
<td>5A - 9C, 12A - 15B</td>
<td>1</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5,94</td>
</tr>
<tr>
<td>A a</td>
<td>26A - 33A</td>
<td>5</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>6-8</td>
</tr>
<tr>
<td>A b</td>
<td>26A - 36A</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4-6</td>
</tr>
<tr>
<td>A c</td>
<td>29A - 32B</td>
<td>10</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>24.76</td>
</tr>
<tr>
<td>A d</td>
<td>32B - 34A</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4-6</td>
</tr>
<tr>
<td>A e</td>
<td>28B - 30A, 32B - 37C</td>
<td>4</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>16-18</td>
</tr>
<tr>
<td>A f</td>
<td>32B - 34A, 35A - 39A</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8-10</td>
</tr>
<tr>
<td>A g</td>
<td>23D - 27A, 28B - 30A, 32B - 37C</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8-10</td>
</tr>
<tr>
<td>B a</td>
<td>42A - 51D</td>
<td>3</td>
<td>33</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8.91</td>
</tr>
<tr>
<td>B b</td>
<td>49A - 53A</td>
<td>2</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8-10</td>
</tr>
<tr>
<td>B c</td>
<td>52A - 55B</td>
<td>1</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8-10</td>
</tr>
<tr>
<td>B d</td>
<td>48C - 50D, 52A - 55B</td>
<td>3</td>
<td>33</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8-10</td>
</tr>
</tbody>
</table>
Valencia, Ruby M., and J. I. Valencia. Evidence for a non-chromosomal origin of dominant lethals. The frequencies which we obtained for dominant lethals induced by X-rays in recently fertilized eggs (5-15 minutes after fertilization, supposedly before union of pronuclei) turned out to be very close to those obtained by Parker (1959, The University of Texas Publication, No. 5914:113) in stage 14 oocytes. Lindsley (personal communication) observed that since adding the male genome has little effect, the majority of dominant lethals induced in these stages apparently are not of chromosomal origin. We considered that this idea was worth a careful test and therefore set up a series of irradiations to test dominant lethal frequency in stage 14 oocytes, using a dose of 500r and exactly the same stocks and crossing schemes that we were using for the fertilized eggs. We also tested mature sperm (ejaculated within 8-10 hours after treatment), using these same stocks and schemes, and have accumulated some data from treated embryos (35-45 minutes after fertilization). The data are as follows:

| Chromosome | Inversion type | Part of inversion | Inversion observed | Frequencies (%) |
|------------|----------------|------------------|-------------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
|            |                |                  |                   | 1              | 2              |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
| C          | a              | 62B - 64A        | 6                 | 17             |                |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
|            | b              | 72B - 77A        | 3                 | 9              | 34.65          |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
|            | c              | 77A - 79D        | 7                 | 20             |                |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
|            | d              | 65C - 72B, 72B - 77A | 13              | 37             |                |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
|            | e              | 67C - 72A, 77A - 79D | 6              | 17             |                |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
| D          | a              | 88A - 93B        | 16                | 62             |                |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
|            | b              | 97A - 96B        | 9                 | 35             | 25.74          |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
|            | c              | 90B - 94B, 96A - 97B | 1              | 4              |                |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |

Treated: 6864 67.5 6972 64.7 2128 69.8 6475 18.2
Control: 6190 7.8 1519 8.0 1519 8.0 5101 8.4
Corrected: 59.7 61.8

* i.e. % unhatched eggs. Corrected value represents death in embryonic stage.

The values we have obtained for stage 14 oocytes are in quite good agreement with the values obtained by Parker (above reference) and the frequency obtained for recently fertilized eggs agrees well with the results of Ulrich (1960, Revue Suisse de Zoologie, 67:287). It seems clear that oocytes, with one nucleus, recently fertilized eggs, with two nuclei, and embryos, with several nuclei, all have about...
the same mortality (around 60%) after 500r. The mortality induced by irradiating sperm is not detectably added to the mortality induced by irradiating oocytes to give a higher mortality after irradiating fertilized eggs. The conclusion that dominant lethals in oocytes and eggs are mostly due to non-chromosomal damage would appear to be in disagreement with the results of Ulrich (DIS 28) who found a drastic difference in LD50 after irradiating the anterior half (containing the nucleus) and the posterior half of the egg, and concluded that the lethality was almost entirely due to nuclear damage. An explanation which would fit both sets of results is that the damage is largely due to injury to some cytoplasmic constituent or constituents, but that these substances are concentrated in the anterior region of the egg during the perifertilization period.

Valencia, Ruby M. Sex ratio after irradiating fertilized eggs.

Fertilized eggs were irradiated with 500r of X-rays (150 kV, 10 mA, 1 mm. Al filter) within 15 minutes after fertilization, presumably well before union of the pronuclei, for the purpose of observing dominant and recessive lethal damage. All adults hatching from the irradiated eggs were counted in order to calculate postembryonic mortality. We took advantage of this situation and counted the various classes of progeny separately, in order to determine whether or not there was an effect on the sex ratio. The crosses from which the eggs were derived were y scS1 B In49 v/y oc ptg females mated with y oc ptg males or with y scS1 B In49 v males. All males hatching from these crosses obviously carry an X chromosome irradiated in the maternal nucleus and a Y chromosome irradiated in the paternal nucleus. The females would carry one X irradiated in the maternal nucleus and one X irradiated in the paternal nucleus. Recessive lethals induced in the maternal X would be expected to result in a lowering of the frequency of males hatching. The results were as follows:

<table>
<thead>
<tr>
<th>Type male used</th>
<th>Females</th>
<th>Males</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>y B</td>
<td>y oc ptg</td>
</tr>
<tr>
<td>y oc ptg</td>
<td>872</td>
<td>822</td>
</tr>
<tr>
<td>y scS1 B In49 v</td>
<td>1567</td>
<td>---</td>
</tr>
<tr>
<td>Total</td>
<td>2439</td>
<td>822</td>
</tr>
<tr>
<td>y oc ptg</td>
<td>1254</td>
<td>1120</td>
</tr>
<tr>
<td>y scS1 B In49 v</td>
<td>1178</td>
<td>---</td>
</tr>
<tr>
<td>Total</td>
<td>2432</td>
<td>1120</td>
</tr>
</tbody>
</table>

A recessive lethal test carried out with the y B females hatching from the irradiated eggs showed that 4.6% of them carried a lethal in the X chromosome which was irradiated in the maternal nucleus. If we assume that an equal number of X chromosomes entered into male zygotes as entered into female zygotes, and that these X chromosomes carried an equal number of recessive lethals, we would expect 4.6% of them to have been eliminated. The number of males expected to have hatched would be 4256, very close to the actual number counted (4266). The observed number represents a reduction of 4.4% from the supposed original number of male zygotes (4461). The controls show no reduction of males, but rather a slight advantage over the females.

It is possible, however, that this apparently good agreement with expectation is spurious. It can be seen that the three classes of females and two classes of males have very different viabilities. It would be preferable to have results similar to these for irradiated isogenic wild type eggs.

The experiment is divided into four groups, each containing at least 60 males. The males were taken from a wild type Canton-S stock, which were stored without females until they were three days old. The males were then given a temperature treatment for half an hour. Immediately after the treatment, each male was mated for a period of 24 hours to five virgin three day old females hybrid for the gene markers, on bw; e".

At the end of the mating period, the females were transferred to black food for egg laying. The males were given a new set of five virgin hybrid females for another 24 hour mating period. This continued over a period of 14 days. In the control group, Group I, the males were stored at a temperature of 22°C. In the experimental groups, Groups II, III and IV, the males were treated for half an hour at a temperature of respectively 70°C, 40°C, and 0°C.

The data given in the table picture the frequency of fertile males on the different days after treatment. As it can be seen from the table, there is no apparent difference between the males treated at 22°C and 70°C. However, for the two other groups of males there is a marked reduction in the frequency of fertile males starting already on the second day after treatment. Thus, the data indicate that temperatures below 70°C induce male sterility.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Temp.</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
<th>13</th>
<th>14 days</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>22°C</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>98</td>
<td>98</td>
<td>96</td>
<td>93</td>
<td>86</td>
<td>86</td>
<td>71</td>
<td>64</td>
<td>53</td>
<td>46</td>
<td>40</td>
</tr>
<tr>
<td>II</td>
<td>70°C</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>95</td>
<td>95</td>
<td>95</td>
<td>78</td>
<td>70</td>
<td>70</td>
<td>56</td>
<td>56</td>
<td>56</td>
<td>50</td>
<td>45</td>
</tr>
<tr>
<td>III</td>
<td>40°C</td>
<td>100</td>
<td>73</td>
<td>51</td>
<td>39</td>
<td>31</td>
<td>26</td>
<td>19</td>
<td>19</td>
<td>12</td>
<td>10</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>IV</td>
<td>0°C</td>
<td>100</td>
<td>50</td>
<td>23</td>
<td>12</td>
<td>7</td>
<td>5</td>
<td>5</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Zürcher, C. Balanced polymorphism and heterosis in crosses of wild type and the mutant eug of D. melanogaster.

Mass-breeding populations of the constitution eug/eug/ + were established at 20°C, 25°C, and 28°C, and the frequencies of the eug allele were determined at intervals of 2-4 weeks. The experiments at 25°C and 28°C were run for 8 months, and those at 20°C, for 4 months. After the first 2 months a constant value of 35 - 40% was obtained for the frequency of the mutant allele. The temperature seems to have no influence on the process of selection under the conditions used in the laboratory. In separate breeding experiments using eug/eug, eug/ + and +/+, it was established that the heterozygotes show a heterotic effect in reference to longevity and resistance to desiccation. Further analyses of viability factors are in progress.
Bender, H. A. Dioxane dehydration of Drosophila tissue. The substitution of dioxane, $0(CH_2CH_2)_2O$, for ethanol in the dehydration of the delicate tissues of Drosophila has proved very successful in reducing shrinkage, and in paraffin embedded material, of decreasing splitting during sectioning. Dioxane is compatible with most fixing agents (including osmic fixation), the majority of stains and the Feulgen technique. Tissues should be fixed and washed as usual, placed in dioxane (with two changes) for at least six hours (usually overnight) and subsequently handled as if in absolute alcohol. (Details of the dioxane method may be found in a critique by Mossman, Stain Technol. 12, 4, October, 1937.)

David, J. A new medium for rearing Drosophila in axenic conditions. The live yeast medium used by many authors in Drosophila studies is often unsuitable for the exact determination of quantitative characters. Different micro-organisms may develop on this medium and, if so, are always the source of important and uncontrolled variations. In contrast, an aseptic, chemically defined medium is very difficult to use when large number of flies are to be studied. Therefore, another medium, more convenient for quantitative research, has been worked out. This new medium is easy to prepare because not chemically defined and easy to use because aseptic conditions are assured in a simple manner by an antiseptic. The composition is the following:

- agar 15 g
- dried brewer's yeast 96 g (dry weight)
- corn meal 96 g (dry weight)
- antiseptic (nipagin or tegosept in alcoholic solution) 6 g
- water: to a total weight of 1200 g

Agar is dissolved in boiling water. The other ingredients are then added, and the medium is autoclaved for 15 minutes at 115 °C. When autoclaved, the medium is supplemented with water up to a total weight of 1200 g, then mixed and poured in culture tubes.

With this diet, larval development and adult fecundity are as good as those observed with the live yeast medium, and accidental variability is much lower. The new medium has been used for seven years in our laboratory and has proved very satisfactory for genetic and physiological studies in Drosophila. More detailed information is available in the following publications:


Hildreth, P. E., and Cole Brunt. Method for collecting large numbers of fertilized D. melanogaster eggs in meiotic stages. The method to be described permits the transfer of females from one egg collection dish to another with no loss of females and with a minimum of agitation to the females, thus disturbing the egg-laying pattern very little. Although we have used the method for collecting large numbers of eggs it would also be satisfactory for collections from single females.
Two days before the eggs are needed, virgin females and males are collected and stored separately with approximately 40 flies in each 1/2-pint culture bottle which contains yeasted standard cornmeal-agar-molasses medium. Also on this day, 600 cc of the same type of culture medium should be mixed with 10 g of live Baker's yeast and allowed to ferment for two days. In the morning, two days after having been collected, males from each bottle are shaken (without etherization) into the bottles, giving about 40 pairs per bottle. After about three hours the flies are shaken, without etherization, into tubes used as egg-laying chambers (one bottle of flies per tube) and the tubes are placed immediately on blotting paper for collection of eggs.

The tubes are clear plastic, about 45 mm long and 22 mm in diameter. One end of the tube is covered with a single layer of dacron gauze (through which the females oviposit) and the other end is plugged with cotton after the flies have been shaken in. The cotton should be pushed down to within about 1/2 cm from the dacron gauze to keep the flies near the food. Dacron gauze is used because the fibers do not absorb moisture and do not shrink or expand with moisture changes. The gauze may be held on the tube with rubber bands or the gauze may be glued to or embedded in the plastic with the proper solvent. We have found the latter method to be most satisfactory.

The collection dish consists of a Petri dish lid or base which has been filled with the previously mentioned fermented culture medium. On the surface of this food is placed a Kimwipe or Kleenex-type tissue on which the blotting paper will be placed. Dark green blotting paper cut into rectangles about three inches by four inches is found to be satisfactory for the egg collection as the eggs are readily visible against the dark background. The blotting paper is first soaked in a vinegar solution (nine parts water to one part commercial white vinegar) before it is placed on the collection dish. Both the fermented food and the vinegar solution are necessary to stimulate rapid egg laying. At the end of the egg-laying interval the tubes may be gently lifted, the blotting paper with the eggs removed, a new piece of blotting paper placed on the fermented food, and the tubes placed on the fresh paper for another collection. The eggs may be treated while on the paper, or removed easily with a needle or small brush, or the paper and eggs may be inserted into a bottle containing culture medium and permitted to develop.

During collecting intervals of five minutes we have occasionally been able to collect over 200 eggs (using nine tubes), and average about 100 eggs. In our experiments we normally make 30 to 40 such collections in an afternoon. In a sample of 190 eggs collected in this manner and then prepared with Feulgen's stain it was observed that slightly more than 75% of the eggs were in meiotic stages. Fixation in some cases did not occur until about 20 minutes after the eggs were laid, so the percentage of eggs in meiotic stages at the end of the five-minute collection period would be higher than is indicated. If small quantities of very young eggs are desired it would be best to use fewer flies in each tube and then to select those tubes in which the eggs are being laid rapidly.

(This work was done under the auspices of the U. S. Atomic Energy Commission.)

Kirschbaum, W. F., and Ruby M. Valencia. Modified egg-collecting technique. We are using a modification of the egg-collecting technique published by Ulrich (DIS 27). Since for our purpose we need not manipulate the eggs, we considered it worth while to avoid any manipulation, thus saving time and perhaps avoiding possible damage to the eggs. When we wish to collect eggs from the flies in the bell jar apparatus, we substitute for the food plate a plate containing a block of wood (to fill the space usually occupied by medium) on which is placed a round piece of thin white blotting paper (diameter 5.6 cm.), wetted in vinegar and spread with a thin film of yeast suspended in vinegar. The paper is previously scored with pencil in rectangles (7.5 x 4.5 mm.) calculated to fit the field of the dissecting microscope. After placing the paper
on the wood block, it is scraped sharply between the pencil lines, in the longitudi-
nal direction of the rectangles. The flies like the rough surface to lay their 
Eggs and lay most of them there, thus facilitating the counting later. The papers 
are simply removed from the plates, iradiated and placed in regular bottles con-
taining medium. The larvae hatching from the eggs very soon crawl down into the 
medium, since they lack sufficient food on the paper. The papers are removed from 
the bottles at 48 hours and the eggs counted. (If time does not permit counting 
at once, the papers may be conveniently stored in petri plates in the refrigerator 
for several days.) For counting, the papers are moistened from the lower surface, 
placed on a clear glass plate and examined with a stereo-microscope, using trans-
mittted light and a magnification of 25X. Hatched and non-hatched eggs are easily 
distinguished.

Levitan, M. Long-distance Drosophila collecting. In collecting particular species of Drosophila 
at some distance from home territory, it is imperative to obtain the largest possible 
sample in the shortest possible time. The problem is particularly acute when seeking 
a species such as D. robusta which is rarely the majority among the flies coming to 
bana traps. Over the past several years I have found that the following technique 
modifications usually increase the efficiency and ease of long-distance collections:

**Bait:** About 5-7 days before they are to be used, very rotten bananas, the 
kind grocery-men want to throw away, are mashed, the cut-up peels added, and placed 
in a bucket which has ice-pick size holes punched in the bottom. At a woods the 
first bucket is hung alone from a branch and the contents allowed to ferment from 
( or absorb) wild yeasts and bacteria, water of fermentation dripping out. When 
traveling this bucket is set into another one so that the fermentation water leaks 
into the one with good bottom. D. robusta and certain other species appear to be 
selectively attracted to bait that is fermented in this way and is not too wet; 
even quite dry bait will attract them better than most other species.

**Traps:** Since D. robusta is most efficiently collected from traps that are 
left in the woods several days, I use quart Mason jars tied to the trees with wire. 
Stove-pipe (for example Sears Roebuck catalog #9H9903, 10 cents for a 50 ft.), 
galvanized, or similar cheap pliable wire is good for this purpose. I attach the 
wire permanently, tightening it around the neck of the jar with pliers. Rain is 
kept out by making the wire long enough so that it can be knotted several times a 
few inches beyond the jar and then passed through the mid-point of a paper plate, 
the plate resting on the knotted portion. Plasticized paper plates are excellent 
for this, especially the ones that present a blue or brown surface to the open mouth 
of the jar. For transport the paper plate can be pulled off, the wire wrapped around 
the jar, and the jars with the bait in them covered by regular jar caps. Thus they 
are ready to hang quickly in the next locality.

Mahowald, A. P. Fixation problems for electron microscopy of 
Drosophila embryos. An electron microscopic study of early 
embryogenesis in Drosophila is nearly 
completed and it seems appropriate at this 
time to pass on information on the fixation 
techniques used. The vitelline membrane is impermeable to the usual fixatives 
used. After trying various chemical and enzymatic methods for attacking the membrane, 
it was found that very brief treatment with ether (1-2 seconds) or slightly longer 
treatments with other organic solvents with lower water affinities, e.g. toluene or 
isopentane, rendered the membrane readily permeable to either OsO₄ or KMnO₄. The 
ether-extracted compound is probably a wax since this substance is frequently the 
water-proofing substance in insects. In electron micrographs a layer about 500Å 
in width was found between the vitelline membrane and the chorion; it is probably 
this layer that was removed by the solvents. Examination with the electron microscope
showed that embryos treated with ether for as little as one second were seriously injured. Consequently further efforts along this line were stopped and the usual micro-puncture with a fine tungsten needle was used.

Concerning the puncture, a compromise must be achieved between one large enough to allow the fixative to penetrate sufficiently fast and one so large that distortion of the embryo results. This latter difficulty is especially common at the early stages studied, i.e. the blastoderm formation and pre-blastoderm stages. In our studies the chorion was routinely removed with NaOCl so that the specific stage desired could be picked out. The removal of the chorion has the added advantage that the puncture can be made more delicately. However, the chorion is no hindrance to the penetration of the fixative, so this is not a necessary step. As soon as the initial fixation has occurred around the puncture wound, the hole should be enlarged. Experience has shown that both OsO₄ and KMnO₄ render the vitelline membrane brittle; consequently care must be taken that excessive pressure is not used which would result in splitting of the membrane. This fact, however, becomes useful for the next procedure. After most of the embryo has become colored with the fixative, it is helpful if the vitelline membrane is dissected off (this is not possible after permanganate fixation for reasons not known). This last difficult step is not necessary for good fixation if the initial penetration of the fixative was rapid enough, but it has other advantages. Bahr et al. (Exp. Cell Research 12:342-355, 1957) have shown that tissues undergo a 15-20% expansion during short fixations and that during subsequent dehydration with alcohol there is an equivalent shrinkage. Because of the vitelline membrane, however, the expansion in the fixative does not occur. Consequently, after the contraction in the alcoholic series, the embryo is 20% smaller than originally. This shrinkage results in increased cytoplasmic density, thus necessitating very thin sections in order to discern the fine structure. If the vitelline membrane is removed within the first 30 minutes of fixation, an expansion of about 15% still occurs. As a result the fine structure is more easily observed and it can be more readily compared to other tissues.

A second advantage is that infiltration with plastics is more uniform. If the vitelline membrane is not removed, the embryo should be cut in two in 95% alcohol in order to facilitate infiltration. This is imperative with the epoxy resins and is necessary for consistent results with methacrylate.

Because of the dangerous fumes of OsO₄, it must be noted that good ventilation is required during these procedures. During the actual operations, especially if groups of embryos are being worked on, a small fan has been successfully used to prevent the accumulation of fumes at the dissecting scope.

No other modifications of standard preparatory procedures were found necessary.

**Mittler, S., and J. Bennett.**

A simple food medium that requires no live yeast with the minimum of variables.

The medium to be described had been developed by Dr. J. Crow at the University of Wisconsin, and has been used for several years in our laboratories with much success. There is no need to add or maintain live yeast whose growth forms a moist sticky layer over the surface of the food which can trap flies. The highly variable molasses and corn meal of the "standard media" has been eliminated. The formation of harmful excess carbon dioxide by the live yeast and movement of media by the gas has been eliminated.

**Food Formula**

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>Amounts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>1000 ml</td>
</tr>
<tr>
<td>Agar</td>
<td>19 gm</td>
</tr>
<tr>
<td>Sucrose</td>
<td>54 gm</td>
</tr>
<tr>
<td>Dried yeast</td>
<td>32 gm</td>
</tr>
<tr>
<td>Propionic acid</td>
<td>5 ml</td>
</tr>
</tbody>
</table>
The agar, sucrose, and dried yeast are added to the warm water and mixed while heated until thoroughly dispersed. The mixture is boiled for 5 to 15 minutes to kill all the yeast (it may be necessary to add 200 ml of additional water at the start to allow for the amount boiled away). The propionic acid is added after the mixture has cooled to 60°C. We use Schlitz Brewing Co.'s "non-debittered dried brewers yeast," obtainable in 100 lb. bags. The large quantity assures long term control of quality of this ingredient.

Sang, James H. A simple method of adding solutions to axenic cultures. The problem of adding measured amounts of nutrients (or of mutagens or other substances) to germ-free cultures of Drosophila larvae at particular ages is simply solved by the device illustrated. Bulb tubes are made by first drawing out cleaned tubing and then blowing a bulb so that a sufficient length of the drawn tube remains to permit its easy fracture by gentle pressure against the side of the culture tube. The bulb is filled by graduated syringes, and stoppered with cotton-wool. Prepared cultures can then be autoclaved and handled in the normal way (Sang, 1956, J. Exper. Biol., 33:45). Alternatively sterile solutions may be introduced into the bulb after autoclaving when this would damage the required additive. The nutrient is added to the medium at the desired time, by breaking the drawn tube against the side of the culture.

Moree, Ray. Simple demonstration of modified ratios using b and e. Laboratory experiments relating to modified ratios and genic interaction can be made both simple and surprising by using stocks of b and e. That the F₁ is wild type is surprising to many. The F₂ is classified by most students into wild type and "dark," in 9:7 ratio if the sample size is large enough. But some students detect what they consider as different degrees of darkness, so the possibilities of getting a 9:3:4 or a 9:6:1 ratio are pointed out: it can also be indicated that if a simple chemical test were available it might even be possible to recognize a 9:3:3:1 ratio. That the ratio of wild to dark may be about 1:1 in F₂ progeny and 1:3 in testcross progeny from the cross F₁ x b/b; e/e, is usually somewhat surprising, too. The results emphasize the way in which inferences as to interaction, epistasis, etc., depend upon the possibilities of discriminating among the individual progeny of a cross.
J. A. Beardmore has moved from the Genetics Department, University of Sheffield, to the Genetic Institute, State University of Groningen, Netherlands.

Professor Hampton L. Carson has recently returned to Washington University, St. Louis, after a period of nine months in Australia as a Senior Fulbright Research Scholar. Most of this time was spent in the Zoology Department at the University of Melbourne; from here, Professor Carson visited other parts of Australia, also New Guinea.

James Divelbiss has moved from the Department of Zoology at the University of Iowa to the Department of Biology, Westmar College, in Le Mars, Iowa, where he is an assistant professor.

Marvin Druger, formerly of the Genetics Group of Columbia University, New York, is visiting the Animal Genetics Laboratory, Sydney University, for one year from mid-August, 1961. Dr. Druger holds a post-doctoral fellowship from the National Institutes of Health and is working on problems of canalization.

Dr. A. C. Fabergé of the Department of Zoology, University of Texas, is visiting the Genetics Laboratory, Biology Department, University of Oregon, until September, 1962.

Lawrence D. Friedman, formerly at the Department of Medical Genetics at the University of Wisconsin, is now Assistant Professor, Department of Biology, Hiram College, Hiram, Ohio.

George D. Hanks has joined the staff of the Department of Genetics, University of Utah, Salt Lake City, Utah.

Jerry Hirsch has been appointed Associate Professor of Psychology in the Department of Psychology, University of Illinois, Urbana, where he will supervise a Ph.D. program in, teach courses in, and direct research on behavior genetics (the major emphasis in the laboratory remains on Drosophila). Reprints of, or references to, studies of heredity and behavior (in all species) will be greatly appreciated.

Benjamin Hochman wishes to express his appreciation to those who supplied him with stocks and trapped flies following the loss of his experimental lines and stocks in October, 1961.

M. E. Jacobs is now located at the Biology Department of Eastern Mennonite College in Harrisonburg, Virginia. He was formerly of Bethany College in Bethany, West Virginia.

Edward C. Keller, Jr., has recently moved to the Genetics Laboratory in the Department of Biochemistry and Nutrition, the University of North Carolina, Chapel Hill, North Carolina.

Professor R. C. Lewontin of the Biology Department, Rochester University, New York, is visiting the Animal Genetics Laboratory, Sydney University, for one year from mid-June, 1961. Professor Lewontin holds a Fulbright Scholarship and is working on problems of canalization and population genetics.

Benedetto Nicoletti is organizing a Drosophila Laboratory in the new Genetics Department, University of Rome, Rome, Italy. He shall be very grateful to the friends who can send him their old reprints or put his name in their mailing list.

T. M. Rizki, formerly in the Biology Department at Reed College, has joined the Department of Zoology at the University of Michigan as Associate Professor.
B. Sakaguchi has moved on the end of August from Dr. Donald F. Poulson's laboratory at Yale University in New Haven to the National Institute of Genetics, Misima, Japan. He is continuing his work on maternal inheritance of "sex-ratio" condition in Drosophila.

L. Sandler will, in June of 1962, move from the Genetics Department, University of Wisconsin, Madison, Wisconsin, to the Genetics Department, University of Washington, Seattle, Washington.

K. C. Sondhi, formerly at the Department of Zoology, University College, London, has been appointed Geneticist at the New England Institute for Medical Research, Ridgefield, Connecticut.

Oistein Strømmane is on leave of absence from the Institute of Genetics, University of Oslo. He is staying as a research associate at the Department of Botany, University of Chicago, through 1962 until May, 1963.

Victor E. Tinderholt. The Department of Genetics, City of Hope Medical Center, and the Department of Zoology, U. C. L. A., report the sad news of the death of Victor E. Tinderholt. His great courage and ability to enjoy the world about him in the face of grave illness will be long remembered by those who knew this intelligent, sensitive person.

Yasuko Toyofuku (Mrs. Tonomura) was appointed a research member of the National Institute of Genetics at Misima on March 15, 1961.

Heinrich Ursprung has been appointed to the Faculty of the Department of Biology, The Johns Hopkins University, Baltimore 18, Maryland, as an Assistant Professor, effective July 1, 1962.

Dr. Marvin Wasserman, from the University of Texas, returns to the United States in 1962. Dr. Wasserman has spent two years in Melbourne as a member of the teaching staff. During this time he has pursued his studies on the repleta group of the genus Drosophila, visiting many regions in Australia and New Guinea.

Yukio Yamada, National Institute of Genetics, Misima, Japan, has joined the Population Genetics Institute, Purdue University, as a visiting research professor. He is especially interested in genotype by environment studies with Drosophila and Tribolium.

MATERIALS REQUESTED OR AVAILABLE

J. A. Beardmore (Genetical Institute, Haren (gr), Netherlands) would like to hear from anyone having stocks of any species of Drosophila showing a morphological polymorphism or knowledge of the occurrence of such polymorphism in natural populations.

J. L. Blount (Department of Biology, Mt. Union College, Alliance, Ohio) would be grateful for wild-type strains of D. melanogaster whose adult longevity is known or suspected to be of either unusually short or long duration.

B. Burnet and J. H. Sang are studying the factors which alter penetrance and expressivity of eyeless. They would be grateful for a stock of ey^D39k which they have been unable to trace, or for any information about this stock which was last reported on by Hinton, 1942, Amer. Nat., 76:219-23.
F. Mainx (Institut f. Allgemeine Biologie, University of Vienna, Wien IX. Schwarzspanierstr. 17) would appreciate obtaining strains of Megaselia scalaris (=Aphiochaeta xanthina) from different places as well as strains of other species of Phoridae easily bred in the laboratory.

George A. Marzluf would appreciate receiving any stocks containing suppressors of vermilion, purple, and black. His address is: Department of Biology, The Johns Hopkins University, Baltimore, Maryland.

Max Planck-Institut für Biologie, Abteilung Beermann (Tübingen, Germany, Spemannstr. 34) would be grateful to obtain: 1) D. nigrohydei; 2) any mutation of D. hydei; 3) any Drosophila species that can be crossed with D. hydei giving either fertile or sterile hybrids.

R. D. Milkman (Department of Zoology, Syracuse University). If anyone finds it desirable to assign a small selection problem to a student, I should like very much to have any true-breeding polygenic crossveinless strains that may be obtained. It should be possible to obtain such a strain by selection of the progeny of even a few wild flies. This has proven easy in the past.

Dr. Yasuhiro Miyoshi would like to have wild strains from various localities in the United States for studies on tolerance of certain salt concentrations. His address is: Department of Zoology, Faculty of Science, Kyoto University, Kyoto, Japan.

QUOTABILITY OF NOTES

Angus, D. 35:71.
Arnold, L. 32:166.
Barish, N. 28:103.
Baumiller, R. 32:113; 33:122.
Bochnig, V. 26:91; 28:108.
Doane, W. W. 32:121; 34:49 (cf Doane 35:45b), 35:45a; 35:78.
Frost, J. N. 35:81a; 35:81b.
Fuscaldo, K. E. 35:84.
Hannah, A., and Ø. Strommaes. 29:121.
Harrison, B. J. 17:60; 28:122a; 28:122b; 28:123.
Jacobs, M. E. 29:126; 31:124; 32:130a; 32:130b; 32:130c; 33:140; 35:89.
Lüers, Th. 28:131; 30:132; 30:133.
Mather, Wharton E. 27:101; 33:147.
Mead, C. G. 35:89.
Oksala, T. A. 31:147; 31:149.
Röhrborn, G. 30:148; 33:156.
Sandler, I. 30:151; 32:154.
Stevenson, R. 33:182.
Strommaes, Ø., and A. Hannah. 29:179.
Telfer, J. D. 28:161.
Volkart, H. D. 33:100.
Ursprung, H. 33:174; 34:110.
The Genetics Training Committee of the University of North Carolina wishes
to announce the availability of the pre- and postdoctoral traineeships for the
study of Drosophila or Medical Genetics. Persons interested should write to
Professor John Graham in care of that institution in Chapel Hill, North Carolina.

King, R. C. A suggestion with respect to translations.
Might it not be useful to have a yearly listing of English translations
of foreign language works dealing with Drosophila and to have duplicate transla-
tions collected in one laboratory (Herskowitz's at St. Louis University, for
example), where they could be made available to everyone? Such a system might
save a great deal of duplicated effort. Each translation should be OKed by the
original author before its release.

Novitski, E., and R. Dorsey. A generalized maximum likelihood program for
the IBM 1620.
We are now programming the IBM 1620 to handle maximum likelihood problems
of the sort that might concern geneticists. The procedure followed will be the
method of least scores as described by Rao, making it possible to solve relatively
complex formulations by iteration. Taking the necessary partial derivatives, and
forming the matrices, transpose and inverse, will be done internally by the program.
Provision will be made to detect insoluble or ambiguous formulations. It would be
helpful to us if anyone with a bona fide likelihood problem at the present time
would let us know its nature so that we might check our concept of what such a
program should be like against the demands of actual cases.

Sokoloff, A. Transfer of Tribolium stocks.
The stocks of Tribolium castaneum, Tribolium confusum and Latheticus oryzae
have been transferred from the Biological Laboratory, Cold Spring Harbor, to the
Department of Genetics, University of California, Berkeley. Several wild type
strains for T. castaneum and T. confusum and one for L. oryzae are being maintained.
In addition a large number of stocks with sex-linked and/or autosomal markers is
available, particularly for T. castaneum. Supplies of some of the mutant and wild
type stocks are available to those who intend to use them in their genetics courses.

Supported by a National Science Foundation Grant, a conference on Behavior
Genetics was held at the Center for Advanced Study of the Behavioral Sciences in
Stanford, California, from August 14 through September 3, 1961. The organizing
committee consisted of Jerry Hirsch (Psychology, University of Illinois), chairman;
Gerald E. McClearn (Psychology, University of California), Benson Ginsburg (Biology,
University of Chicago), Howard Hunt (Psychology, University of Chicago).
The other members of the conference were Gordon Allen (Genetics, National
Institute of Health), Peter L. Broadhurst (Psychiatry, University of London, England),
Jan. H. Bruell (Psychology, Western Reserve University), Ernst W. Caspari (Biology,
University of Rochester), Eckhard Hess (Psychology, University of Chicago),
John A. King (Zoology, Michigan State University), Daniel S. Lehrman (Animal
Behavior, Rutgers University), Gardner Lindzey (Psychology, University of Minnesota),
Aubrey Manning (Zoology, University of Edinburgh, Scotland), Robert C. Roberts
(Animal Genetics, University of Edinburgh, Scotland), and W. Robert Thompson
(Psychology, Wesleyan University).
Guests at some sessions included Sherwood Washburn (Anthropology, University of California), James McGaugh (Psychology, San Jose State College), Mark Rosenzweig (Psychology, University of California), J. Anthony Deutsch (Psychiatry and Psychology, Stanford University), Kenneth Calby (Psychiatry, Center for Advanced Study of the Behavioral Sciences), Leon Otis (Stanford Research Institute), John Clausen (Sociology, University of California), William Meredith (Psychology, University of California), Frank A. Beach (Psychology, University of California), Francis Palmer (Social Science Research Council), and Louise Erlenmeyer-Kimling (Medical Genetics, Psychiatric Institute, Columbia University).

In 1962 a second and final meeting of the conference will be held to complete a volume on Behavior Genetics. It will consist of chapters that were stimulated by last summer's discussions.

At the September, 1961, meeting of the Social Science Research Council, a committee for genetics and social behavior was established. The members of this committee are: Gardner Lindzey (Psychology, University of Minnesota), chairman; Ernst Caspari (Biology, University of Rochester), Theodosius Dobzhansky (Zoology, Columbia University), David Hamburg (Psychiatry, Stanford University), Jerry Hirsch (Psychology, University of Illinois), Gerald McClearn (Psychology, University of California), James Spuhler (Anthropology, University of Michigan).

The expressed purpose and functions of the new committee are "to facilitate and expedite research in Behavior Genetics in whatever manner seems appropriate with particular reference to the application of new knowledge and advanced methods and techniques to the study of human behavior."

DIRECTORY

Geographical

(Alphabetically arranged according to country, city, laboratory.)

ARGENTINA

Buenos Aires
Comision Nacional de Energia Atomica, Claustro de Investigaciones Cientificas, Laboratorio de Genetica

Kirschbaum, Werner F. Research Assistant.
Leon, Williams N. Technical Assistant.
de Marinic, Susana Ercolini. Research Assistant.
Paz, Bonifacia del Carmen. Curator of Stocks.

Buenos Aires
Universidad de Buenos Aires, Facultad de Ciencias Exactas y Naturales

Cachero, Nestor. Research Assistant.
Diez, Julio. Research Assistant.
Kaufman, Beatriz. Stocks Curator.
Mazar-Barnett, Beatriz. Student Investigator.
Valencia, Juan I. Professor. Head of Department of Biology.
AUSTRALIA

Adelaide, South Australia
University of Adelaide, Department of Genetics

Mayo, M. Jean. Ph.D. Lecturer.
Hayman, D. L. Ph.D. Lecturer.

Brisbane
The University of Queensland, Department of Zoology

Mather, Wharton B. Ph.D. Senior Lecturer. Population Genetics, Cytogenetics, Chromosomal Polymorphism.
Angus, D. B.Sc. (Hons.) Graduate Student. Population genetics.
Spurway, Rosalyne. Research Assistant.

Hobart, Tasmania
The University of Tasmania, Department of Zoology

Brink, N. G. Research student. Mutation.
Campbell, R. (Mrs.) Research assistant.
Clark, A. M. Professor. Radiation and chemical mutagenesis.
Clark, E. G. (Mrs.) Research assistant.
Knott, B. Technical assistant.

Melbourne, Victoria
The University of Melbourne, Department of Zoology

Gunson, Mary M. M.Sc. Lecturer. Salivaries.
Wasserman, M. Ph.D. Senior Lecturer. Cytology; evolution.

Sydney, New South Wales
Sydney University, CSIRO Animal Genetics Division, Animal Genetics Laboratory, Department of Zoology

Sheldon, B. L. B.Sc. Agr., Ph.D. Research Officer. Selection; induction of mutations.

Sydney, New South Wales
University of Sydney, Department of Animal Husbandry

Barker, J. S. F. Ph.D. Senior Lecturer. Population genetics.

AUSTRIA

Vienna IX (Wien IX)
Institut f. Allgemeine Biologie, Schwarzspanierstrasse 17

Mainz, Felix. M.D., Ph.D. Professor. Head of department.
Ruttner, Friedrich. M.D., Ph.D. Genetics of the honey bee.
BELGIUM

Louvain
The University, Agricultural Institute, Laboratory of General Genetics

See DIS 34:127.

BRAZIL

Pôrto Alegre
Universidade do Rio Grande do Sul, Departamento de Genética, Instituto de Ciências Naturais, Av. Paulo Gama.


Cordeiro, E. R. Technician.

Bitadi, T. F. (Miss) Technician. Genetic analysis of irradiated populations.


Ludwig, Maria. Technician. Stockkeeper.

Ludwig, Nilda Conceição. Technician.


Alophia (Iridaceae): chromosomal polymorphism.


Ramila, D. Technician. Foodmaker.


Salzano, F. M. Ph.D. Assistant Professor. Head of the Human Genetics Division of the Department. Human blood groups. Indian population genetics.

Santos, Alda T. D. Administrative Assistant.

Silva, Luiz C. Technician Electronics.

Silva, Tereza M. Technician. Stockkeeper.

Simões, G. V. Technician. Field worker. Human genetics.

Thedy, O. Technician.

Tondo, C. V. E.B. Bc.Sc. Head of Biophysical Division of the Department. Electrophoresis; chromatography in Drosophila mutants and human blood groups; development of new techniques.

Trogildo, D. N. Technician. Stockkeeper.


São Paulo
Universidade de São Paulo, Faculdade de Filosofia, Ciências e Letras, Departamento de Biologia Geral, Caixa Postal 8.105

Basile, R. Graduate Student. Cytogenetics.


Camba, C.A. Graduate Student. Speciation.

Prota-Pessoa, O. Ph.D. Assistant Professor. Human genetics.
Matos, N. S. Graduate Student. Population genetics.
Magalhães, L. E. de. Ph.D. Assistant Professor. Population genetics and speciation.
Pavan, C. Ph.D. Professor. Head of the Department. Population genetics, radiation genetics and cytogenetics.
Poletto, D. Graduate Student. Human genetics.
Wajntal, A. Graduate Student. Human genetics.

CANADA

Toronto
University of Toronto, Department of Zoology

Butler, L. Ph.D. Associate Professor. Director of the Laboratory. Population genetics.
Tallan, I. Ph.D. Assistant Professor. Genetics of antigens.
Seiger, M. B. M.A. Graduate Student. Quantitative inheritance.
Mileiko, V. V. B.A. Technical Assistant. Curator of Stocks.

Vancouver, B. C.
The University of British Columbia, Department of Biology and Botany

Cole, Kathleen M. M.A., Ph.D. Assistant Professor. Mutations and cytogenetics.

Vancouver, B. C.
The University of British Columbia, Department of Zoology

Band, Henretta T. (Mrs.) Ph.D. Research Associate. Population genetics
Ogawa, Tomoye. Technical Assistant
Tabata, Kazi. Technical Assistant

CHILE

Santiago
Universidad de Chile, Instituto de Biología "Juan Noé," Cátedra de Biología, Zañartu 1042

Brncic, D. Associate Professor. Population genetics.
Covarrubias, Edmundo. M.D. Research Assistant.
Pellcic, M. Dolores. Technical Assistant.

COLOMBIA

Bogotá
University of the Andes, Department of Biology, Apartado Aereo 4976

Hunter, Alice S. Ph.D. Physiology, taxonomy--Drosophila.
Newball, Sarah. Assistant.

Bogotá D. E.
University of the Andes, Department of Genetics

Cortés, Blanca Inés (Miss). Laboratory technician.
Díaz, Napoleón (Mr.). Laboratory technician.
Duplat, Hermán (Mr.). Laboratory technician.
Chejne, Abraham (Mr.). Laboratory technician.
Rios, Cesar (Mr.). Technician's help.

DENMARK

Copenhagen
University of Copenhagen, Institute of Genetics, 2A Øster Farimagsgade

See DIS 34:128.

FINLAND

Helsinki
University of Helsinki, Institute of Genetics, P. Rautatiekatu 13

Suomalainen, Esko. Ph.D. Professor. Head of Department.
Tiivola, Airi. (Mrs.) Technical assistant. Curator of Stocks.

Turku
University of Turku, Institute of Genetics

PROST, Justin N. Ph.D. M.I.H. Postdoctoral Fellow. Melanogaster: interchromosomal effects.
Hannah-Alava, Aloha (Mrs.). Ph.D. Research Associate. Melanogaster: developmental genetics; mutations.
Harmoinen, Liisa (Miss). Research Assistant. Melanogaster: mutations.
Heinonen, Pirkko (Miss). Research Assistant. Melanogaster: mutations.
Oksala, T. A. Ph.D. Professor. Head of Department. Melanogaster: mechanism of segregation; interchromosomal effects.
Savolainen, Salme (Mrs.) Technical Assistant.
Wallenius, Marja-Liisa (Miss). Research Assistant. Melanogaster: mechanism of segregation; interchromosomal effects.

FRANCE

Gif-sur-Yvette (S. et O.)
Centre National de la Recherche Scientifique, Laboratoire de Génétiqne evolutie et de Biométrie

BERGERARD, J. Professor. Cytogenetics.
BIGLER, J. (Miss) Technician.
BOSIGE, E. Ph.D. Chargé de recherches. Heterosis, sexual selection.
LANGÉ, G. (Miss) Assistant. Triploid intersexes of Drosophila.
LÉON, M. (Miss) Graduate student. Irradiation effects on development.
LOUIS, M. (Mrs.) Technician.
PIVA, A. Graduate student. Quantitative inheritance.
QUEIROZ, J. (Mrs.) Attachée de recherches. Quantitative inheritance.
TEISSIER, G. Professor. Head of the department. Population genetics, quantitative inheritance, biometry.

Gif-sur-Yvette (S. et O.)
Centre National de la Recherche Scientifique, Laboratoire de Génétique Formelle

LESTRAANGE, M.-Th. de (Miss). Attachée de recherches. CO₂ sensitivity in Drosophila.
L'HERITIER, Ph. Professor. Head of the Department. CO₂ sensitivity in Drosophila.
Ohanessian-Guillemain, A. (Mrs.) Chargée de recherches. CO₂ sensitivity in Drosophila.
Plus, N. (Mrs.) Chargée de recherches. CO₂ sensitivity in Drosophila.
Froust, J. (Mrs.) Attachée de recherches. Quantitative inheritance in Drosophila.
Vigier, Ph. Maître-assistant. CO₂ sensitivity in Drosophila.

Lyon (Rhône)
Laboratoire de Zoologie Expérimentale, Faculté des Sciences, 16, quai C. Bernard
Brun, J. Maître-Assistant. Cytology and genetics of nematodes.
Dalmon, J. Assistant. Nucleic acid metabolism.
Godej, J. (Mrs.) Assistant. Ovogenesis in Drosophila.
Guerrier, P. Cytology of nematodes.
Legay, J. M. Maître de conférences. Physiology and genetics of phytophagous insects.
Nigon, V. Professor. Head of the department. Nucleic acid metabolism.

Orsay (S. et O.)
Université de Paris, Faculté des Sciences, Biologie Générale
Bernard, J. (Miss) Assistante. CO₂ sensitivity in Drosophila.
Bregliano, J-C. Assistant. CO₂ sensitivity in Drosophila.
Brun, G. Chef de Travaux Pratiques. CO₂ sensitivity in Drosophila.

Paris
Faculté des Sciences, Laboratoire de Zoologie, 1 rue Victor Cousin, Paris 5 ème

See DIS 34:129.

Strasbourg (Bas-Rhin)
Université de Strasbourg, Faculté des Sciences

See DIS 34:129.

GERMANY

Berlin-Buch
Deutsche Akademie der Wissenschaften, Institut für experimentelle Krebsforschung,
Genetische Abteilung, Lindenberger Weg 70
Bender, Erhard. Dr. Microbial genetics: Chemical mutagenesis.
Geissler, Erhard. Dr. Head of Department. Microbial genetics: Lysogeny.
Pasternak, Luise. Melanogaster: Chemical mutagenesis.

Berlin-Dahlem
Institut für Genetik der Freien Universität Berlin, Rudeloffweg 9
Bellitz, Hans-Joachim (Dr.). Research Assistant. Melanogaster: induced mutations.
Bochnig, Veronika (Dr.). Research Assistant. Melanogaster: physiological genetics, radiation genetics.
Lüers, Herbert (Prof. Dr.). Director. Comparative genetics; mutagens.
Lüers, Thea (Mrs., Dr.). Guest Associate. Drosophila neurology.
Nöthel, Horst. Graduate student. Radiation genetics.
Ravasani, Chapour. Graduate student. Melanogaster: radiation genetics.
Röhrborn, Gunter (Dr.). Research Assistant. Drosophila tumors; chemical mutagens.
Struck, Eva (Mrs., Dr.). Research Assistant. Insects: cytology.
Wolf, Erich (Dr.). Associate. Insects: cytology.

Darmstadt
Botanisches Institut der Technischen Hochschule
Ziegler, Irmgard (Mrs., Dr.). Physiology of pteridines under the influence of genes.

Hamburg 13
Zoologisches Staatsinstitut und Zoologisches Museum, von-Melle-Park 10
Koske-Westphal, Thea (Mrs.). Ph.D. Study of hybrids between triploid melanogaster females and x-rayed simulans males.
Kosswig, Curt. Prof. Dr. Director.

Hamburg-Eppendorf
Universitäts-Frauenklinik, Strahlenbiologische Abteilung
See DIS 34:130.

Heidelberg
Universität Heidelberg, Zoologisches Institut, Sofienstr. 6
See DIS 34:130.

Karlsruhe
Institut für Strahlenbiologie, Kernforschungszentrum Karlsruhe
Apitzsch, Ursula. Curator of Stocks.
Catsch, Alexander. Prof. Dr. Drosophila genetics.
Dittrich, Wolfgang. Prof. Dr. Molecular genetics.
Hotz, Gerhart. Dr. Bacteriophage genetics.
Kiechelsen, Gerda. Dr. Drosophila genetics.
Müller, Adolf. Dr. Radiation biology.
Traut, Horst. Dr. Drosophila genetics.
Ufholz, Ilse. Technical Assistant.
Zimmer, Karl Günther. Prof. Dr. Radiation genetics.

Marburg/Lahn
Zoologisches Institut der Phillips-Universität, Ketzerbach 63
Becker, Gweneth L. Ph.D. Independent investigator. Lethals.
Scriba, Martin. Graduate student. Deficiencies and early embryology.

Mariensee über Wunstorf
Max-Planck-Institut für Tierzucht und Tierernährung
Gellert, Heidemarie. Technical Assistant.
Gottschewski, G. H. M. Prof. Dr. Head of department. Developmental and physiological genetics.
Querner, Waltraud. Dr. Assistant, Stockkeeper. Tissue culture.
Schwinck, Ilse. Dr. Guest investigator. Physiological genetics.
Zimmermann, Wolfgang. Dr. Assistant. Genetics.
Münster (Westf.)
Institut für Humangenetik der Universität Münster

Graebner, Erika. Technical Assistant.
Ostertag, Wolfram. Ph.D. Radiation genetics (somatic damage).

Tübingen
Max Planck-Institut für Biologie, Spemannstr. 34

Beermann, Wolfgang. Prof. Dr. Director. Physiology of salivary gland chromosomes.
Hess, Oswald. Dr. Research Assistant. Physiology of chromosomes (Y-chromosome).
Joneleit, Christa (Miss). Technical Assistant. Curator of stocks.
Meyer, Günther F. Dr. Research Assistant. Gametogenesis, light and electron microscopy.
Seidel, Sigrid (Miss). Graduate student. Sex determination (tra-mutation).

Ghana

Legon, Accra
University of Ghana, Department of Zoology

See DIS 34:131.

Legon, Accra
University of Ghana, Department of Chemistry

Blair, J. A. Ph.D. Lecturer. Origin of pteridine compounds in Drosophila.

Great Britain

Aberdeen, Scotland
University of Aberdeen, Department of Zoology

See DIS 34:131.

Bayfordbury, Hertford, Herts, England
John Innes Horticultural Institution

Harrison, B. S. Multiple insecticide resistance.

Birmingham 15, England
The University, Department of Genetics

See DIS 34:131.

Cambridge, England
University of Cambridge, Department of Genetics, Milton Road

Alderson, T. Ph.D. Research worker. Chemical mutagenesis.
Batten, J. L. Research student.
Gibson, J. B. Ph.D. Assistant in Research. Analysis of selected lines.
Nash, D. Research student. Developmental genetics.
Pelecanos, M. Research student. Chemical mutagenesis.
Spickett, S. G. Research student. Developmental genetics of quantitative characters.
Thoday, J. M. Ph.D. Professor. Selection, particularly disruptive. Location of polygenes.

Chalfont St. Giles, Bucks, England
Institute of Cancer Research: Royal Cancer Hospital, Pollards Wood

Fahmy, Myrtle J. Ph.D. Mutagenesis.
Fahmy, O. G. M.Sc., Ph.D. Cytogenetics.
Cleaves, C. Technical Assistant.
Hope, L. Technical Assistant.
Knight, E. B.Sc. Research Assistant.
Sweron, M. Technical Assistant.

Edinburgh 9, Scotland
Agricultural Research Council Poultry Research Centre, King's Buildings

Burnet, B. Ph.D. Physiological genetics.
Pratt, G. Stock keeper.
Sang, J. H. Ph.D., F. R. S. E. Drosophila nutrition and physiological genetics.
Strachan, I. Technical assistant.

Edinburgh 9, Scotland
University of Edinburgh, Institute of Animal Genetics

Auerbach, C. A. D.Sc., F. R. S. Reader, Chemical and induced mutagenesis.
Allan, J. Graduate student. Selection.
Basden, E. B. Research assistant. Wild species.
Clayton, G. Lecturer. Selection.
Kilsall, P. J. Graduate student. Spontaneous and induced nondisjunction.
Khishin, A. Guest investigator. Formaldehyde and radiation induced mutagenesis.
Knight, G. R. Research assistant. Subobscura salivaries.
Leigh, B. Graduate student. Chemical and induced mutagenesis. On leave with Professor Sobels.
Mostafa, A. Graduate student. Selection.
Nafei, H. Graduate student. Formaldehyde induced mutagenesis.
Osman, H. Graduate student. Selection.
Perry, M. Research assistant. Autoradiography.
Robertson, A. D.Sc. Quantitative genetics.
Robertson, F. W. D.Sc. Population and physiological genetics.
Royes, V. Graduate student. Drosophila nutrition.
Scharloo, W. Ph.D. Guest investigator. Quantitative inheritance.
Sen, B. K. Quantitative genetics.
Slizynska, H. (Mrs.) Ph.D. Cytological analysis.
Slizynski, B. M. Ph.D. Salivaries.
Snyder, L. A. Ph.D. Guest investigator. Chemical and induced mutagenesis.
Strachan, K. (Miss) Stock-keeper.

Glasgow, Scotland
University of Glasgow, Department of Genetics

Pontecorvo, G. F. R. S. Professor.
Forbes, E. C. Chief Technician.
Dorn, G. L. Ph.D. Research Associate.

Harwell, Didcot, Berks, England
Medical Research Council, Radiobiological Research Unit

Gale, C. Technical Assistant.
Jempson, J. Technical Assistant.
Lamerton, M. Technical Assistant.
Purdon, C. E. Ph.D. Radiation Genetics.

London, E. C. 1, England
St. Bartholomew's Hospital Medical College, Department of Zoology and Comparative Anatomy

Hollingsworth, M. Ph.D. Lecturer. Inbreeding and infertility. Bristle patterns in intersexes.
London, W. C. 1, England
University College, Department of Biometry, Eugenics and Genetics
Grüneberg, H. Professor.

London, W. C. 1, England
University College, Department of Zoology
Clarke, Jean M. Research Assistant. Ageing in Drosophila.
Lamb, Marion J. Research student. Radiation and ageing in Drosophila.
Maynard Smith, J. Lecturer. Genetics of pattern formation.

Manchester, England
Christie Hospital and Holt Radium Institute, Cytogenetics Department
See DIS 34:133.

Manchester 13, England
The University, Departments of Botany and Zoology
Hartshorne, John N. Lecturer in Genetics.

Nottingham, England
The University, School of Agriculture, Department of Agricultural Science
See DIS 34:133.

Sheffield 10, England
The University, Department of Genetics
Boam, T. B. Chief Technician. Stockkeeper.
Roper, J. A. Professor. Microbial Genetics.

INDIA
Calcutta 19
Calcutta University, Department of Zoology, Cytogenetics Laboratory
See DIS 34:133.

Calcutta 35
Indian Statistical Institute, 203, Barrackpore Trunk Road
See DIS 34:133.

Hyderabad
Osmania University, Radiation Genetics Project, aided by Department of Atomic Energy
(Government of India)
Reddi, O. S. Dr. Investigator.
Mathew, C. (Mr.) Senior Scientific Assistant.
Prabhakara Rao (Mr.). Junior Scientific Assistant.

Research problems under investigation:
1. Induction of translocations in the spermatogonia of Drosophila melanogaster by CB 1506.
2. Studies on the specific effect of phenylalanine mustard on the II chromosomal lethals of Drosophila melanogaster.
3. Induction of translocations in the spermatogonia of Drosophila melanogaster by X-rays.
Madras 7
Veterinary College, Department of Animal Genetics, Vepery


New Delhi 12
Institute of Agricultural Research Statistics (I. C. A. R.), Library Avenue

Marain, Prem. Professor.

Israel

Jerusalem
Hebrew University, Department of Zoology

Barak, Elisheva. Research student. Induced chromosome breaks.
Blum, Sonya. Laboratory Assistant.
Fattal, S. Laboratory Assistant.
Falk, R. Ph.D. Instructor. Induced mutations: viability effects and mechanisms.
Friedlander, M. Research student. Cytogenetics.
Goldschmidt, Elisabeth. Ph.D. Associate Professor. Pteridines.
Hinell, Nechama. Laboratory Assistant. Chromosome breakage.
Rappaport, Sarah. Research student.
Wahrman, J. Ph.D. Lecturer. Cytogenetics.

Italy

Milano
Università di Milano, Istituto di Genetica, Via Celoria 10

Di Pasquale, A. D.Sc. Assistant. Genetics of "brown spots."
Galucci, E. M.D. Research Fellow. Induced mutations in Drosophila.
Giavelli, S. M.D. Research Fellow. Induced mutations in Drosophila.
Locatelli, F. (Miss) Technician. Curator of Stocks.
Sironi, G. P. Student Assistant. Induced mutations in Drosophila.
Zambruni, L. D.Sc. Assistant. Genetics of "brown spots."

Napoli
Dell'Università, Istituto di Genetica

See DIS 34:147.
Pavia
Università di Pavia, Istituto di Genetica

See DIS 34:134.

Roma
Istituto di Genetica, Città Universitaria

Micheli, Aldo. Curator of stocks.
Montalenti, Giuseppe. Professor of Genetics. General genetics.
Nicoletti, Benedetto. Assistant Professor. Melanogaster cytogenetics, mutagenesis.
Olivieri, Mancini Angela. Research Fellow. Melanogaster.

JAPAN

Aichi
Nagoya University, Faculty of Agriculture, Department of Animal Breeding

Bito, J. Graduate student. Melanogaster; mutation.
Esaki, K. Instructor. Melanogaster; mutation.
Hayakawa, J. Graduate student. Melanogaster; mutation.
Kondo, K. (Dr.) Professor. General genetic problems.
Nozawa, K. (Dr.) Assistant Professor. Melanogaster, other species; population genetics and mutation.
Ota, N. (Miss) Technical Assistant. Curator of stocks.

Chiba-shi
National Institute of Radiological Sciences, Biology Division

See DIS 34:135.

Hiroshima
Hiroshima University, Faculty of Science, Zoological Laboratory

Minamori, Sumio. Dr. Assistant Professor (on leave, 1961-1962, National Institute of Genetics, Mishima). Melanogaster: population genetics.

Kobe
Kobe University, Biological Laboratory

Fujii, S. Dr. Professor. Chromosomal aberrations; salivary chromosomes; developmental genetics.
Kanehisa, T. Dr. Research Assistant. Biochemical genetics of tumor.
Kawabe, M. Dr. Assistant Professor. Developmental genetics; variations; human genetics.
Kitazume, Y. Research Assistant. Cytological studies of lethal mutations.
Maeda, Y. Assistant in Research. Melanogaster; mutation.

Kyoto
Kyoto University, Faculty of Science, Department of Zoology

Imazumi, Tadashi. Assistant. Physiological genetics and embryology.
Kato, Masaru. Dr. Assistant Professor. Biochemical genetics and embryology.
Kato, Mikio. Dr. Research associate. Biochemical genetics. (Present address: Department of Zoology, University of Ottawa, Ottawa 2, Canada.)
Miyoshi, Yasuhiro. Graduate student. Physiological genetics.
Nakamura, Kenji. Dr. Professor. Cytogenetics and physiology.
Okuda, Chizuko (Miss). Technical Assistant. Curator of stocks.
Misima, Sizuoka-ken
National Institute of Genetics

Fuwa, K. Research Assistant. Population genetics; deleterious genes in natural populations.
Hirayumi, Y. Dr. Research Member. Population genetics.
Imai, Y. (Miss) Technical Assistant.
Iyama, S. Dr. Research Member. Population genetics; competition and migration (in University of Minnesota, Department of Agronomy, St. Paul).
Kimura, M. Ph.D. Research Member. Population genetics; theoretical (in University of Wisconsin, Madison).
Masuda, H. (Miss) Technical Assistant.
Minamori, S. Dr. (Assistant Professor of Hiroshima University, Visiting Researcher) Population genetics; deleterious genes in natural populations.
Mukai, T. Ph.D. Research Member. Population genetics; radiation and polygene.
Nakamura, K. (Miss) Technical Assistant.
Naruse, T. Dr. Research Member. Population genetics; competition and migration.
Nawa, S. Dr. Research Member. Biochemical genetics; pteridine and nucleic acid (in University of Texas, Austin).
Oshima, C. Dr. Head of Department. Population genetics; resistance, radiation and deleterious genes in natural populations.
Sakaguchi, B. Dr. Research Member. Biochemical genetics; enzymes.
Sakai, K. Dr. Head of Department. Population genetics; competition and migration.
Taira, T. Dr. Research Member. Biochemical genetics; eye pigment formation and metamorphosis.
Toyofuku, Y. (Mrs. Tonomura) Dr. Research Member. Cytogenetics.
Yamada, Y. Dr. Research Member. Population genetics; mutation and selection (in Purdue University, Population Genetics Institute, Lafayette).

Mitaka, Tokyo
International Christian University, Biology Department

Sinoto, Y. Professor. Salivary chromosomes.
Shoji, T. Instructor. Salivary chromosomes.
Kaminishi, H. (Mrs.) Research Fellow. Salivary chromosomes.

Okayama, Kobe
Konan University, Biological Laboratory

Inouye, I. Research Associate. Melanogaster, selection.
Kaji, S. Dr. Assistant Professor. Melanogaster, selection, physiological genetics.
Takaya, H. Dr. Professor. Melanogaster, selection.

Osaka
Osaka University, Faculty of Medicine, Department of Genetics; and
Osaka University, Faculty of Science, Biological Institute

Fujio, Y. Graduate Student. Drosophila: embryological genetics.
Hiraga, S. Graduate Student. Musca: biochemical genetics.
Ichikawa, S. (Miss) Technical Assistant. Curator of stocks.
Kikawara, H. Dr. Professor. Drosophila and Musca: chemical genetics and resistance to insecticides.
Kuroda, Y. Dr. Lecturer. Drosophila: embryological genetics. (Present address: Department of Zoology, University of Chicago, Chicago, Illinois, U. S. A.)
Nobuki, R. (Miss) Technical Assistant. Curator of stocks.
Ogita, Z. Dr. Assistant. Drosophila and Musca: chemical genetics and resistance to insecticides.
Ottori, Y. (Mrs.) Graduate Student. Musca: cytogenetics and sex-determination.
Seki, T. Dr. Lecturer. Drosophila, Musca and Bombyx: chemical genetics.
Tsukamoto, M. Dr. Assistant. Drosophila and Musca: mutations, chemical genetics and resistance to insecticides.
Watanabe, H. (Mrs.) Graduate Student. Musca: chemical genetics.
Sakai, Osaka
University of Osaka Prefecture, Department of Biology

Ogaki, M. Dr. Assistant Professor. Melanogaster: genetics of physiological character.
Tanaka, E. Assistant. Melanogaster: physiological genetics.

Sapporo
Hokkaido University, Faculty of Science, Department of Zoology

Kansko, A. Research assistant. Geographical distribution; cytogenetics.
Makino, S. Dr. Professor. Cytogenetics; population genetics.
Momma, E. Dr. Assistant Professor. Geographical distribution; cytogenetics; population genetics.
Shima, T. Research assistant. Geographical distribution; cytogenetics.
Takada, H. Dr. Research assistant. Taxonomy; ecology.

Tokyo
Tokyo Metropolitan University, Faculty of Science, Department of Biology, Setagaya-ku

Ichida, H. (Miss) Graduate student. Melanogaster: biochemical genetics, tumor.
Ikeda, H. Graduate student. Bifasciata, other species; population genetics, cytoplasmic sex-ratio.
Kitagawa, O. Dr. Research Assistant. Bifasciata, melanogaster: population genetics, radiation genetics.
Kurokawa, H. Dr. Lecturer. Auraria, other species; population genetics, taxonomy.
Moriwaki, D. Dr. Professor. Melanogaster, bifasciata, other species: population genetics, gene analysis, radiation genetics.
Ohba, S. Assistant Professor. Melanogaster, other species: population genetics, ecology, tumor.
Ohnishi, E. Dr. Research Assistant. Melanogaster, virilis: biochemistry.
Okada, T. Dr. Professor. Various species: variations, taxonomy, ecology.
Tobari, I. Research Assistant. Melanogaster: radiation genetics.
Tobari (Nakajima), Y. (Mrs.) Research Assistant. Ananassae, other species: population genetics, heterosis, gene analysis.
Tsukamoto, H. (Miss) Technical assistant.

KOREA

Kongju, Chung Cheong Nam Do
Kongju National Teachers' College, Department of Biology

See DIS 34:137.

Kwangju, Chunnam
National Chunnam University, College of Liberal Arts and Sciences, Department of Biology

Kim, D. U. Assistant Professor. Microbial genetics.
Kim, K. W. Assistant Professor. Drosophila taxonomy.
Park, M. S. Instructor.
Wui, I. S. Instructor.

Seoul
Chung-Ang University, College of Liberal Arts and Sciences, Department of Biology

Chun, W. S. Graduate student. Geographical survey.
Lee, C. S. Graduate Research Assistant. Cytology.
Lee, T. J. Assistant Professor. Population genetics, geographical distribution.
Seoul
Seoul National University, Department of Zoology

Kang, Yung Sun. Dr. Professor. Cytology.
Chung, Chil K. Instructor. Cytogenetics.

Seoul
Sung Kyun-Kwan University, College of Arts and Science, Department of Biology

Kim, D. S. Graduate student. Migration and competition.
Paik, Y. K. Dr. Consultant. (Permanent address: Yonsei University, Department of Biology, Seoul)
Sung, K. C. Graduate student. Migration and competition.

Seoul
Yonsei University, College of Science and Engineering, Department of Biology

Kim, D. S. Graduate student. (Permanent address: Sung Kyun-Kwan University) Migration and competition.
Paik, Y. K. Dr. Associate Professor. Chairman. Population genetics.
Sung, K. C. Graduate student. (Permanent address: Sung Kyun-Kwan University) Migration and competition.
Yoo, C. S. Undergraduate assistant.
Youn, J. S. Graduate Research Assistant. Radiation genetics.

NETHERLANDS

Groningen
State University, Genetical Institute, Haren (Gr)

Bult, P. Graduate student. Competition.
du Pui, L. (Miss) Technical assistant.
Fockens, W. (Miss) Technical assistant.

Leiden
Genetisch Laboratorium der Rijksuniversiteit

Gloor, H. J. Professor. Developmental genetics.
Heerkens, G. M. (Miss) Technical Assistant.
Jacobs, A. A. C. M. (Miss) Assistant. Localization mutants D. hydei.
Schepers, A. H. (Miss) Assistant. Eye pigments.
Schulten, C. G. M. Research student. "Sex-ratio."
Volkers, W. S. Assistant.

Leiden
State University, Department of Radiation Genetics, Wassenaarseweg 62

Goedhart, A. (Miss) Technical assistant.
den Hollander, C. J. M. (Miss) Technical assistant.
de Klerk, T. H. (Miss) Technical assistant.
Leigh, Barry. B.Sc. Radiation mutagenesis.
Lommerse, M. A. H. (Miss) Technical assistant.
de Ruiter, F. J. (Miss) Technical assistant.
Sobels, F. H. Ph.D. Professor. Radiation mutagenesis, repair mechanism.
Utrecht
State University, Institute of Genetics, Opaalweg 20

Dykstra, W. T. (Mrs.) Technical assistant.
Hemel, J. O. van. Demonstrator.
Rühke, C. L. Professor, Director.
Tuinstra, E. J. (Mrs.) Stockkeeper.

NORWAY

Bergen
University of Bergen, Zoological Laboratory

Abro, Arnold. c.r. Melanogaster, radiation effects.
Brinkmann, Aug. Jr. Professor of Zoology, Director of the Laboratory.

Blindern
University of Oslo, Institute of Genetics

Smith, Bina W. (Miss) B.Sc. Curator of Stocks.
Sollunn, Frank-Jørgen. Graduate student. Radiation genetics.
Strömaaes, Øistein. Ph.D. Assistant Professor. Radiation genetics.
Wedvik, Hans. Graduate student. Radiation genetics.

Oslo
Norsk Hydro’s Institute for Cancer Research, The Norwegian Radium Hospital

Mossige, Jeanne Coyne. Research Fellow. Radiosensitivity in sperm.

PANAMA

Panama
The Gorgas Memorial Laboratory, Balboa Heights Post Office Box 65, Canal Zone

Pipkin, Sarah Bedichek.

SOUTH AFRICA

Johannesburg
South African Institute for Medical Research

See DIS 34:139.

Johannesburg
University of the Witwatersrand, Department of Zoology

Hartmann, Ingeborg J. Ph.D. Lecturer. Zaprionus; cytogenetics.
Nolte, D. J. D.Sc. Senior Lecturer. Eye pigmentary system; polygenes in geographic strains.
Pillmann, Loré. Curator of Stocks.
Pretoria
University of Pretoria, Department of Genetics

Geerthsen, J. M. P. B.S. Senior lecturer.
Hofmeyr, J. D. J. M.S., Ph.D., D.Phil. Professor.
Nel, P. M. B.S., B.S.(For.) Graduate student. Chromosomal polymorphism.
Van Niekerk, Brenda. Technician.
Van Schaik, Nancy W. M.S., Ph.D. Lecturer.

SPAIN

Barcelona
Universidad, Centro de Genética Animal y Humana del C. S. I. C.

Alcobé, S. (Mr.) Dr. Director of the Centro. Professor of Anthropology.
Cama, J. (Mr.) Technical Assistant. Curator of Stocks.
Fusté, M. (Miss) Graduate student. D. subobscura populations.
Nadal, A. (Miss) Graduate student. Lethals in natural populations.
Pons, J. (Mr.) Dr. Research worker. Human genetics.
Prevosti, A. (Mr.) Dr. Head of Drosophila Department. Population genetics.

Madrid 6
Centro de Investigaciones Biológicas, Laboratorio de Genética, Velázquez 138

Miralles, L. Graduate student. Cytogenetics.
Morey, M. Research Assistant. Mutagenesis.
Ortiz, E. Dr. Head of Department. Mutagenesis.
Ramírez, P. Technical Assistant.
Rodríguez, C. Graduate student. Cytogenetics.
Solana, I. Technical Assistant.
Torroja, E. Research Assistant. Mutagenesis.

SWEDEN

Stockholm
University of Stockholm, Institute of Genetics

Eiche, A. Ph.K. Research Assistant. Melanogaster: population genetics and mutations.
Montelius, I. Ph.K. Research Assistant. Melanogaster: population genetics.
Ramel, C. Ph.D. Research Associate. Melanogaster: interchromosomal effects, viability mutations.

Uppsala 7
University of Uppsala, Institute of Genetics

Lund, B. Curator of Stocks.
Oholendorff, Helga. Ph.D. Research Assistant.
Rasmussen, B. Ph.D. Research Associate. Melanogaster: physiological genetics.
SWITZERLAND

Bern
Zoologisches Institut der Universität

Rosin, Siegfried. Ph.D. Professor. Developmental genetics.

Zürich
Röntgeninstitut der Universität, Strahlenbiologisches Laboratorium

See DIS 34:140.

Zürich
Zoologisches Institut der Eidgenössischen Technischen Hochschule

Kroeger, Heinrich. Dr. Research assistant. Chromosome metabolism; pattern formation.
Müller, Melanie (Miss). Technical assistant.
Schneider, Annemarie (Mrs.). Graduate student. Cytology.
Ulrich, Hans. Dr. Professor. Differential radiation effects on nucleus and cytoplasm; oxygen effect.

Zürich
Zoologisches Institut der Universität

Buck, Dieter. Graduate student. Imaginal discs.
Burla, Hans. Ph.D. Professor. Taxonomy, population genetics.
Chen, Pei Shen. Ph.D. Professor. Physiology and development.
Diem, Claudia. Graduate student. Enzymes.
Greuter, Mark. Graduate student. Release experiments with Drosophila species.
Hadorn, Ernst. Ph.D. Professor. Developmental and biochemical genetics; lethals.
Munz, Peter. Graduate student. Enzymes.
Nöthiger, Rolf. Assistant. Imaginal discs.
Schläpfer, Theo. Graduate student. Imaginal discs.
Weismann, Hanspeter. Graduate student. Metabolism.
Zürcher, Christian. Graduate student. Wild type allele of ebony (e).

UNITED ARAB REPUBLIC

Alexandria, Egypt
University of Alexandria, Faculty of Agriculture

Dawood, M. M. Ph.D. Lecturer. Lethals in natural populations of Drosophila. On study leave at the Department of Genetics, University of California, Berkeley 4, U. S. A.


Moawad, H. B.Sc. Graduate student. Heritability under severe conditions.


Shoeb, Y. Z. Dipl. Agric. Technical assistant.


Tantawy, A. O. Ph.D. Associate professor and acting head of the division. Population genetics; radiation genetics and physiological genetics; studies on natural populations of Drosophila melanogaster and D. simulans.

Assuit
University of Assuit, Department of Genetics

See DIS 34:141.

UNITED STATES

Alliance, Ohio
Mount Union College, Department of Biology

Blount, Jerry L. Ph.D. Associate Professor. Chairman of Department. Chemical mutagenesis; longevity factors.

Savage, Ellery. Technical Assistant.

Ames, Iowa
Iowa State University, Genetics Department

Gowen, John W. Ph.D. Professor. Melanogaster: crossing over, gene structure and physiological action; heterosis.

Hollander, W. F. Ph.D. Professor. General genetics.

Kloos, Wesley E. Graduate student. Simulans.

Stadler, Janice (Miss). Ph.D. Assistant Professor. Melanogaster: agents for mutations, heterosis.

Thompson, Peter E. Ph.D. Assistant Professor. Melanogaster: mutation.

Amherst, Massachusetts
Amherst College, Department of Biology

Casey, Lucy (Mrs.). Curator of Stocks, Research Assistant.

Hexter, W. M. Ph.D. Associate Professor. Genetic fine structure and crossing over.

Ives, P. T. Ph.D. Research Associate. Radiation and population genetics.

Plough, H. H. Ph.D. Professor Emeritus. Mutation and environmental effects.

Russell, Phyllis (Mrs.). Research Assistant.

Tiffany, Barbara (Miss). Technical Assistant.

Yost, H. T. Jr. Ph.D. Associate Professor. Cell particulates and radiation effects.

Ann Arbor, Michigan
The University of Michigan, Department of Zoology

File, Sharon. Undergraduate student.

Randerson, Sherman. Graduate student.

Rizki, Rose M. Research.

Rizki, T. M. Associate Professor.

Argonne, Illinois
Argonne National Laboratory, Division of Biological and Medical Research

See DIS 34:142.
Athens, Georgia
University of Georgia, Department of Zoology

See DIS 34:142.

Austin, Texas
University of Texas, Department of Zoology, Genetics Foundation

Bunde, Daryl. N. I. H. Predoctoral Fellow.
Burmesiter, Maritha (Mrs.). Welch Foundation Predoctoral Fellow.
Chertkoff, Lynn (Mrs.). Research Assistant. Position effect; pseudalleles.
Dickerman, Richard C. N. I. H. Training Grant Predoctoral Fellow.
Elequin, Flora T. M.A. N. I. H. Training Grant Predoctoral Fellow.
Fabergé, A. C. Ph.D. Research Associate. General genetics.
Forrest, H. S. Ph.D. Associate Professor. Biochemical genetics.
Futch, David G. M.A. N. I. H. Training Grant Predoctoral Fellow.
Gerstenberg, Virginia L. (Mrs.). Technical Assistant.
Judd, Burke H. Ph.D. Associate Professor. Position effect; pseudalleles.
Lagowski, Joanna M. (Mrs.). Ph.D. Research Associate. Biochemical genetics.
Norwood, Sharon. Technical Assistant.
Oliver, C. P. Ph.D. Professor. Gene action; human genetics.
Resch, Kathleen. Technical Assistant.
Rinehart, Robert R. N. I. H. Training Grant Predoctoral Fellow.
Schmid, Werner. M.D. Research Associate (Switzerland). Radiation genetics, general genetics.
Wagner, Robert P. Ph.D. Professor. Gene action; biochemical genetics.
Welch, Robert M. Ph.D. Research Associate. Cytochemistry.
Wilson, Florence D. (Mrs.). Research Assistant. Radiation effects.

Baltimore 18, Maryland
Johns Hopkins University, Department of Biology

Caples, Susan W. (Mrs.) B.A. Research Assistant. Melanogaster; comparative study of induced mutation in males and females.
Glass, H. Bentley. Ph.D. Professor. Melanogaster; population genetics of suppressor systems (erupt and tumor); gene action of su-er and su-tu; tryptophan metabolism in Drosophila; radiation and oxygen effects; comparative effects of mutagens on males and females at different ages.
Mahowald, Anthony P. B.S. Graduate student. Electron microscopy of early embryogenesis; developmental cytology of early embryonic lethals.
Marzuf, George S. B.S., M.S. Graduate student. Nature of gene action and interactions with specific suppressors; tryptophan metabolism in D. melanogaster.
Ritterhoff, Rebecca K. (Mrs.) B.S. Research Staff Assistant. Melanogaster: comparative study of induced mutation in males and females; effect of very low doses of ionizing radiation; Minutes: recessive lethals and spontaneous visibles in males and females; effects of oxygen concentration.
Ursprung, Heinrich. Ph.D. Research Associate. Imaginal discs; xanthine dehydrogenase.
Wright, Eileen X. (Mrs.) B.A. Research Assistant. Ontogeny of gene-enzyme systems; phenogenetics of embryonic lethals.
Wright, Theodore R. F. Ph.D. Assistant Professor. Ontogeny of gene-enzyme systems; esterases and xanthine dehydrogenase; phenogenetics of embryonic lethals.
Baton Rouge, Louisiana
Louisiana State University, Department of Zoology

Brannon, James R., M.S. Graduate student.
Iyengar, Shanta V., Ph.D. Assistant Professor.
Prestridge, Martha Ann. Undergraduate research worker.

Berea, Kentucky
Berea College, Department of Biology

McCune, Thomas. Undergraduate technical assistant.
Seto, Frank. Ph.D. Developmental genetics.

Berkeley, California
University of California, College of Agriculture, Department of Genetics

Brown, Spencer W., Ph.D. Cytogenetics.
Dempster, Everett R., Ph.D. Population genetics.
Sokoloff, Alexander, Ph.D. Population genetics. Comparative genetics of Coleoptera.
Walen, Kirsten H., Ph.D. Cytogenetics.

Berkeley, California
University of California, Department of Zoology

Brunt, Cole M., A.B. Laboratory Technician.
Gottlieb, Frederick, M.A. N.I.H. Predoctoral Trainee. Developmental genetics.
Hildreth, Philip, Ph.D. Research Associate. Mutation, mating behavior.
Horn, Selina, M.A. Graduate student. Sex ratio.
King, Jack, M.A. Graduate student. Developmental genetics.
Stern, Curt, Ph.D. Professor. General.
Tokunaga, Chiyoko, Ph.D. Visiting investigator. Developmental genetics.

Bloomington, Indiana
Indiana University, Department of Zoology

Barbour, Evelyn, M.A. Research Assistant.
Bart, Carol, B.S. Research Assistant.
Edmonson, Margaret (Mrs.), M.A. Graduate Investigator.
Meyer, Helen Unger (Mrs.), Ph.D. Research Associate.
Muller, H. J., D.Sc. Professor.
Oster, Irwin I., Ph.D. Consultant. (Permanent address: Institute for Cancer Research, Philadelphia, Pennsylvania)
Thomas, Sandra (Mrs.), A.B. Research Assistant.
Wagoner, Dale E., A.B. Predoctoral N.I.H. Fellow.
Zimmering, Stanley, Ph.D. Research Executive.

Buffalo 14, New York
University of Buffalo, Department of Biology

Farnsworth, Marjorie W., Ph.D. Lecturer and Research Associate. Melanogaster developmental genetics and biochemistry.
Goldin, Herbert, A.B. Graduate student.
Luchowski, Elizabeth (Mrs.), A.B. Research technician.
Treonar, Katherine, A.B. Graduate student.
Cambridge 38, Massachusetts
Harvard University, The Biological Laboratories

Emrich, Nancy (Mrs.). Research Assistant. Melanogaster.
Jonsson, Ulla-Britt (Miss). Senior Research Assistant. Curator of Stocks. Mutation
and fertility in melanogaster.
Lefevre, George. Ph.D. Radiation genetics; mating behavior in melanogaster.
Rose, Barbara (Miss). Research Assistant. Melanogaster.

Cambridge, Massachusetts
Massachusetts Institute of Technology
See DIS 34:144.

Chambersburg, Pennsylvania
Wilson College
See DIS 34:144.

Chapel Hill, North Carolina
University of North Carolina, Medical School, Department of Biochemistry

Glassman, Edward. Ph.D. Biochemical genetics.
Hodge, Lon. D.V.M. Biochemical genetics.
Karam, J. A.B. Biochemical genetics.
McLean, Janice. B.S. Research Laboratory Supervisor.
Moore, R. Laboratory Assistant.
Parish, J. Laboratory Assistant.
Yen, Terrence T. A.B. Biochemical genetics.

Chapel Hill, North Carolina
University of North Carolina, Department of Zoology

Henderson, Ann S. Graduate Assistant.
Hubbard, William B. M.Ed. Predoctoral Fellow.
Kiesselbach, Theodore H. Honors Student.
James, Judy McNease (Mrs. Wm. S.). A.B. Research Assistant.
Price, Mary Jane (Mrs. Robt. E., Jr.). Research Assistant.
Wall, Lynn. A.B. Research Assistant.
Whittinghill, Maurice. Ph.D. Professor. Irradiation; chemical mutagens; crossing
over.

Chicago 11, Illinois
Loyola University, College of Arts and Sciences

Arnold, Lloyd L. Ph.D. Aging.

Chicago 37, Illinois
University of Chicago, Department of Zoology

Batt, Murray. Research Assistant.
Hubby, Jack L. Ph.D. Instructor. Biochemical genetics.
Roberts, Paul A. M.D. Graduate student. Nondisjunction, developmental genetics.
Sims, Maureen (Miss). B.S. Graduate student.
Spofford, Janice B. Ph.D. Research Associate. Parental effects on phenotype.
Throckmorton, Lynn H. (Mr.) Ph.D. Research Associate. Pteridine metabolism and
protein differences in Drosophila, general Dipteran and Drosophila taxonomy.
Cleveland 15, Ohio
Fenn College, Department of Biology

See DIS 34:145.

Cleveland 15, Ohio
Western Reserve University, Biological Laboratory

See DIS 34:145.

Cold Spring Harbor, New York
Carnegie Institution of Washington, Department of Genetics

Buchanan, Jennie (Mrs. Paul). Research Assistant, Curator of Stocks.
Das, C. C. Ph.D. Guest Investigator (on leave from Allahabad University, Allahabad, India). Cytogenetics.
Gillies, Gloria (Mrs.). Research Assistant.

Cold Spring Harbor, New York
Long Island Biological Association, Biological Laboratory

Krauss, Marian (Miss). B.S. Research Assistant.
Prokop, Barbara (Miss). B.S. Research Assistant. Curator of Stocks.
Talrna, Joy (Mrs.) M.A. Research Assistant.
Taylor, Albert. Technical Assistant.

Columbus, Ohio
Ohio State University, Department of Zoology and Entomology

See DIS 34:145.

Corvallis, Oregon
Oregon State University, Department of Zoology

Heath, Gloria (Mrs.) Student.
Mohler, J. D. Ph.D. Associate Professor.
Neeley, John R. B.S. Graduate student.
Smith, Sheila (Mrs.). Assistant in Zoology.
Thompson, Steven R. B.S. Graduate student.

Davis, California
University of California, Department of Genetics

Bowman, J. T. B.S. N. I. H. Predoctoral Fellow.
Eggert, J. B.S. Laboratory Technician.
Geer, B. W. M.S. Research Fellow.
Green, M. M. Ph.D. Professor.
DeKalb, Illinois
Northern Illinois University, Department of Biological Sciences

Bennett, Jack. Ph.D. Assistant Professor. Selection, insecticide resistance, populations.
Bennett, Katherine Wilson. B.A. Cytogenetics.
Capek, Ronald. B.S. Graduate student. Selection, behavior.
Gianopulos, Harold W. B.S. Graduate student. Selection, insecticide resistance.
Landy, Ronald. B.S. Graduate student. Wild population.
Monkman, Marie (Mrs.). B.S. Graduate student. Research Technician.
Martin, Robert J. B.S. Graduate student. Research Assistant. Radiation.
Weideman, Jeannine. Research Assistant.
Wei, Irene Y. L. Undergraduate student. (Summer N. S. F. Undergraduate Research Participant) Research Technician.
Wu, Ching-kuel. B.S. Graduate student. Populations.

Duarte, California
City of Hope Medical Center, Department of Genetics

Brawley, Mary Anne. Stockkeeper.
Gugler, David H. Research Technician.
Kaplan, William D. Ph.D. Mutagenesis, cytology.
Tanaka, Tatsuya. Ph.D. Cytology.

Durham, North Carolina
Duke University, Department of Zoology

Bird, Margaret Ann (Miss). B.A. Research Assistant.
Burnham, Deborah (Miss). Research Assistant.
Ward, Calvin L. Ph.D. Associate Professor.

East Lansing, Michigan
Michigan State University, Department of Biochemistry

Burnett, Jean B. Ph.D. Research Associate.
Bernhard, Karen L. Technician.
Fox, Allen S. Ph.D. Professor.
Puch, Morton S. M.S. Graduate Research Assistant.
Kan, James L. Ph.D. N. I. H. Postdoctoral Fellow.
Kang, Suk Hee. B.S. Graduate Research Assistant.
Kapetan, Anne S. Technician.
Parzen, Sheldon D. B.S. Graduate Research Assistant.
Yoon, Sei Byung. Ph.D. Research Associate.

East Lansing, Michigan
Michigan State University, Department of Zoology

Camp, Herbert L. Technician.
DeVries, JoAnne K. Graduate student.
Myszewski, Michael E. Graduate student.
Nugent, Karen L. Technician.
Seaton, Robert K. N. S. F. Research Participant.
Stanich, Gloria J. N. S. F. Research Participant.
Trosko, James E. N. D. E. A. Predoctoral Fellow.
Yanders, Armon F. Associate Professor. Radiation effects; mutagenesis; fertilization.
Eugene, Oregon
University of Oregon, Department of Biology

Clancy, C. W. Ph.D. Professor. Developmental genetics.
Dorsey, R. Graduate student. Statistician.
Ehrlich, Elizabeth (Mrs.). Research Assistant Adj. Characteristics of sex-linked lethals.
Erickson, J. M.S. Instructor. Meiotic drive.
Farhang, M. Helper.
Foster, T. B.S. Graduate student. Mutations and chromosomal aberrations.
Hamilton, J. (Miss) Technician. Tandem metacentrics.
Johnson, R. B.S. Graduate student. X-linked non-autonomous lethals.
Landenberger, M. (Mrs.) B.S. Research Assistant.
Masters, N. M.S. Research Assistant. Developmental genetics.
Mickel, S. (Miss) Undergraduate Research Participant. Statistician.
Novitski, E. Ph.D. Professor. (On leave at Zoologisches der Universität, Zürich, Switzerland)
Parker, D. M. (Mrs.) Research Assistant Adj. D. simulans.
Teviotdale, F. (Miss) B.S. Graduate student.

Evanston, Illinois
Northwestern University

Butterworth, Francis M. B.A. Graduate student. Melanogaster, cytochemistry and ultrastructure of the fat body.
Falk, Gretchen J. B.A. Graduate student. Autoradiography.
Green, Christopher C. Undergraduate research student.
King, Robert C. Ph.D. Associate Professor. Melanogaster oogenesis.
Koch, Elizabeth A. B.S. Graduate student. Melanogaster, ultrastructure of fes ovaries.
Mills, Richard P. Undergraduate. Willistoni, ultrastructure.
Packalis, Helen. B.S. Curator of stocks.
Smith, Patricia A. B.S. Graduate student.

Fayetteville, Arkansas
University of Arkansas, Department of Zoology

Bryniarski, Teresa. Research assistant.
Clayton, Frances E. Ph.D. Associate Professor. Radiation effects; development.
Halpern, Lynda S. (Mrs.) B.S. Graduate assistant. Radiation effects.

Gainesville, Florida
University of Florida, Department of Biology

Wallbrunn, Henry M. Mutation rates, population genetics.

Harrisonburg, Virginia
Eastern Mennonite College, Department of Biology

Jacobs, M. E. Ph.D. Professor. Melanism.

Hiram, Ohio
Hiram College, Department of Biology

Friedman, Lawrence D. Ph.D. Assistant Professor. General genetics.

Houston, Texas
Rice University, Department of Biology

See DIS 34:147.
Houston, Texas
University of Texas, M. D. Anderson Hospital and Tumor Institute, Department of Biology
Alexander, Mary L. Ph.D. Radiation; population genetics.
Bergendahl, Janet (Mrs.). M.A. Research Assistant.
Duval, Donya (Miss). B.A. Research Technician II.
Haas, Felix L. Ph.D. Radiation; biochemical genetics.
McKinley, Kay (Miss). B.A. Research Technician II.

Iowa City, Iowa
University of Iowa, Department of Zoology
Brosseau, George E., Jr. Ph.D. Assistant Professor. Melanogaster: genetics of the Y chromosome.
LeVier, Robert L. Undergraduate student assistant.
Gilmore, G. Thomas. Undergraduate student assistant.

Ithaca, New York
Cornell University, Plant Breeding Department
Baumann, James L. Graduate Research Assistant.
Everett, Herbert L. Ph.D. Associate Professor. General genetics.
Gutermann, Hilda (Mrs.). Research Assistant.
Loomis, Margaret (Mrs.). Technical Assistant.
Myers, Oval. Graduate Teaching Assistant.
Sanderson, K. E. Research Associate. General genetics.
Schafrick, Carol. Graduate Teaching Assistant.
Silberman, June (Mrs.). Research Assistant.
Suska, Jadwiga (Mrs.). Research Assistant.
Thompson, Margaret Emmerling. Ph.D. Assistant Professor. General genetics.
Vanoucek, E. G. Graduate Research Assistant.

Jamaica, New York
St. John's University, Department of Biology, Graduate School
Fuscaldo, Kathryn E. Ph.D. Assistant Professor. Biochemical genetics.
Siracusano, Vincent C. Graduate Research Assistant. Biochemical genetics.
Gonnella, Victoria M. Graduate Research Assistant. Immunogenetics.

Johnson City, Tennessee
East Tennessee State College, Department of Biology

Lafayette, Indiana
Purdue University, Department of Biological Sciences
See DIS 34:148.

Lafayette, Indiana
Purdue University, Population Genetics Institute
Bartlett, A. C. M.S. Instructor. Radiation genetics.
Bell, A. E. Ph.D. Professor. Population genetics, selection, G x E interactions.
Bhat, P. N. M.S. Graduate Assistant. Population genetics.
Englert, D. C. M.S. Graduate fellow. Population genetics.
Hardin, R. T. M.S. Graduate Research Assistant. G x E interactions.
Pare, J. P. M.S. Graduate fellow. Selection methods.
Shideler, Doris (Mrs.). Research assistant.

Lawrence, Kansas
University of Kansas, Department of Entomology
See DIS 34:149.

Le Mars, Iowa
Westmar College, Department of Biology
Divelbiss, J. E. Ph.D. Assistant Professor. Complex loci, red eye pigments.

Lexington, Kentucky
University of Kentucky, Department of Zoology
Carpenter, John M. Ph.D. Professor and Department Head. Seasonal fluctuations of Drosophila in relation to wild yeast populations, reproductive potential, gene ecology.
Gilliland, Karen P. Student Assistant.
Semp, Bernard A. Graduate Assistant.
Stewart, Walter H. Graduate Assistant.

Lincoln, Nebraska
The University of Nebraska, Department of Zoology
Lund, Douglas E. Graduate student. D. obscura group CO₂ sensitivity
Miller, Dwight D. Professor.
Stone, Larrie E. Graduate student. D. affinis cytology.
Sulerud, Ralph L. Graduate student. D. melanogaster CO₂ sensitivity.

Logan, Utah
Utah State University, Department of Zoology
Egbert, Larre N. B.S. N. I. H. Fellow. Biometrical genetics.
Gardner, Eldon J. Ph.D. Professor. Melanogaster: mutants of the head region.
Johnson, George R. M.A. Research Assistant. Melanogaster: population studies on genes related to maternal effects.
Simmons, John R. Ph.D. Assistant Professor. Melanogaster: biochemical genetics.
Sorensen, William K. B.S. Graduate student. Melanogaster: development of head abnormalities.

Los Angeles, California
University of California, Department of Botany
De Young, Patricia. Laboratory Assistant.
Epling, Carl C. Ph.D. Professor. Pseudoobscura: population genetics.
Mayhew, Stephen. Undergraduate Technical Assistant.
McCullough, Marilyn. Undergraduate Technical Assistant.
Whitesel, Barbara. Laboratory Assistant.

Los Angeles, California
University of California, Department of Zoology
Carlson, Elof A. Ph.D. Assistant Professor. The dumpy locus; comparative mutagenesis.
Corwin, Harry. B.A. Graduate student.
Falk, Peter. Student assistant, University High School.
Hawkins, Evelyn. Technical assistant.
Hendrickson, Robert. B.A. Graduate student.
Phillips, Barry. B.A. Summer investigator from Queen's University, Canada.
Phillips, Claire. B.A. Stock-keeper and research assistant.
Sederoff, Ronald. B.A. Graduate student.

Madison, Wisconsin
University of Wisconsin, Departments of Genetics and Medical Genetics and Zoology

Abrahamson, Seymour. Ph.D. Assistant Professor. Radiation genetics.
Baumiller, Robert. S.J., Ph.D. Post-doctoral Fellow of the National Foundation.
Chung, Yong Jai. B.S. Graduate student.
Coifman, Robert. B.E.P. Graduate student.
Crow, James F. Ph.D. Professor.
Davis, Brian. B.A. Research Assistant.
Greenberg, Hayla (Miss). M.S. Graduate student.
Lux, Edith (Mrs.). Project Assistant.
Maruyama, Takeo. M.S. Graduate student.
Mattson, Thomas. B.A. Graduate student.
Rosenfeld, Averil (Mrs.). B.S. Project Assistant.
Sandler, L. Ph.D. Assistant Professor.
Thomas, Constance (Miss). M.S. Project Assistant.
Voynow, Nancy (Mrs.). B.A. Research Assistant.

Minneapolis 14, Minnesota
University of Minnesota, Departments of Zoology and Animal Husbandry

See DIS 34:150.

Moscow, Idaho
University of Idaho, Department of Biological Sciences

See DIS 34:150.

Newark, New Jersey
Rutgers, The State University, 40 Rector Street

See DIS 34:150.

New Haven 11, Connecticut
Albertus Magnus College, Department of Biology

Cullen, Sister Mary Urban. O.P., Ph.D. Professor. Developmental genetics.

New Haven, Connecticut
Yale University, Department of Zoology

Counce, Sheila J. (Mrs. R. Bruce Nicklas) Ph.D. Research Associate. Developmental genetics, experimental embryology.
Grabicki, Eugenia (Mrs.). Curator of Stocks and Technician.
(On leave from Department of Zoology, Jagiellonian University, Cracow, Poland, until September, 1962.)
Leventhal, Elaine (Mrs.). M.S. N.I.H. Pre-doctoral Trainee. Developmental genetics and cytology.
Maxim, Peter. Undergraduate. N. S. F. Undergraduate Research Program. Population genetics.

Mills, Richard P. Undergraduate. Developmental genetics, heritable infections.

Nicklas, R. Bruce. Ph.D. Assistant Professor. Cytology of Diptera.

Passano, Kari Nordback (Mrs.). Cand. Real. Guest.

Poulson, D. F. Ph.D. Professor. Physiological and developmental genetics, hereditary infection.

Rosner, J. L. B.S. Graduate Teaching Assistant. Microbial genetics.


New York 27, New York
Columbia University, Department of Zoology

Barker, J. S. F. Ph.D. (University of Sydney, Australia) Fullbright Fellow. Interspecific competition.

Carmody, George. Graduate student. Reproductive isolating mechanisms.

Dobzhansky, Th. Professor. Population genetics: pseudoobscura, persimilis, willistoni, prosaltans, and other species.


Mishara, Joan. Graduate student. Population genetics.

Mourad, Abd el Khalik. Graduate student. (University of Alexandria, Egypt) Population genetics and radiation.

Pavlovsky, O. A. Research Assistant. Cytology; population genetics.


Sankaranarayan, Krishna. Graduate student. (Annamalai University, India) Population genetics.

Solima, Angela. Ph.D. (University of Naples, Italy) Population genetics.

Spassky, Boris. Research Associate. Comparative genetics of species.

Spassky, N. P. (Mrs.) Research Assistant. Population genetics.


Tidwell, Thomas. Graduate student. Reproductive isolating mechanisms.


Weisbrot, David. Graduate student. Melanogaster, simulans.

New York 21, New York
The Rockefeller Institute

See DIS 34:150.

Norman, Oklahoma
University of Oklahoma, Department of Zoology

See DIS 34:150.

Notre Dame, Indiana
University of Notre Dame, Department of Biology

Bender, Harvey A. Ph.D. Assistant Professor. Developmental genetics.

Craig, George B. Ph.D. Associate Professor. Population genetics (Aedes).

Moskowski, Theresa (Miss). Curator of Stocks, Technician.

Oak Ridge, Tennessee
Oak Ridge National Laboratory, Biology Division, P. O. Box Y

Grell, E. H. Ph.D. Chromosome behavior and biochemical genetics.
Grell, Rhoda F. Ph.D. Chromosome behavior.
Lindsley, Dan L. Ph.D. Chromosome behavior and radiation genetics.
Petty, John. B.S. Research assistant.
Pratt, Guthrie T. (Mrs.). M.S. Research assistant.
Scandlyn, Bobbie J. (Miss) B.S. Research assistant.
Suzuki, David T. Ph.D. Chromosome behavior.
von Borstel, R. C. Ph.D. Radiation genetics.
Von Halle, Elizabeth S. (Mrs.) B.A. Research consultant.
Welshons, William J. Ph.D. Pseudoallelism.
Willerson, Ruby D. (Mrs.) Curator of stocks.

Pasadena, California
California Institute of Technology, Division of Biology

Del Campo, Gladys. B.S. Research Assistant.
Kiger, John. Student.
Lewis, E. B. Ph.D. Professor.
Markowitz, E. Student.
Mitchell, Annamarie (Mrs.) Dipl. Lab.
Mitchell, H. K. Ph.D. Professor.
Mora, Sergio. M.S. Curator of Stocks.
Seeof, R. L. Ph.D. Research Fellow.
Sturtevant, A. H. Ph.D. Professor.

Philadelphia 11, Pennsylvania
The Institute for Cancer Research, Fox Chase, Division of Chemotherapy

See DIS 34:151.

Philadelphia 11, Pennsylvania
The Institute for Cancer Research, Fox Chase, Department of Genetics and Cytochemistry

See DIS 34:152.

Philadelphia 22, Pennsylvania
Temple University, Department of Biology

See DIS 34:152.

Philadelphia 29, Pennsylvania
Woman's Medical College, Department of Anatomy

Campbell, Shirley. Cytology Assistant.
Levitan, Max. Ph.D. Associate Professor. Population genetics.
Schiller, Ruth. Research Assistant.
White, Susan. Technical Assistant.

Pittsburgh 13, Pennsylvania
University of Pittsburgh, Department of Biological Sciences

Carver, James E., Jr. M.S. Graduate student. Research Assistant. Lethals in melanogaster populations.
Langer, Bozena (Mrs.). Ph.D. Research Associate. Mating propensity, persimilis.
Spiess, Eliot B. Ph.D. Associate Professor. Population genetics.
Spiess, Luretta D. (Mrs.) Ph.D. Research Associate. Population genetics.
Sweet, Edward E. Ph.D. Research Associate. Sterility in populations.
Portland, Oregon
Reed College, Department of Biology

See DIS 34:152.

Pullman, Washington
Washington State University, Department of Zoology

Hudson, James E. Student.

Raleigh, North Carolina
North Carolina State College, Department of Genetics

Brown, J. C. (Mrs.) Research Assistant.
Bruck, David. Graduate Research Assistant.
Collins, F. F. (Mrs.) Research Assistant.
Council, S. B. (Mrs.) Research Assistant.
Dobie, N. B. (Mrs.) Research Assistant (Stockkeeper).
Dyson, J. G. (Mrs.) Graduate Research Assistant.
Kojima, Ken-ichi. Ph.D. Experimental and theoretical population genetics; quantitative genetics.
Mettler, Lawrence E. Ph.D. Experimental population genetics; cytogenetics.
Richardson, R. H. N. S. F. Cooperative Fellow (Graduate Research Assistant).
Schaffer, H. E. N. D. E. A. Predoctoral Fellow.
Wing, M. S. (Mrs.) Research Assistant.

Richmond 19, Virginia
Medical College of Virginia, Department of Biology and Genetics

Bridges, Elizabeth P. B.S. Research Assistant.
Hughes, Roscoe D. Ph.D. Professor. Cytogenetics.
Townsend, J. Ives. Ph.D. Assistant Professor. Population genetics; marginal populations.

Ridgefield, Connecticut
New England Institute for Medical Research

Freeborn, John. Technical Assistant.
Mahler, Marilyn (Mrs.). B.A. Research Assistant.
Mickey, George H. Ph.D. Cytogeneticist. Mutations.
Sondhi, Gunthild (Mrs.). Technician.

Riverside, California
University of California, Department of Biology

See DIS 34:153.

Rochester 20, New York
University of Rochester, Department of Biology

See DIS 34:153.

St. Louis, Missouri
Saint Louis University, Department of Biology

See DIS 34:153.

St. Louis, Missouri
Washington University, Department of Zoology

See DIS 34:153.
Salt Lake City, Utah  
University of Utah, Department of Genetics

Hanks, George D. Ph.D. Population genetics; meiotic drive.
Hochman, Benjamin. Ph.D. Population genetics; isoalleles, lethals.
Prows, Ronald. B.S. Technician.

Salt Lake City, Utah  
University of Utah, College of Medicine, Department of Surgery

Burdette, Walter J. Ph.D., M.D. Professor and Head, Department of Surgery.
Mukherjee, Barid B. Ph.D. Research Associate.
Pilgrim, H. Ira. Ph.D. Research Assistant Professor.
Anderson, Betty. Laboratory technician.
Baumgart, Gerda Isolde. Laboratory technician.
Bigelow, Robert R. Laboratory technician.
Hegewald, Eva. Laboratory technician.
Hegewald, Rudolph J. Laboratory technician.
Janke, Hannelore. Laboratory technician.
Nomura, Koji. Laboratory technician.
Paul, Lloyd A. Laboratory technician.
Pilar, Beatriz M. M. Laboratory technician.
Steinitz, John. Diener.
Stratopoulos, George. Laboratory technician.
Thomas, Carol B. Research Assistant.

San Diego, California  
San Diego State College, Department of Zoology

Johns, Ruth E. B.A. Graduate student.
Lovellette, Edward J. B.A. Graduate student.
Ratty, Frank J. Ph.D. Associate Professor.

Staten Island 1, New York  
Wagner College

Annan, Murvel E. Ph.D.
Reitan, Phillip J. Ph.D. Drosophila development.

Storrs, Connecticut  
University of Connecticut, Department of Zoology and Entomology


Syracuse 10, New York  
Syracuse University, Department of Zoology and Division of Science Teaching

Milkman, Roger D. Ph.D. Associate Professor. Drosophila genetics: population, developmental, and physiological.
Phillips, Donald. B.A. Research Assistant.
Petersen, Kathleen L. (Mrs.) B.A. Graduate student. Punebris.
Gallatte, Alfred T. Ph.D. Professor. Virilis.

Tallahassee, Florida  
Florida State University, Department of Biological Sciences

Edington, C. W. Ph.D. Radiation genetics.
Epler, J. L. M.S. Radiation genetics and chemical mutagenesis.
Tucson, Arizona
University of Arizona, Department of Zoology

See DIS 34:154.

University, Alabama
University of Alabama, College of Arts and Sciences, Department of Biology

Guest, William C. Ph.D. Assistant Professor. Cytogenetics.

University Park, Pennsylvania
Pennsylvania State University, Buckhout Laboratory

Come, Thomas V. M.A. Graduate student.
Crain, Paul. Ph.D. Associate Professor. Cytogenetics.
Nash, Donald J. Ph.D. Assistant Professor. Population genetics.

Upton, New York
Brookhaven National Laboratory, Department of Biology

See DIS 34:155.

Urbana, Illinois
University of Illinois, Department of Psychology, Behavior Genetics Laboratory

Hosteller, Roy C. B.A. Research Assistant.

Urbana, Illinois
University of Illinois, Department of Zoology

Luce, Wilbur M. Ph.D. Professor. Bar series; effect of environmental agents; radiation; effect of chemicals; physiological genetics.
Olson, John B. B.S. Research Assistant. Curator of stocks. Location of mutants.
Tanaka, Eiji. New address: Department of Biology, University of Osaka Prefecture, Sakai, Japan.

Washington 25, D.C.
National Science Foundation, Genetic Biology Program

See DIS 34:155.

Wellesley 81, Massachusetts
Wellesley College, Department of Zoology and Physiology

Bull, Alice Louise. Ph.D. Assistant Professor. Developmental genetics.
<table>
<thead>
<tr>
<th>Alphabetical</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abrahamson, S.</td>
</tr>
<tr>
<td>Aber, A.</td>
</tr>
<tr>
<td>Aeppli, L.</td>
</tr>
<tr>
<td>Aita, Y.</td>
</tr>
<tr>
<td>Ako, S.</td>
</tr>
<tr>
<td>Alderson, T.</td>
</tr>
<tr>
<td>Alexander, M.</td>
</tr>
<tr>
<td>Allan, J.</td>
</tr>
<tr>
<td>Allen, A.C.</td>
</tr>
<tr>
<td>Altenburg, E.</td>
</tr>
<tr>
<td>Altmann, J.</td>
</tr>
<tr>
<td>Anderson, B.</td>
</tr>
<tr>
<td>Anderson, Betty</td>
</tr>
<tr>
<td>Anderson, P.</td>
</tr>
<tr>
<td>Anderson, J.</td>
</tr>
<tr>
<td>Anderson, R.</td>
</tr>
<tr>
<td>Andersson, M.</td>
</tr>
<tr>
<td>Anger, A.</td>
</tr>
<tr>
<td>Anan, M.</td>
</tr>
<tr>
<td>Antochevsky, N.</td>
</tr>
<tr>
<td>Apitzsch, U.</td>
</tr>
<tr>
<td>Arden, L.</td>
</tr>
<tr>
<td>Arnold, L.</td>
</tr>
<tr>
<td>Aronson, M.</td>
</tr>
<tr>
<td>Auerbach, C.</td>
</tr>
<tr>
<td>Bairati, A.</td>
</tr>
<tr>
<td>Baker, W.</td>
</tr>
<tr>
<td>Ball, F.</td>
</tr>
<tr>
<td>Band, H.</td>
</tr>
<tr>
<td>Banerjee, S.</td>
</tr>
<tr>
<td>Barak, E.</td>
</tr>
<tr>
<td>Barber, R.</td>
</tr>
<tr>
<td>Barbour, E.</td>
</tr>
<tr>
<td>Barigeloni, O.</td>
</tr>
<tr>
<td>Barker, J.</td>
</tr>
<tr>
<td>Bart, C.</td>
</tr>
<tr>
<td>Bartlett, J.</td>
</tr>
<tr>
<td>Bartlett, A.</td>
</tr>
<tr>
<td>Bass, E.</td>
</tr>
<tr>
<td>Basile, R.</td>
</tr>
<tr>
<td>Beaman, A.</td>
</tr>
<tr>
<td>Batt, M.</td>
</tr>
<tr>
<td>Betten, J.</td>
</tr>
<tr>
<td>Beaman, J.</td>
</tr>
<tr>
<td>Baumgart, G.</td>
</tr>
<tr>
<td>Beuville, G.</td>
</tr>
<tr>
<td>Beaudoin, J.</td>
</tr>
<tr>
<td>Becker, G.</td>
</tr>
<tr>
<td>Becker, H.</td>
</tr>
<tr>
<td>Behrendt, W.</td>
</tr>
<tr>
<td>Beggs, C.</td>
</tr>
<tr>
<td>Bell, H.</td>
</tr>
<tr>
<td>Bell, A.</td>
</tr>
<tr>
<td>Bender, E.</td>
</tr>
<tr>
<td>Bender, H.</td>
</tr>
<tr>
<td>Bennett, J.</td>
</tr>
<tr>
<td>Bennett, K.</td>
</tr>
<tr>
<td>Bentvelzen, P.</td>
</tr>
<tr>
<td>Berendsen, H.</td>
</tr>
<tr>
<td>Berenzain, J.</td>
</tr>
<tr>
<td>Bergerard, J.</td>
</tr>
<tr>
<td>Bernard, J.</td>
</tr>
<tr>
<td>Bernhard, K.</td>
</tr>
<tr>
<td>Bert, G.</td>
</tr>
<tr>
<td>Bhat, P.</td>
</tr>
<tr>
<td>Bigelow, R.</td>
</tr>
<tr>
<td>Bigler, J.</td>
</tr>
<tr>
<td>Bird, M.</td>
</tr>
<tr>
<td>Bito, J.</td>
</tr>
<tr>
<td>Blair, J.</td>
</tr>
<tr>
<td>Blair, F.</td>
</tr>
<tr>
<td>Blake, F.</td>
</tr>
<tr>
<td>Blout, J.</td>
</tr>
<tr>
<td>Blum, S.</td>
</tr>
<tr>
<td>Boss, T.</td>
</tr>
<tr>
<td>Boehm, V.</td>
</tr>
<tr>
<td>Bond, C.</td>
</tr>
<tr>
<td>Böcking, E.</td>
</tr>
<tr>
<td>Bowman, J.</td>
</tr>
<tr>
<td>Brannin, J.</td>
</tr>
<tr>
<td>Braver, G.</td>
</tr>
<tr>
<td>Braver, N.</td>
</tr>
<tr>
<td>Brewley, M.</td>
</tr>
<tr>
<td>Bregliano, J.</td>
</tr>
<tr>
<td>Breuer, M.</td>
</tr>
<tr>
<td>Bridges, E.</td>
</tr>
<tr>
<td>Brink, N.</td>
</tr>
<tr>
<td>Brinkman, A.</td>
</tr>
<tr>
<td>Brasie, D.</td>
</tr>
<tr>
<td>Brosseau, G.</td>
</tr>
<tr>
<td>Brown, E.</td>
</tr>
<tr>
<td>Brown, J.</td>
</tr>
<tr>
<td>Brown, M.</td>
</tr>
<tr>
<td>Brown, W.</td>
</tr>
<tr>
<td>Brown, L.</td>
</tr>
<tr>
<td>Bruck, D.</td>
</tr>
<tr>
<td>Brun, G.</td>
</tr>
<tr>
<td>Brun, J.</td>
</tr>
<tr>
<td>Brunt, G.</td>
</tr>
<tr>
<td>Bryniarski, T.</td>
</tr>
<tr>
<td>Buchanan, J.</td>
</tr>
<tr>
<td>Buck, D.</td>
</tr>
<tr>
<td>Bull, K.</td>
</tr>
<tr>
<td>Bull, S.</td>
</tr>
<tr>
<td>Bult, F.</td>
</tr>
<tr>
<td>Bunco, A.</td>
</tr>
<tr>
<td>Bunde, D.</td>
</tr>
<tr>
<td>Bunker, M.</td>
</tr>
<tr>
<td>Burdette, W.</td>
</tr>
<tr>
<td>Burdick, A.</td>
</tr>
<tr>
<td>Burger, C.</td>
</tr>
<tr>
<td>Burla, G.</td>
</tr>
<tr>
<td>Burnside, M.</td>
</tr>
<tr>
<td>Burnett, B.</td>
</tr>
<tr>
<td>Burnett, J.</td>
</tr>
<tr>
<td>Burnham, B.</td>
</tr>
<tr>
<td>Butler, L.</td>
</tr>
<tr>
<td>Butcher, P.</td>
</tr>
<tr>
<td>Buzzati-Traverso, A.</td>
</tr>
<tr>
<td>Cachero, N.</td>
</tr>
<tr>
<td>Cana, J.</td>
</tr>
<tr>
<td>Cami, C.</td>
</tr>
<tr>
<td>Camp, H.</td>
</tr>
<tr>
<td>Campbell, R.</td>
</tr>
<tr>
<td>Name</td>
</tr>
<tr>
<td>---------------</td>
</tr>
<tr>
<td>Campbell, S.</td>
</tr>
<tr>
<td>Canuti, N.</td>
</tr>
<tr>
<td>Capek, R.</td>
</tr>
<tr>
<td>Caples, S.</td>
</tr>
<tr>
<td>Caraglio, M.</td>
</tr>
<tr>
<td>Carlson, E.</td>
</tr>
<tr>
<td>Carney, G.</td>
</tr>
<tr>
<td>Carpenter, J.</td>
</tr>
<tr>
<td>Carver, J.</td>
</tr>
<tr>
<td>Casagrande, A.</td>
</tr>
<tr>
<td>Casey, L.</td>
</tr>
<tr>
<td>Castro, J.</td>
</tr>
<tr>
<td>Castignoni, M.</td>
</tr>
<tr>
<td>Castro, C.</td>
</tr>
<tr>
<td>Cattan, A. G.</td>
</tr>
<tr>
<td>Cestari, E.</td>
</tr>
<tr>
<td>Cheyn, A.</td>
</tr>
<tr>
<td>Cheyn, J.</td>
</tr>
<tr>
<td>Cheyn, K.</td>
</tr>
<tr>
<td>Chiquita, J.</td>
</tr>
<tr>
<td>Chisom, A.</td>
</tr>
<tr>
<td>Choi, J.</td>
</tr>
<tr>
<td>Chovnick, A.</td>
</tr>
<tr>
<td>Chun, W.</td>
</tr>
<tr>
<td>Chung, J.</td>
</tr>
<tr>
<td>Chung, C.</td>
</tr>
<tr>
<td>Chu, Y.</td>
</tr>
<tr>
<td>Chioffi, E.</td>
</tr>
<tr>
<td>Cividalli, L.</td>
</tr>
<tr>
<td>Clancy, C.</td>
</tr>
<tr>
<td>Clark, A.</td>
</tr>
<tr>
<td>Clark, E.</td>
</tr>
<tr>
<td>Clarke, J. G.</td>
</tr>
<tr>
<td>Clayton, F.</td>
</tr>
<tr>
<td>Clayton, G.</td>
</tr>
<tr>
<td>Cliquet, R.</td>
</tr>
<tr>
<td>Cohen, B.</td>
</tr>
<tr>
<td>Cohen, J.</td>
</tr>
<tr>
<td>Coifman, M.</td>
</tr>
<tr>
<td>Collins, F.</td>
</tr>
<tr>
<td>Cole, K.</td>
</tr>
<tr>
<td>Collette, A.</td>
</tr>
<tr>
<td>Cox, T.</td>
</tr>
<tr>
<td>Comstock, R.</td>
</tr>
<tr>
<td>Cooke, F.</td>
</tr>
<tr>
<td>Coon, H.</td>
</tr>
<tr>
<td>Cordeiro, A.</td>
</tr>
<tr>
<td>Cordeiro, E.</td>
</tr>
<tr>
<td>Corn, J.</td>
</tr>
<tr>
<td>Cortés, B.</td>
</tr>
<tr>
<td>Cortés, Y.</td>
</tr>
<tr>
<td>Corwin, H.</td>
</tr>
<tr>
<td>Coughlin, A.</td>
</tr>
<tr>
<td>Cousine, S.</td>
</tr>
<tr>
<td>Council, S.</td>
</tr>
<tr>
<td>Covarrubias, E.</td>
</tr>
<tr>
<td>Courreges, B.</td>
</tr>
<tr>
<td>Coye, M.</td>
</tr>
<tr>
<td>Craig, G.</td>
</tr>
<tr>
<td>Craig, J. F.</td>
</tr>
<tr>
<td>Cruickshank, W.</td>
</tr>
<tr>
<td>Cullen, M.</td>
</tr>
<tr>
<td>Cunnings, E.</td>
</tr>
<tr>
<td>da Cunha, A.</td>
</tr>
<tr>
<td>Dalilie, J.</td>
</tr>
<tr>
<td>Dalmon, France</td>
</tr>
<tr>
<td>Das, C.</td>
</tr>
<tr>
<td>David, J.</td>
</tr>
<tr>
<td>Davis, E.</td>
</tr>
<tr>
<td>Davis, D.</td>
</tr>
<tr>
<td>Dawson, M.</td>
</tr>
<tr>
<td>Dearden, M.</td>
</tr>
<tr>
<td>de Capea, A.</td>
</tr>
<tr>
<td>Deconinck, C.</td>
</tr>
<tr>
<td>de Finot, W.</td>
</tr>
<tr>
<td>de Fresqueville, J.</td>
</tr>
<tr>
<td>de Klerk, T.</td>
</tr>
<tr>
<td>del Campo, G.</td>
</tr>
<tr>
<td>del Solar, E.</td>
</tr>
<tr>
<td>de Martinis, S.</td>
</tr>
<tr>
<td>de Martinis, P.</td>
</tr>
<tr>
<td>Dempster, B.</td>
</tr>
<tr>
<td>den Hollander, C.</td>
</tr>
<tr>
<td>de Ruiter, F.</td>
</tr>
<tr>
<td>de Vries, J.</td>
</tr>
<tr>
<td>de Young, P.</td>
</tr>
<tr>
<td>Dharwarajam, M.</td>
</tr>
<tr>
<td>Diament, B.</td>
</tr>
<tr>
<td>Días, N.</td>
</tr>
<tr>
<td>Dickerman, R.</td>
</tr>
<tr>
<td>Dienes, C.</td>
</tr>
<tr>
<td>Díaz, J.</td>
</tr>
<tr>
<td>Dierer, S.</td>
</tr>
<tr>
<td>Dign, A.</td>
</tr>
<tr>
<td>Dilley, E.</td>
</tr>
<tr>
<td>Di Pasquale, A.</td>
</tr>
<tr>
<td>Ditadi, T.</td>
</tr>
<tr>
<td>Dittrich, W.</td>
</tr>
<tr>
<td>Dittrich, W.</td>
</tr>
<tr>
<td>Divalbiss, J.</td>
</tr>
<tr>
<td>Doane, W.</td>
</tr>
<tr>
<td>Dobie, N.</td>
</tr>
<tr>
<td>Dobzhansky, Th.</td>
</tr>
<tr>
<td>Dodds, J.</td>
</tr>
<tr>
<td>Dorn, G.</td>
</tr>
<tr>
<td>Dorsey, R.</td>
</tr>
<tr>
<td>Druker, M.</td>
</tr>
<tr>
<td>Duplat, H.</td>
</tr>
<tr>
<td>du Pui, L.</td>
</tr>
<tr>
<td>Dupvse, F.</td>
</tr>
<tr>
<td>Duval, D.</td>
</tr>
<tr>
<td>Dyer, W.</td>
</tr>
<tr>
<td>Dykkstra, W.</td>
</tr>
<tr>
<td>Dyson, J.</td>
</tr>
<tr>
<td>Ebeling, W.</td>
</tr>
<tr>
<td>Edington, C.</td>
</tr>
<tr>
<td>Edmondson, M.</td>
</tr>
<tr>
<td>Edwards, J.</td>
</tr>
<tr>
<td>Egbert, L.</td>
</tr>
<tr>
<td>Eggert, J.</td>
</tr>
<tr>
<td>Emrlich, E.</td>
</tr>
<tr>
<td>Ehrman, L.</td>
</tr>
<tr>
<td>Name</td>
</tr>
<tr>
<td>------</td>
</tr>
<tr>
<td>A. Eiche</td>
</tr>
<tr>
<td>A. Elens</td>
</tr>
<tr>
<td>A. Elequin</td>
</tr>
<tr>
<td>A. Emara</td>
</tr>
<tr>
<td>N. Enrigh</td>
</tr>
<tr>
<td>D. Epler</td>
</tr>
<tr>
<td>F. Epping</td>
</tr>
<tr>
<td>R. Emara</td>
</tr>
<tr>
<td>N. Emrich</td>
</tr>
<tr>
<td>D. Engert</td>
</tr>
<tr>
<td>J. Epling</td>
</tr>
<tr>
<td>C. Erickson</td>
</tr>
<tr>
<td>K. Esaki</td>
</tr>
<tr>
<td>H. Everett</td>
</tr>
<tr>
<td>A. Fabergé</td>
</tr>
<tr>
<td>M. Fahy</td>
</tr>
<tr>
<td>M. Fahy</td>
</tr>
<tr>
<td>G. Falk</td>
</tr>
<tr>
<td>P. Falk</td>
</tr>
<tr>
<td>R. Falk</td>
</tr>
<tr>
<td>H. Fannele</td>
</tr>
<tr>
<td>H. Farhang</td>
</tr>
<tr>
<td>N. Farnsworth</td>
</tr>
<tr>
<td>S. Fernandez</td>
</tr>
<tr>
<td>S. File</td>
</tr>
<tr>
<td>D. Finlay</td>
</tr>
<tr>
<td>M. Fletcher</td>
</tr>
<tr>
<td>A. Fockens</td>
</tr>
<tr>
<td>G. Forbes</td>
</tr>
<tr>
<td>E. Forbes</td>
</tr>
<tr>
<td>H. Forrest</td>
</tr>
<tr>
<td>T. Foster</td>
</tr>
<tr>
<td>J. Fox</td>
</tr>
<tr>
<td>J. Freeborn</td>
</tr>
<tr>
<td>J. Freed</td>
</tr>
<tr>
<td>O. Freund</td>
</tr>
<tr>
<td>M. Friedlaender</td>
</tr>
<tr>
<td>L. Friedman</td>
</tr>
<tr>
<td>J. Frithson</td>
</tr>
<tr>
<td>J. Fritz-Neglil</td>
</tr>
<tr>
<td>J. Frost</td>
</tr>
<tr>
<td>O. Protase-Pessa</td>
</tr>
<tr>
<td>O. Frydendberg</td>
</tr>
<tr>
<td>M. Fuchs</td>
</tr>
<tr>
<td>J. Fuji</td>
</tr>
<tr>
<td>Y. Fuji</td>
</tr>
<tr>
<td>K. Fussaido</td>
</tr>
<tr>
<td>M. Fuster</td>
</tr>
<tr>
<td>D. Futch</td>
</tr>
<tr>
<td>K. Fuka</td>
</tr>
<tr>
<td>Y. Fuyama</td>
</tr>
<tr>
<td>G. Gale</td>
</tr>
<tr>
<td>E. Gallucci</td>
</tr>
<tr>
<td>A. García-Beallid</td>
</tr>
<tr>
<td>E. Gardner</td>
</tr>
<tr>
<td>H. Gay</td>
</tr>
<tr>
<td>B. Geer</td>
</tr>
<tr>
<td>J. Geerthsen</td>
</tr>
<tr>
<td>B. Geissler</td>
</tr>
<tr>
<td>H. Gelert</td>
</tr>
<tr>
<td>M. Gerletti</td>
</tr>
<tr>
<td>E. Gersh</td>
</tr>
<tr>
<td>V. Gerstenberg</td>
</tr>
<tr>
<td>C. Ghini</td>
</tr>
<tr>
<td>H. Ghosh</td>
</tr>
<tr>
<td>H. Gianopulos</td>
</tr>
<tr>
<td>S. Glavelli</td>
</tr>
<tr>
<td>A. Glavelli</td>
</tr>
<tr>
<td>A. Gibson</td>
</tr>
<tr>
<td>J. Gibson</td>
</tr>
<tr>
<td>K. Gidholm</td>
</tr>
<tr>
<td>K. Gill</td>
</tr>
<tr>
<td>G. Gillies</td>
</tr>
<tr>
<td>K. Gilliland</td>
</tr>
<tr>
<td>J. Gilmore</td>
</tr>
<tr>
<td>H. Glass</td>
</tr>
<tr>
<td>S. Glass</td>
</tr>
<tr>
<td>E. Glassman</td>
</tr>
<tr>
<td>O. Gleaves</td>
</tr>
<tr>
<td>H. Glor</td>
</tr>
<tr>
<td>R. Gloor</td>
</tr>
<tr>
<td>J. Godet</td>
</tr>
<tr>
<td>A. Goldhart</td>
</tr>
<tr>
<td>W. Goetz</td>
</tr>
<tr>
<td>H. Goldin</td>
</tr>
<tr>
<td>E. Goldschmidt</td>
</tr>
<tr>
<td>J. Gomella</td>
</tr>
<tr>
<td>F. Gottlieb</td>
</tr>
<tr>
<td>G. Gottschewski</td>
</tr>
<tr>
<td>J. Gowen</td>
</tr>
<tr>
<td>E. Grabiski</td>
</tr>
<tr>
<td>E. Graeser</td>
</tr>
<tr>
<td>A. Grassmann</td>
</tr>
<tr>
<td>C. Green</td>
</tr>
<tr>
<td>M. Green</td>
</tr>
<tr>
<td>R. Greenberg</td>
</tr>
<tr>
<td>E. Grell</td>
</tr>
<tr>
<td>R. Grell</td>
</tr>
<tr>
<td>E. Greuter</td>
</tr>
<tr>
<td>H. Grisch</td>
</tr>
<tr>
<td>A. Griffen</td>
</tr>
<tr>
<td>C. Grisseau</td>
</tr>
<tr>
<td>L. Grün</td>
</tr>
<tr>
<td>H. Grüneberg</td>
</tr>
<tr>
<td>C. Guerrier</td>
</tr>
<tr>
<td>W. Guest</td>
</tr>
<tr>
<td>D. Gufler</td>
</tr>
<tr>
<td>A. Giglischmi</td>
</tr>
<tr>
<td>M. Guillaumim</td>
</tr>
<tr>
<td>M. Gunson</td>
</tr>
<tr>
<td>H. Gutenmann</td>
</tr>
<tr>
<td>F. Hass</td>
</tr>
<tr>
<td>N. Hadler</td>
</tr>
<tr>
<td>W. Hadorn</td>
</tr>
<tr>
<td>H. Hagens</td>
</tr>
<tr>
<td>J. Haldane</td>
</tr>
<tr>
<td>C. Halper</td>
</tr>
<tr>
<td>O. Halper</td>
</tr>
<tr>
<td>L. Halpern</td>
</tr>
<tr>
<td>J. Hamilton</td>
</tr>
<tr>
<td>A. Hamps</td>
</tr>
<tr>
<td>C. Hanks</td>
</tr>
<tr>
<td>E. Hanly</td>
</tr>
<tr>
<td>A. Hannah-Alava</td>
</tr>
<tr>
<td>A. Hansen</td>
</tr>
<tr>
<td>I. Hansten</td>
</tr>
<tr>
<td>R. Hardin</td>
</tr>
<tr>
<td>H. Harlock</td>
</tr>
<tr>
<td>Name</td>
</tr>
<tr>
<td>-----------------</td>
</tr>
<tr>
<td>Harmoinen, L.</td>
</tr>
<tr>
<td>Harrow, C.</td>
</tr>
<tr>
<td>Harrington, D.</td>
</tr>
<tr>
<td>Harrison, B.</td>
</tr>
<tr>
<td>Hartzmann, J.</td>
</tr>
<tr>
<td>Hartshorne, J.</td>
</tr>
<tr>
<td>Hauser, A.</td>
</tr>
<tr>
<td>Hawkins, N.</td>
</tr>
<tr>
<td>Hawkins, E.</td>
</tr>
<tr>
<td>Hasekawa, T.</td>
</tr>
<tr>
<td>Hayman, D.</td>
</tr>
<tr>
<td>Heath, C.</td>
</tr>
<tr>
<td>Heed, W.</td>
</tr>
<tr>
<td>Heerkens, C.</td>
</tr>
<tr>
<td>Hegewald, E.</td>
</tr>
<tr>
<td>Hegewald, R.</td>
</tr>
<tr>
<td>Heinonen, E.</td>
</tr>
<tr>
<td>Heinsso, M.</td>
</tr>
<tr>
<td>Hessel, J.</td>
</tr>
<tr>
<td>Hendrickson, R.</td>
</tr>
<tr>
<td>Henderson, A.</td>
</tr>
<tr>
<td>Henke, H.</td>
</tr>
<tr>
<td>Henningsen, K.</td>
</tr>
<tr>
<td>Herzkowitz, J.</td>
</tr>
<tr>
<td>Hess, O.</td>
</tr>
<tr>
<td>Heuts, M.</td>
</tr>
<tr>
<td>Hexter, W.</td>
</tr>
<tr>
<td>Higgins, W.</td>
</tr>
<tr>
<td>Hildreth, F.</td>
</tr>
<tr>
<td>Hillman, R.</td>
</tr>
<tr>
<td>Hime, N.</td>
</tr>
<tr>
<td>Hinton, C.</td>
</tr>
<tr>
<td>Hiraga, S.</td>
</tr>
<tr>
<td>Hiraiizumi, Y.</td>
</tr>
<tr>
<td>Hiroyoshi, T.</td>
</tr>
<tr>
<td>Hirsh, J.</td>
</tr>
<tr>
<td>Hirsch, D.</td>
</tr>
<tr>
<td>Hochmenn, R.</td>
</tr>
<tr>
<td>Hodge, L.</td>
</tr>
<tr>
<td>Hoeningberg, H.</td>
</tr>
<tr>
<td>Hofstey, J.</td>
</tr>
<tr>
<td>Höhne, G.</td>
</tr>
<tr>
<td>Hollander, W.</td>
</tr>
<tr>
<td>Holler, A.</td>
</tr>
<tr>
<td>Hollingsworth, W.</td>
</tr>
<tr>
<td>Hope, L.</td>
</tr>
<tr>
<td>Horn, S.</td>
</tr>
<tr>
<td>Horowitz, A.</td>
</tr>
<tr>
<td>Hosteller, R.</td>
</tr>
<tr>
<td>Hots, G.</td>
</tr>
<tr>
<td>House, M.</td>
</tr>
<tr>
<td>House, V.</td>
</tr>
<tr>
<td>Howe, M.</td>
</tr>
<tr>
<td>Hubbard, W.</td>
</tr>
<tr>
<td>Hubby, J.</td>
</tr>
<tr>
<td>Hudson, J.</td>
</tr>
<tr>
<td>Hughes, R.</td>
</tr>
<tr>
<td>Hungerford, D.</td>
</tr>
<tr>
<td>Hunter, A.</td>
</tr>
<tr>
<td>Hurwitz, D.</td>
</tr>
<tr>
<td>Ibrahim, S.</td>
</tr>
<tr>
<td>Ichida, H.</td>
</tr>
<tr>
<td>Ichikawa, S.</td>
</tr>
<tr>
<td>Ikeda, H.</td>
</tr>
<tr>
<td>Imai, Y.</td>
</tr>
<tr>
<td>Imaizumi, T.</td>
</tr>
<tr>
<td>Inagaki, E.</td>
</tr>
<tr>
<td>Inouye, I.</td>
</tr>
<tr>
<td>Ives, P.</td>
</tr>
<tr>
<td>Iyama, S.</td>
</tr>
<tr>
<td>Iyengar, S.</td>
</tr>
<tr>
<td>Jacobs, A.</td>
</tr>
<tr>
<td>Jacobs, M.</td>
</tr>
<tr>
<td>James, J.</td>
</tr>
<tr>
<td>Janke, H.</td>
</tr>
<tr>
<td>Jayakar, S.</td>
</tr>
<tr>
<td>Jemson, J.</td>
</tr>
<tr>
<td>Johansen, I.</td>
</tr>
<tr>
<td>Johansson, K.</td>
</tr>
<tr>
<td>Johns, R.</td>
</tr>
<tr>
<td>Johnsen, R.</td>
</tr>
<tr>
<td>Johnson, G.</td>
</tr>
<tr>
<td>Johnson, W.</td>
</tr>
<tr>
<td>Jonselit, C.</td>
</tr>
<tr>
<td>Jonsson, U.</td>
</tr>
<tr>
<td>Judd, B.</td>
</tr>
<tr>
<td>Jur, C.</td>
</tr>
<tr>
<td>Kaji, S.</td>
</tr>
<tr>
<td>Kaminishi, H.</td>
</tr>
<tr>
<td>Kanesio, T.</td>
</tr>
<tr>
<td>Kaneko, A.</td>
</tr>
<tr>
<td>Kang, S.</td>
</tr>
<tr>
<td>Kang, Y.</td>
</tr>
<tr>
<td>Kapetan, A.</td>
</tr>
<tr>
<td>Kaplan, W.</td>
</tr>
<tr>
<td>Karaz, J.</td>
</tr>
<tr>
<td>Karlik, A.</td>
</tr>
<tr>
<td>Kasai, I.</td>
</tr>
<tr>
<td>Kato, M.</td>
</tr>
<tr>
<td>Kato, Masaru, J.</td>
</tr>
<tr>
<td>Kato, S.</td>
</tr>
<tr>
<td>Kaufman, E.</td>
</tr>
<tr>
<td>Kaufmann, B.</td>
</tr>
<tr>
<td>Kawale, M.</td>
</tr>
<tr>
<td>Kaye, R.</td>
</tr>
<tr>
<td>Keller, E.</td>
</tr>
<tr>
<td>Kelsall, P.</td>
</tr>
<tr>
<td>Kernaghan, R.</td>
</tr>
<tr>
<td>Kessler, S.</td>
</tr>
<tr>
<td>Khairalla, A.</td>
</tr>
<tr>
<td>Khishin, A.</td>
</tr>
<tr>
<td>Kieiselbach, T.</td>
</tr>
<tr>
<td>Kiger, J.</td>
</tr>
<tr>
<td>Kiil, T.</td>
</tr>
<tr>
<td>Kikkawa, H.</td>
</tr>
<tr>
<td>Kim, D.S.</td>
</tr>
<tr>
<td>Kim, D. V.</td>
</tr>
<tr>
<td>Kim, K.</td>
</tr>
<tr>
<td>Kim, C.</td>
</tr>
<tr>
<td>Kimura, M.</td>
</tr>
<tr>
<td>King, Jack</td>
</tr>
<tr>
<td>King, James, N.</td>
</tr>
<tr>
<td>King, R.</td>
</tr>
<tr>
<td>Kirshbaum, W.</td>
</tr>
<tr>
<td>Kirchofien, G.</td>
</tr>
<tr>
<td>Kitagawa, C.</td>
</tr>
<tr>
<td>Last Name</td>
</tr>
<tr>
<td>-----------</td>
</tr>
<tr>
<td>Kitazumi</td>
</tr>
<tr>
<td>Kloos</td>
</tr>
<tr>
<td>Knight</td>
</tr>
<tr>
<td>Knight</td>
</tr>
<tr>
<td>Knott</td>
</tr>
<tr>
<td>Koch</td>
</tr>
<tr>
<td>Koch</td>
</tr>
<tr>
<td>Khulein</td>
</tr>
<tr>
<td>Kojima</td>
</tr>
<tr>
<td>Kornarski</td>
</tr>
<tr>
<td>Kondo</td>
</tr>
<tr>
<td>Koref-Santibanez</td>
</tr>
<tr>
<td>Kosak-Westphal</td>
</tr>
<tr>
<td>Kosswig</td>
</tr>
<tr>
<td>Krause</td>
</tr>
<tr>
<td>Krauss</td>
</tr>
<tr>
<td>Kraus</td>
</tr>
<tr>
<td>Krawinkel</td>
</tr>
<tr>
<td>Krebs</td>
</tr>
<tr>
<td>Kribsenke</td>
</tr>
<tr>
<td>Kroeger</td>
</tr>
<tr>
<td>Kromm</td>
</tr>
<tr>
<td>Kyvland</td>
</tr>
<tr>
<td>Kinkel</td>
</tr>
<tr>
<td>Kunze-Mühl</td>
</tr>
<tr>
<td>Kuroda</td>
</tr>
<tr>
<td>Kurokawa</td>
</tr>
<tr>
<td>Kyle</td>
</tr>
<tr>
<td>Lagowski</td>
</tr>
<tr>
<td>Laird</td>
</tr>
<tr>
<td>Lakovara</td>
</tr>
<tr>
<td>Lamb</td>
</tr>
<tr>
<td>Lamerton</td>
</tr>
<tr>
<td>Landenberger</td>
</tr>
<tr>
<td>Landy</td>
</tr>
<tr>
<td>Langer</td>
</tr>
<tr>
<td>Leaqué</td>
</tr>
<tr>
<td>Lauffer</td>
</tr>
<tr>
<td>Laureys</td>
</tr>
<tr>
<td>Lawrence</td>
</tr>
<tr>
<td>Leaes</td>
</tr>
<tr>
<td>Le Blang</td>
</tr>
<tr>
<td>Lederman-Klein</td>
</tr>
<tr>
<td>Lee</td>
</tr>
<tr>
<td>Lee</td>
</tr>
<tr>
<td>Lee</td>
</tr>
<tr>
<td>Lafaye</td>
</tr>
<tr>
<td>Legay</td>
</tr>
<tr>
<td>Leigh</td>
</tr>
<tr>
<td>Leigh</td>
</tr>
<tr>
<td>Iacon</td>
</tr>
<tr>
<td>Leon</td>
</tr>
<tr>
<td>Lesher</td>
</tr>
<tr>
<td>Lestrangé</td>
</tr>
<tr>
<td>Leventhal</td>
</tr>
<tr>
<td>LeVier</td>
</tr>
<tr>
<td>Levine</td>
</tr>
<tr>
<td>Levine</td>
</tr>
<tr>
<td>Levine</td>
</tr>
<tr>
<td>Levitin</td>
</tr>
<tr>
<td>Lewgoy</td>
</tr>
<tr>
<td>Lewis</td>
</tr>
<tr>
<td>Lewis</td>
</tr>
<tr>
<td>Lewis</td>
</tr>
<tr>
<td>Lewontin</td>
</tr>
<tr>
<td>Lepzi</td>
</tr>
<tr>
<td>L'Héritier</td>
</tr>
<tr>
<td>Lichtwardt</td>
</tr>
<tr>
<td>Lindsley</td>
</tr>
<tr>
<td>Locatelli</td>
</tr>
<tr>
<td>Löfler</td>
</tr>
<tr>
<td>Lommers</td>
</tr>
<tr>
<td>Loomis</td>
</tr>
<tr>
<td>Louis</td>
</tr>
<tr>
<td>Lovellette</td>
</tr>
<tr>
<td>Lucchesi</td>
</tr>
<tr>
<td>Luce</td>
</tr>
<tr>
<td>Luchowski</td>
</tr>
<tr>
<td>Ludwig</td>
</tr>
<tr>
<td>Ludwig</td>
</tr>
<tr>
<td>Ludwig</td>
</tr>
<tr>
<td>Liuers</td>
</tr>
<tr>
<td>Liuers</td>
</tr>
<tr>
<td>Lund</td>
</tr>
<tr>
<td>Lund</td>
</tr>
<tr>
<td>Lüning</td>
</tr>
<tr>
<td>Lux</td>
</tr>
<tr>
<td>McClarahan</td>
</tr>
<tr>
<td>McCullough</td>
</tr>
<tr>
<td>McCune</td>
</tr>
<tr>
<td>McDowell</td>
</tr>
<tr>
<td>McKean</td>
</tr>
<tr>
<td>McKinley</td>
</tr>
<tr>
<td>McLean</td>
</tr>
<tr>
<td>McSheehy</td>
</tr>
<tr>
<td>Maclay</td>
</tr>
<tr>
<td>Magda</td>
</tr>
<tr>
<td>Magalhaes</td>
</tr>
<tr>
<td>Magaribuchi</td>
</tr>
<tr>
<td>Magni</td>
</tr>
<tr>
<td>Mahler</td>
</tr>
<tr>
<td>Mahowald</td>
</tr>
<tr>
<td>Mainz</td>
</tr>
<tr>
<td>Makino</td>
</tr>
<tr>
<td>Mallah</td>
</tr>
<tr>
<td>Malogolowkin</td>
</tr>
<tr>
<td>Markowitz</td>
</tr>
<tr>
<td>Marques</td>
</tr>
<tr>
<td>Martin</td>
</tr>
<tr>
<td>Martin</td>
</tr>
<tr>
<td>Maruyama</td>
</tr>
<tr>
<td>Marzluft</td>
</tr>
<tr>
<td>Massamallo</td>
</tr>
<tr>
<td>Masterson</td>
</tr>
<tr>
<td>Masuda</td>
</tr>
<tr>
<td>Mother</td>
</tr>
<tr>
<td>Mother</td>
</tr>
<tr>
<td>Mathew</td>
</tr>
<tr>
<td>Matos</td>
</tr>
<tr>
<td>Mattson</td>
</tr>
<tr>
<td>Maxim</td>
</tr>
<tr>
<td>Mayhew</td>
</tr>
<tr>
<td>Maynard Smith</td>
</tr>
<tr>
<td>Mayo</td>
</tr>
<tr>
<td>Mazur-Barnett</td>
</tr>
<tr>
<td>Need</td>
</tr>
<tr>
<td>Medina</td>
</tr>
<tr>
<td>Melov</td>
</tr>
<tr>
<td>Mercier</td>
</tr>
</tbody>
</table>
January 1962  Directory - Alphabetical  36:175

Merrell, D. see DIS 34:180
Mettler, L. Raleigh, North Carolina
Meyer, G. Germany, Tübingen
Meyer, H. Bloomington, Indiana
Michell, A. Italy, Rome
Mickel, S. Eugene, Oregon
Mickey, G. Ridgefield, Connecticut
Mileiko, V. Canada, Toronto
Mills, Richard Evanston, Illinois
Mills, R.P. New Haven, Connecticut
Mincari, A. Japan, Misima
Miralles, L. Spain, Madrid
Mishara, J. New York, New York
Mitchell, A. Pasadena, California
Mitchell, H. Pasadena, California
Mitra, J. see DIS 34:133
Mittler, S. Dekalb, Illinois
Miyoshi, Y. Japan, Tokyo
Monawad, H. United Arab Rep., Alexandria
Mohler, J. Corvallis, Oregon
Mohr, O. Norway, Blindern
Momaya, M. Japan, Sapporo
Monclús, M. Spain, Barcelona
Montalenti, G. Italy, Rome
Montelius, I. Sweden, Stockholm
Montgomery, R. see DIS 34:147
Moore, R. Chapel Hill, North Carolina
More, S. Pasadena, California
Morales, N. Brazil, Porto Alegre
Morre, R. Pullman, Washington
Mrey, M. Spain, Madrid
Morikawa, D. Japan, Tokyo
Moskvinwski, T. Notre Dame, Indiana
Mossige, J. Norway, Oslo
Mostafa, A. Gr. Britain, Edinburgh
Mourad, A. United Arab Rep., Alexandria
Mouravieff, A. see DIS 34:127
Moyer, S. see DIS 34:150
Moynehan, J. see DIS 34:148
Mukai, T. Japan, Misima
Mukherjee, A. Berkeley, California
Mukherjee, B. Salt Lake City, Utah
Mukherjee, M. see DIS 34:133
Müller, A. Germany, Karlsruhe
Müller, H. J. Bloomington, Indiana
Müller, M. Switzerland, Zürich
Munz, C. Brazil, Porto Alegre
Munoz, E. Argentina, Buenos Aires
Munoz, P. Switzerland, Zürich
Myers, O. Ithaca, New York
Wyszomrs, W. East Lansing, Michigan
Nadad, A. Spain, Barcelona
Nafei, H. Gr. Britain, Edinburgh
Nakamura, K. Japan, Misima
Nakamura, Kenji Japan, Kyoto
Nakao, Y. see DIS 34:135
Napp, M. Brazil, Porto Alegre
Narain, P. India, Delhi
Narayana Rao, N. India, Madras
Narise, T. Japan, Misima
Nash, D. Gr. Britain, Cambridge
Nash, Donald J. University Park, Penn.

Nama, S. Japan, Misima
Neeley, J. Corvallis, Oregon
Net, P. South Africa, Pretoria
Noulat, M. France, Lyon
Nowball, S. Colombia, Bogotá
Nicklas, R. New Haven, Connecticut
Nicoletti, B. Italy, Rome
Nigon, V. France, Lyon
Nobuki, R. Japan, Osaka
Nold, D. S. Africa, Johannesburg
Nomura, K. Salt Lake City, Utah
Norwood, S. Austin, Texas
Nöthel, H. Germany, Berlin-Dahlem
Nöthiger, R. Switzerland, Zürich
Novitski, E. Eugene, Oregon
Nozawa, K. Japan, Anzyo-Shi
Nugent, K. East Lansing, Michigan

Oftedal, P. Norway, Oslo
Ogaki, M. Japan, Sakai
Ogita, Z. Japan, Osaka
Ogawa, T. Canada, Vancouver
Ohanessian-Chilemain, A. France, Gif-sur-Yvette
Ohba, S. Japan, Tokyo
Ohlendorff, N. Sweden, Uppsala
Oishi, E. Japan, Tokyo
Okada, T. Japan, Tokyo
Okazaki, T. Japan, Tokyo
Okeke, M. Japan, Tokyo
Okuda, C. Japan, Kyoto
Olivieri, G. Italy, Rome
Olivieri, M. Italy, Rome
Olson, J. Urbana, Illinois
O'Neal, L. see DIS 34:150
Orillard, C. see DIS 34:129
Orde, J. Brazil, São Paulo
Ortiz, E. Spain, Madrid
Oshima, C. Japan, Misima
Oster, J. Bloomington, Indiana

Ostertag, W. Germany, Münster
Ota, N. Japan, Anzyo-Shi
Otujy, Y. Japan, Osaka

Paek, Y. Korea, Seoul
Pakeltis, H. Evanston, Illinois
Paolini, F. see DIS 34:140
Paro, J. Lafayette, Indiana
Parish, J. Chapel Hill, N. Carolina
Park, M. Korea, Kwangju
Parker, Dean see DIS 34:153
Parker, D. M. Eugene, Oregon
Parsons, P. Gr. Britain, Cambridge
Parson, S. East Lansing, Michigan
Passman, K. New Haven, Connecticut
Pasternak, L. Germany, Berlin-Buch
Paterson, H. see DIS 34:139
Paul, L. Salt Lake City, Utah
Pavan, C. Brazil, São Paulo
Pavlovsky, O. New York, New York
Paz, R. Argentina, Buenos Aires
Peat, W. see DIS 34:133
Pelecosos, M. Gr. Britain, Cambridge
Pellicer, M. Chile, Santiago
Perdrix, S. France, Lyon
Perry, M. Gr. Britain, Edinburgh
Perry, T. Johnson City, Tennessee
Pesci, J., Italy
Peterson, W. Chicago, Illinois
Peterson, K. Syracuse, New York
Petit, C. see DIS 34:129
Pettijohn, S. Africa, Johannesburg
Phillips, B. Los Angeles, California
Phillips, C. Los Angeles, California
Phillips, D. Syracuse, New York
Pila, M. Salt Lake City, Utah
Pilgrim, S. Africa, Johannesburg
Pillman, L. France, Gif-sur-Yvette
Piluso, A. France, Gif-sur-Yvette
Plagne, H. see DIS 34:145
Plough, H. D. Cambridge, Massachusetts
Plus, N. France, Gif-sur-Yvette
Pollock, R. see DIS 34:160
Polzin, W. Germany, Berlin-Dahlem
Pons, J. Spain, Barcelona
Ponte Corvo, G. Gr. Britain, Glasgow
Pooley, M. see DIS 34:152
Pollack, R. see DIS 34:150
Polzin, W. Germany, Berlin-Dahlem
Ponente, D. Brazil, São Paulo
Polivanov, S. New York, New York
Pollack, R. see DIS 34:150
Pons, J. Spain, Barcelona
Polivanov, S. New York, New York
Pollet, D. Brazil, São Paulo
Poli, V. Italy, Milano
Pratt, G. Gr. Britain, Edinburgh
Pratt, Cathie, Oak Ridge, Tennessee
Prestridge, M. Baton Rouge, La.
Prevosti, A. Spain, Barcelona
Price, M. Chapel Hill, North Carolina
Prokop, B. Cold Spring Harbor, N.Y.
Proust, J. France, Gif-sur-Yvette
Proust, J. France, Gif-sur-Yvette
Proux, A. France, Gif-sur-Yvette
Queiroz, J. France, Gif-sur-Yvette
Quermer, W. Germany, Mariensee
Rahat, A. Israel, Jerusalem
Rai, K. Notre Dame, Indiana
Rakha, F. United Arab Rep., Alexandria
Rasmussen, C. Sweden, Stockholm
Ramial, D. Brazil, Porto Alegre
Ramirez, P. Spain, Madrid
Ranzheimer, S. Ann Arbor, Michigan
Rao, H. India, Hyderabad
Rappaport, S. Israel, Jerusalem
Rasmussen, E. Sweden, Upsala
Rasmussen, M. Sweden, Upsala
Rasmussen, I. see DIS 34:154
Ratty, F. San Diego, California
Rausani, C. Germany, Berlin-Dahlem
Ray-Chaudhuri, S. see DIS 34:133
Reddi, C. India, Hyderabad
Reeve, E. Gr. Britain, Edinburgh
Regaly, M. Brazil, Porto Alegre
Reuten, P. Staten Island, New York
Hendel, J. Australia, Sydney
Resch, K. Austin, Texas
Reszonski, Raimondi, G. Italy, Milano
Richardson, R. Raleigh, North Carolina
Ridgway, T. see DIS 34:163
Riles, L. see DIS 34:163
Rinehart, R. Austin, Texas
Rinkel, R. see DIS 34:149
Ritte, C. Israel, Jerusalem
Rios, C. Colombia, Bogótá
Ritossa, F. see DIS 34:134
Ritterhoff, R. Baltimore, Maryland
Riski, R. Ann Arbor, Michigan
Riski, T. Ann Arbor, Michigan
Roberts, P. Chicago, Illinois
Robertson, A. Gr. Britain, Edinburgh
Robertson, F. Gr. Britain, Edinburgh
Roderick, T. see DIS 34:143
Rodriguez, C. Spain, Madrid
Rohlf, F. see DIS 34:149
Rühborn, G. Germany, Berlin-Dahlem
Roisemberg, I. Brazil, Porto Alegre
Ronen, A. Israel, Jerusalem
Roper, J. Gr. Britain, Sheffield
Rosenfeld, A. Madison, Wisconsin
Rosen, S. Switzerland, Bern
Rosen, J. New Haven, Connecticut
Royo, V. Gr. Britain, Edinburgh
Rudio, C. Colombia, Bogótá
Ruderer, E. Austria, Vienna
Rudkin, G. see DIS 34:162
Rudolph, E. Germany, Berlin-Dahlem
Ruske, C. Netherlands, Utrecht
Russell, J. see DIS 34:154
Russell, P. Gr. Britain, Edinburgh
Rutter, J. Madison, Wisconsin
Russell, E. Austria, Vienna
Saarinen, L. see DIS 34:152
Sakaguchi, B. Japan, Misima
Sakai, K. Japan, Misima
Sakitey, A. Ghana, Legon
Sakurai, W. see DIS 34:142
Salzano, F. Brazil, Porto Alegre
Sammalisto, L. Finland, Helsinki
Sanderson, K. Ithaca, New York
Sanderson, K. Ithaca, New York
Sang, J. Gr. Britain, Edinburgh
Sankaranayanan, K. New York, New York
Santos, A. Brazil, Porto Alegre
Sarkar, S. see DIS 34:133
Savage, E. Alliance, Ohio
Savhagen, R. Sweden, Stockholm
Savolainen, S. Finland, Helsinki
Scheid, H. Oak Ridge, Oak Ridge
Schafer, H. Raleigh, North Carolina
Schafrir, C. Ithaca, New York
Schael, A. Cold Spring Harbor, N.Y.
Schario, W. Gr. Britain, Edinburgh
Schepers, A. Netherlands, Leiden
Scheren, A. Philadelphia, Penn.
Schindler, D. Lafayette, Indiana
Schiller, R. Philadelphia, Penn.
Schlagel, G. see DIS 34:149
Schlappfer, T. Switzerland, Zürich
Schmidt, W. Austin, Texas
Schneider, A. Switzerland, Zürich
Schneider, A. Switzerland, Zürich
Schneider, I. Switzerland, Zurich
Schotten, S. Netherlands, Utrecht
Schubert, G. see DIS 34:130
Schuelen, R. see DIS 34:146
Schultz, J. see DIS 34:152
Schwarz, R. see DIS 34:152
Schweda, M. see DIS 34:149
Schwilck, I. Germany, Mariensee
Schulten, G. Netherlands, Leiden
Scossiroli, R. see DIS 34:134
Scriba, M. Germany, Marburg/Lahn
Seaton, R. East Lansing, Michigan
Sederoft, R. Los Angeles, California
Seecof, R. Pasadena, California
Seidel, F. Germany, Marburg/Lahn
Seidel, S. Germany, Tübingen
Seiger, M. Canada, Toronto
Seki, T. Japan, Osaka
Semp, E. Lexington, Kentucky
Sen, B. Gr. Britain, Edinburgh
Seto, F. Berea, Kentucky
Shapard, P. see DIS 34:155
Shapiro, N. see DIS 34:148
Sharlot, W. Netherlands, Leiden
Sheldon, E. Australia, Sydney
Sheridan, B. Australia, Sydney
Sherwood, E. Berkeley, California
Shidel, D. Lafayette, Indiana
Shima, T. Japan, Sapporo
Shiomi, M. see DIS 34:135
Shoeb, Y. United Arab Rep., Alexandria
Shoji, T. Japan, Mitaka
Sick, K. see DIS 34:123
Sieux, J. Chicago, Illinois
Silberman, J. Ithaca, New York
Silva, L. Brazil, Porto Alegre
Silva, T. Brazil, Porto Alegre
Simmons, J. Logan, Utah
Simões, G. Brazil, Porto Alegre
Singer, K. see DIS 34:144
Singleton, J. see DIS 34:148
Simo, Y. Japan, Mitaka
Simonesano, V. Jamaica, New York
Sironi, G. Italy, Milano
Slatia, H. see DIS 34:142
Silzynski, B. Gr. Britain, Edinburgh
Silzynski, H. Gr. Britain, Edinburgh
Smith, E. Norway, Blindern
Smith, E. Norway, Blindern
Smith, F. Evanston, Illinois
Smith, S. Corvallis, Oregon
Snyder, L. Gr. Britain, Edinburgh
Sobels, F. Netherlands, Leiden
Sokal, A. see DIS 34:149
Sokoloff, A. Berkeley, California
Solana, I. Spain, Madrid
Solima, A. New York, New York
Soliman, G. United Arab Rep., Alexandria
Soliman, M. United Arab Rep., Alexandria
Sollum, F. Norway, Blindern
Sondhi, G. Ridgefield, Connecticut
Sondhi, K. Ridgefield, Connecticut
Sonnenblick, B. see DIS 34:150
Sorensen, W. Logan, Utah
Southin, J. Los Angeles, California
Spassky, B. New York, New York
Sperlich, D. Austria, Vienna
Spickett, S. Gr. Britain, Cambridge
Spieler, R. Chicago, Illinois
Spies, E. Pittsburgh, Pennsylvania
Spiess, L. Pittsburgh, Pennsylvania
Spielth, J. see DIS 34:153
Spofford, J. Chicago, Illinois
Springer, R. Austria, Vienna
Spurway, H. see DIS 34:133
Spurway, R. Australia, Brisbane
Stadler, J. Ames, Iowa
Stalker, H. see DIS 34:153
Stanich, G. East Lansing, Michigan
Steffenson, D. see DIS 34:155
Steenberg, A. see DIS 34:145
Steinitz, J. Salt Lake City, Utah
Stern, Curt Berkeley, California
Stevenson, R. Johnson City, Tennessee
Stewart, W. Lexington, Kentucky
Stiers, R. see DIS 34:127
Stone, L. Lincoln, Nebraska
Stone, W. Austin, Texas
Strangio, V. Australia, Melbourne
Strachan, I. Gr. Britain, Edinburgh
Strachan, K. Gr. Britain, Edinburgh
Stratopoulos, G. Salt Lake City, Utah
Strickberger, M. New York, New York
Strömmann, O. Norway, Blindern
Struck, E. Germany, Berlin-Dahlem
Sturtevant, A. Pasadema, California
Suguna, S. India, Madras
Sulerud, R. Lincoln, Nebraska
Sullivan, R. see DIS 34:149
Sung, K. Korea, Seoul
Sumanalainen, E. Finland, Helsinki
Suiza, J. Ithaca, New York
Suzuki, L. Oak Ridge, Tennessee
Svensson, M. Sweden, Uppsala
Sweet, E. Pittsburgh, Pennsylvania
Sworon, M. Gr. Britain, Chalfont
Tabata, K. Canada, Vancouver
Tachibana, H. see DIS 34:135
Taira, T. Japan, Misima
Takada, H. Japan, Sapporo
Takaya, H. Japan, Okamoto
Takken, I. Canada, Toronto
Talsma, J. Cold Spring Harbor, New York
Tanaka, E. Japan, Sakai
Tanaka, T. Duarte, California
Tano, S. see DIS 34:148
Tanawy, A. United Arab Rep., Alexandria
Tate, A. Netherland, Leiden
Taylor, A. Cold Spring Harbor, New York
Teissier, G. France, Gif-sur-Yvette
Telleman, A. see DIS 34:150
Teviotdale, F. Eugene, Oregon
Thady, G. Brazil, Porto Alegre
Thoday, J. Gr. Britain, Cambridge
Thomas, Carol Madison, Wisconsin
Thomas, Constance, Salt Lake City, Utah
Thomas, S. Bloomington, Indiana
Thompson, M. Ithaca, New York
Thompson, P. Ames, Iowa
<table>
<thead>
<tr>
<th>Name</th>
<th>Country</th>
<th>City</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thompson, S.</td>
<td>Australia, Oregon</td>
<td>Corvallis</td>
</tr>
<tr>
<td>Thomson, J.</td>
<td>Australia, Melbourne</td>
<td></td>
</tr>
<tr>
<td>Throckmorton, L.</td>
<td>Chicago, Illinois</td>
<td></td>
</tr>
<tr>
<td>Tiffany, B.</td>
<td>Massachusetts</td>
<td>Amherst</td>
</tr>
<tr>
<td>Tiivola, A.</td>
<td>Finland</td>
<td>Helsinki</td>
</tr>
<tr>
<td>Tobari, I.</td>
<td>Japan, Tokyo</td>
<td></td>
</tr>
<tr>
<td>Tobari, Y.</td>
<td>Japan, Tokyo</td>
<td></td>
</tr>
<tr>
<td>Tokunaga, C.</td>
<td>Berkeley, California</td>
<td></td>
</tr>
<tr>
<td>Toledo, J.</td>
<td>Brazil, São Paulo</td>
<td></td>
</tr>
<tr>
<td>Toledo, S.</td>
<td>Brazil, São Paulo</td>
<td></td>
</tr>
<tr>
<td>Tondo, G.</td>
<td>Brazil, São Paulo</td>
<td>Porto Alegre</td>
</tr>
<tr>
<td>Torroja, E.</td>
<td>Spain, Madrid</td>
<td></td>
</tr>
<tr>
<td>Townsend, J.</td>
<td>Virginia, Richmond</td>
<td></td>
</tr>
<tr>
<td>Trout, H.</td>
<td>Germany, Karlsruhe</td>
<td></td>
</tr>
<tr>
<td>Travaglini, E.</td>
<td></td>
<td>see DIS 34:152</td>
</tr>
<tr>
<td>Tresman, K.</td>
<td>New York</td>
<td>Buffalo</td>
</tr>
<tr>
<td>Trogildo, D.</td>
<td>Brazil, São Paulo</td>
<td></td>
</tr>
<tr>
<td>Trosko, J.</td>
<td>Michigan, East Lansing</td>
<td></td>
</tr>
<tr>
<td>Trout, W.</td>
<td>Indiana, Bloomington</td>
<td></td>
</tr>
<tr>
<td>Tschumi, P.</td>
<td>Switzerland, Bern</td>
<td></td>
</tr>
<tr>
<td>Tsukamoto, H.</td>
<td>Japan, Tokyo</td>
<td></td>
</tr>
<tr>
<td>Tsukamoto, M.</td>
<td>Japan, Osaka</td>
<td></td>
</tr>
<tr>
<td>Twinnstra, E.</td>
<td>Netherlands, Utrecht</td>
<td></td>
</tr>
<tr>
<td>Twisten, M.</td>
<td></td>
<td>see DIS 34:152</td>
</tr>
<tr>
<td>Ufholz, I.</td>
<td>Germany, Karlsruhe</td>
<td></td>
</tr>
<tr>
<td>Ulrich, H.</td>
<td>Switzerland, Zürich</td>
<td></td>
</tr>
<tr>
<td>Ursprung, H.</td>
<td>Maryland, Baltimore</td>
<td></td>
</tr>
<tr>
<td>Uscati, J.</td>
<td></td>
<td>see DIS 34:129</td>
</tr>
<tr>
<td>Valencia, J.</td>
<td>Argentina, Buenos Aires</td>
<td></td>
</tr>
<tr>
<td>Valencia, R.</td>
<td>Argentina, Buenos Aires</td>
<td></td>
</tr>
<tr>
<td>Van Hooff, J.</td>
<td>South Africa, Pretoria</td>
<td>Netherlands, Leiden</td>
</tr>
<tr>
<td>Van Niekerk, B.</td>
<td>South Africa, Pretoria</td>
<td></td>
</tr>
<tr>
<td>Van Schaik, N.</td>
<td>South Africa, Pretoria</td>
<td></td>
</tr>
<tr>
<td>Van Valen, L.</td>
<td>New York, New York</td>
<td></td>
</tr>
<tr>
<td>Vigier, F.</td>
<td>France, Gif-sur-Yvette</td>
<td></td>
</tr>
<tr>
<td>Virzo, A.</td>
<td></td>
<td>see DIS 34:134</td>
</tr>
<tr>
<td>Vloeberghs, J.</td>
<td></td>
<td>see DIS 34:127</td>
</tr>
<tr>
<td>Volkerst, W.</td>
<td>Michigan, East Lansing</td>
<td>Netherlands, Leiden</td>
</tr>
<tr>
<td>von Borstel, R.</td>
<td>Tennessee, Oak Ridge</td>
<td></td>
</tr>
<tr>
<td>von Halle, R.</td>
<td>Tennessee, Oak Ridge</td>
<td></td>
</tr>
<tr>
<td>Voynow, N.</td>
<td>Wisconsin, Madison</td>
<td></td>
</tr>
<tr>
<td>Waddington, C.</td>
<td>Britain, Edinburgh</td>
<td>Gr, Oxford</td>
</tr>
<tr>
<td>Wagner, R.</td>
<td>Texas, Austin</td>
<td></td>
</tr>
<tr>
<td>Wagoner, D.</td>
<td>Indiana, Bloomington</td>
<td></td>
</tr>
<tr>
<td>Wajntal, A.</td>
<td>Brazil, São Paulo</td>
<td></td>
</tr>
<tr>
<td>Wale, K.</td>
<td>California, Berkeley</td>
<td></td>
</tr>
<tr>
<td>Wall, L.</td>
<td>North Carolina, Chapel Hill</td>
<td></td>
</tr>
<tr>
<td>Wallace, E.</td>
<td>New York</td>
<td>Ithaca</td>
</tr>
<tr>
<td>Wallbrum, C.</td>
<td>Florida, Gainesville</td>
<td></td>
</tr>
<tr>
<td>Wallenda, M.</td>
<td>Turkia, Finland</td>
<td></td>
</tr>
<tr>
<td>Wang, S.</td>
<td></td>
<td>see DIS 34:145</td>
</tr>
<tr>
<td>Ward, C.</td>
<td>North Carolina, Durham</td>
<td></td>
</tr>
<tr>
<td>Wasserman, M.</td>
<td>Australia, Melbourne</td>
<td></td>
</tr>
<tr>
<td>Watanabe, H.</td>
<td>Japan, Osaka</td>
<td></td>
</tr>
<tr>
<td>Wettlaufer, J.</td>
<td></td>
<td>see DIS 34:127</td>
</tr>
<tr>
<td>Wedvik, H.</td>
<td>Norway, Blindern</td>
<td></td>
</tr>
<tr>
<td>Wei, L.</td>
<td>Illinois, Dekalb</td>
<td></td>
</tr>
<tr>
<td>Weideman, J.</td>
<td>Illinois, Dekalb</td>
<td></td>
</tr>
<tr>
<td>Weingart, E.</td>
<td>New York, Cold Springs Harbor</td>
<td>N.Y.</td>
</tr>
<tr>
<td>Weinmann, H.</td>
<td>Zürich, Switzerland</td>
<td></td>
</tr>
<tr>
<td>Weisbrod, D.</td>
<td>New York, New York</td>
<td></td>
</tr>
<tr>
<td>Welch, R.</td>
<td>Texas, Austin</td>
<td></td>
</tr>
<tr>
<td>Welshons, W.</td>
<td>Tennessee, Oak Ridge</td>
<td></td>
</tr>
<tr>
<td>Westergaard, M.</td>
<td></td>
<td>see DIS 34:128</td>
</tr>
<tr>
<td>Wette, R.</td>
<td></td>
<td>see DIS 34:150</td>
</tr>
<tr>
<td>Wharton, M.</td>
<td>Texas, Austin</td>
<td></td>
</tr>
<tr>
<td>Wheeler, M.</td>
<td>Texas, Austin</td>
<td></td>
</tr>
<tr>
<td>White, R.</td>
<td></td>
<td>see DIS 34:147</td>
</tr>
<tr>
<td>White, S.</td>
<td>Pennsylvania, Philadelphia</td>
<td></td>
</tr>
<tr>
<td>Whitesel, B.</td>
<td>California, Los Angeles</td>
<td></td>
</tr>
<tr>
<td>Whittinghill, M.</td>
<td>chapel Hill, North Carolina, USA</td>
<td></td>
</tr>
<tr>
<td>Whittington, N.</td>
<td></td>
<td>see DIS 34:153</td>
</tr>
<tr>
<td>Wilhelm, H.</td>
<td></td>
<td>see DIS 34:152</td>
</tr>
<tr>
<td>Wilkerson, R.</td>
<td>Tennessee, Oak Ridge</td>
<td></td>
</tr>
<tr>
<td>Wells, G.</td>
<td>Canada, Vancouver</td>
<td></td>
</tr>
<tr>
<td>Williams, T.</td>
<td></td>
<td>see DIS 34:153</td>
</tr>
<tr>
<td>Williamson, D.</td>
<td>Connecticut, New Haven</td>
<td></td>
</tr>
<tr>
<td>Wilson, F.</td>
<td>Texas, Austin</td>
<td></td>
</tr>
<tr>
<td>Wilson, L.</td>
<td>Massachusetts, Wellesley</td>
<td></td>
</tr>
<tr>
<td>Wing, W.</td>
<td>North Carolina, Raleigh</td>
<td></td>
</tr>
<tr>
<td>Winge, H.</td>
<td>Brazil, São Paulo</td>
<td></td>
</tr>
<tr>
<td>Winterfeldt, G.</td>
<td>Germany, Berlin-Dahlem</td>
<td></td>
</tr>
<tr>
<td>Wolf, E.</td>
<td>Germany, Berlin-Dahlem</td>
<td></td>
</tr>
<tr>
<td>Wolf, J.</td>
<td></td>
<td>see DIS 34:150</td>
</tr>
<tr>
<td>Wolff, S.</td>
<td>Tennessee, Oak Ridge</td>
<td></td>
</tr>
<tr>
<td>Woods, P.</td>
<td></td>
<td>see DIS 34:155</td>
</tr>
<tr>
<td>Wrathall, C.</td>
<td>Utah, Salt Lake City</td>
<td></td>
</tr>
<tr>
<td>Wright, E.</td>
<td>Maryland, Baltimore</td>
<td></td>
</tr>
<tr>
<td>Wright, T.</td>
<td>Maryland, Baltimore</td>
<td></td>
</tr>
<tr>
<td>Wu, C.</td>
<td>Illinois, Dekalb</td>
<td></td>
</tr>
<tr>
<td>Wui, I.</td>
<td>Korea, Kwangju</td>
<td></td>
</tr>
<tr>
<td>Würgler, F.</td>
<td>Switzerland, Zürich</td>
<td></td>
</tr>
<tr>
<td>Wylie, C.</td>
<td></td>
<td>see DIS 34:148</td>
</tr>
<tr>
<td>Yamada, Y.</td>
<td>Indiana, Lafayette</td>
<td></td>
</tr>
<tr>
<td>Yanders, A.</td>
<td>Michigan, East Lansing</td>
<td></td>
</tr>
<tr>
<td>Yasuda, N.</td>
<td></td>
<td>see DIS 34:135</td>
</tr>
<tr>
<td>Yeatts, V.</td>
<td></td>
<td>see DIS 34:146</td>
</tr>
<tr>
<td>Yen, T.</td>
<td>North Carolina, Chapel Hill</td>
<td></td>
</tr>
<tr>
<td>Yoo, C.</td>
<td>Korea, Seoul</td>
<td></td>
</tr>
<tr>
<td>Yoon, S.</td>
<td>Michigan, East Lansing</td>
<td></td>
</tr>
<tr>
<td>Yost, H.</td>
<td>Massachusetts, Amherst</td>
<td></td>
</tr>
<tr>
<td>Your, J.</td>
<td>Korea, Seoul</td>
<td></td>
</tr>
<tr>
<td>Ytterborn, K.</td>
<td>Sweden, Stockholm</td>
<td></td>
</tr>
<tr>
<td>Zambrucchi, L.</td>
<td>Italy, Milan,</td>
<td></td>
</tr>
<tr>
<td>Zebe, F.</td>
<td></td>
<td>see DIS 34:130</td>
</tr>
<tr>
<td>Zigler, L.</td>
<td>Germany, Darmstadt</td>
<td></td>
</tr>
<tr>
<td>Zimmer, K.</td>
<td>Germany, Karlsruhe</td>
<td></td>
</tr>
<tr>
<td>Zimmering, S.</td>
<td>Indiana, Bloomington</td>
<td></td>
</tr>
<tr>
<td>Zimmermann, W.</td>
<td>Germany, Mariensee</td>
<td></td>
</tr>
<tr>
<td>Zürcher, C.</td>
<td>Switzerland, Zürich</td>
<td></td>
</tr>
</tbody>
</table>
(Editor's Comment--continued from page 8)

In any case, one must at the present time regard DIS primarily as a medium for the dissemination of stock lists and related technical data. However, I would regard the strictest adherence to this as unnecessarily stultifying, even if it were possible to define the above terms unambiguously. If one regards DIS instead as an informal means of promoting and facilitating the work of Drosophila geneticists, filling a niche not occupied by any other means of communication, then one can readily justify a broader interpretation of its function.

We are now including a section in which workers are given the opportunity to give approval to the quotation of specific notes of theirs by others. This may eliminate a good deal of unnecessary letter writing. I hope that the notes are mentioned specifically by DIS number and page number, for the following reason: when casual informal notes are written, sometimes covering work not yet completed, there are bound to be errors and it would help greatly if each worker would look closely at his past contributions and indicate (in the form suggested in the call) where further information about a preliminary note might be found. It is not really a question of whether the worker is willing to stand by any criticism for an incomplete or inaccurate note, as it is a matter of offering positive help to workers who want to get the maximum amount of correct information possible. Similarly I wonder about the desirability of the flat statement made by a number of workers that "all my notes, past present and future, may be quoted." The first presumes an infallibility that is more appropriate for publications rather than DIS notes and it is my considered judgment that this infallibility can be rightfully claimed by no more than three workers in the entire field. The second statement wills the infallibility to the second or third scientific generation and may be even more suspect. For these same reasons, I would, for the present at least, like to defer granting permission to quote current notes, in order to give each worker some time to consider carefully his own judgment of the quotability of that note.

Finally let me extend my thanks to the staff in Eugene, Oregon, who have put out this issue while I have been sojourning in Switzerland. These include: Mrs. Dorothy Parker, who has had over-all responsibility for the operation, Mrs. Elizabeth Ehrlich, Miss Jan Hamilton, Mrs. Mary Helen Landenberger, and Miss Hermia Ehrlich.

E. Novitski