tion of segregation ice and frozen sediments may represent annual temperature cycles, although this type of stratification can also occur under constant temperature conditions.

Because formation of segregation ice involves drawing additional water into the system, and because segregation ice can continue to form even under great pressure, ice island heights of 7 m are not unexpected. At most lakes, the islands probably were even higher in the past and definitely were larger in area. These ice deposits are now disappearing, and the most spectacular ones, such as those at Laguna Colorada, will probably not persist for more than a decade or two. However, their future will be strongly influenced by even small changes in lake water level, air temperature, or geothermal activity.

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Drosophila Males Contribute to Oogenesis in a **Multiple Mating Species**

Abstract. Two species of Drosophila that differ in their ecology and mating systems have been compared with respect to male contribution to the somatic tissues and developing oocytes of females. In the species Drosophila mojavensis females remate daily, exhibit a copulatory plug, and have been shown to obtain a contribution from the male ejaculate. In contrast, Drosophila melanogaster males do not contribute to females. Female Drosophila melanogaster do not remate as frequently as Drosophila mojavensis females nor is a copulatory plug formed.

Species of Drosophila show wide variability in mating systems (1). One of the most striking differences observed is the frequency at which females of various species remate. For example, mated females of the cosmopolitan species D. melanogaster, when provided with males every morning, usually will not remate for about 5 to 7 days (2). Under similar conditions, females of the Sonoran Desert endemic cactiphilic species D. mojavensis remate daily. The pattern of daily remating in D. mojavensis is observed even when mated females do not oviposit and even though the ventral receptacle may contain numerous sperm (1). Another difference between these two species is the formation of a reaction mass or copulatory plug following mating in D. mojavensis but not in D. melanogaster (3). The evolutionary significance of this difference in remating frequency is postulated to be linked to an interspecific difference in parental investment by males (1); specifically, D. mojavensis females, living in a harsh environment often with limited resources, are predicted to remate more frequently in order to obtain nutrients from the male ejaculate. We now present evidence that, during copulation, males of D. mojavensis contribute nutrients to oocytes and to female somatic tissues while D. melanogaster males apparently do not.

Males were isotopically labeled by placing freshly oviposited eggs on Carolina instant Drosophila medium containing 3 H-labeled amino acids (4). Virgin males were separated upon eclosion and stored at $24^{\circ} \pm 1^{\circ}C$ until they were required for mating experiments. After 4 days, labeled D. melanogaster males were mated to unlabeled females. Whole D. melanogaster males showed an average of 28,509 disintegrations per minute at the time of mating. Labeled D. mojavensis males were stored for 8 days before being mated to unlabeled females; these males showed an average of about 350,000 disintegrations per minute. The developmental time of D. mojavensis is 50 percent longer than D. melanogaster, which most likely accounts for the larger size of D. mojavensis and their higher concentration of radioactivity. The presence of isotope in mated females was determined immediately after copulation and again 24 hours later (5). The body parts analyzed are shown in Table 1.

In both species a large amount of radioactivity was seen in the female reproductive tract (uterus, ventral recepta-

Table 1. Radioactivity found in females at two times after mating. Data presented are averages for single females. At least three replications of three females per replication were performed for each time point. Counts per minute were converted to disintegrations per minute according to a standard quench curve. Controls consisted of body parts from unlabeled females. The results are given as means ± standard errors.

Radioactivity (disintegrations per minute)		
0 hours	24 hours	t*
Drosophila meland	ogaster	
18.9 ± 0.8	23.8 ± 6.1	1.39
23.7 ± 4.6	31.1 ± 8.7	1.31
39.5 ± 12.2	25.1 ± 3.1	1.09
$1112.4 \pm 63.1 \ddagger$	50.4 ± 23.9	6.47§
25.9 ± 9.8	23.9 ± 5.4	0.32
Drosophila mojav	ensis	
31.6 ± 6.6	$56.23 \pm 3.6 \ddagger$	5.211
50.4 ± 8.6	98.0 ± 5.5	7.77§
28.9 ± 6.4	$72.2 \pm 5.3 \ddagger$	10.24§
$1426.7 \pm 118.6 \ddagger$	$67.4 \pm 9.1 \ddagger$	84.44§
32.2 ± 8.5	$191.7 \pm 20.4 \ddagger$	8.622§
	$\begin{tabular}{ c c c c c } \hline Radioactivit\\ \hline 0 hours\\ \hline $Drosophila melano\\ 18.9 \pm 0.8\\ 23.7 \pm 4.6\\ 39.5 \pm 12.2\\ 1112.4 \pm 63.1 \ddagger\\ 25.9 \pm 9.8\\ \hline $Drosophila mojar\\ 31.6 \pm 6.6\\ 50.4 \pm 8.6\\ 28.9 \pm 6.4\\ 1426.7 \pm 118.6 \ddagger\\ 32.2 \pm 8.5\\ \hline \end{tabular}$	Radioactivity (disintegrations per minu0 hours24 hoursDrosophila melanogaster18.9 \pm 0.823.8 \pm 6.123.7 \pm 4.631.1 \pm 8.739.5 \pm 12.225.1 \pm 3.11112.4 \pm 63.1 \ddagger 50.4 \pm 23.925.9 \pm 9.823.9 \pm 5.4Drosophila mojavensis31.6 \pm 6.656.23 \pm 3.6 \ddagger 50.4 \pm 8.698.0 \pm 5.5 \ddagger 28.9 \pm 6.472.2 \pm 5.3 \ddagger 1426.7 \pm 118.6 \ddagger 67.4 \pm 9.1 \ddagger 32.2 \pm 8.5191.7 \pm 20.4 \ddagger

*When appropriate, the Welch method was used (11); otherwise two-tailed t-tests were used to compare sample means. †Minus reproductive tracts. ‡Differs significantly from unlabeled con-†Minus reproductive tracts. ||P < 0.05.‡Differs significantly P < 0.01.trols

cle, and spermathecae) immediately after copulation. At this point the amount of isotope in other parts did not differ significantly from that in control. Twenty-four hours later a large amount of isotope still remained in the reproductive tracts, although the level was reduced. In D. melanogaster females radioactivity decreased in the reproductive tract, and no significant amounts of isotope were detected in any other tissues after 24 hours. Drosophila females have been observed to expel material from their reproductive tract after mating (6), and we have assumed that this accounts for the reduction in the amount of isotope after 24 hours.

However, D. mojavensis females showed significant radioactivity in other body parts, especially unfertilized ovarian oocytes. Since oogenesis requires approximately the same length of time in both these species, and the same female reproductive state existed in both experiments, differential rates of oogenesis cannot be the underlying cause of the presence of isotope in D. mojavensis oocytes. The amount of isotope also increases in the somatic tissues of D. mojavensis females within 24 hours. The molecular nature of the substances transferred is still unknown, but since male contribution has also been documented with 14 C-labeled amino acids (7) any major role of tritium exchange in the observation of male contribution in D. mojavensis can be ruled out.

After two or three consecutive matings, D. melanogaster males are temporarily sterile (2, 8). This sterility is caused by a reduction in male accessory gland secretions, not by a reduction in sperm number. After 24 hours, abstinent males regain their fertility (2). In contrast, D. mojavensis males may mate seven or more times consecutively without any observable reduction in fertility (1). It is possible that D. mojavensis males transfer less material at each copulation in order to take advantage of the increased mating opportunities in their population.

The phenomenon of males transferring nutrients to females which then appear in female somatic tissues and oocytes has been demonstrated in several species of Lepidoptera (7). In these two Drosophila species, which differ in their ecology and mating systems, a difference in male contribution to egg production is also apparent. Drosophila melanogaster, a cosmopolitan species, can use a variety of substrates for breeding. However, D. mojavensis uses necrotic tissue of organ pipe cactus in Sonora, Mexico and

southern Arizona, and agria cactus in Baja California. At certain times of the year these resources are limited, and even during times of abundant resources females are selective about the stage and condition of the necrotic tissue on which they will oviposit (9). Females must be able to manufacture eggs and also survive until finding an appropriate place to oviposit, and male nutrient contribution may help them do this. The discovery of male nutrient contribution in a genus whose phylogenetic relationships are well defined, whose mating systems vary, and whose ecology is under intensive study (10) provides a new opportunity to inquire into the evolution of mating strategies in insects.

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Monoclonal Antibody to Thy-1 Enhances Regeneration of **Processes by Rat Retinal Ganglion Cells in Culture**

Abstract. Ganglion cells were dissociated from postnatal rat retinas, identified by specific fluorescent labels, and maintained in culture on a variety of substrates. Regeneration of processes by retinal ganglion cells was enhanced when the cells were plated on glass coated with a monoclonal antibody against the Thy-1 determinant. Plain glass and glass coated with polylysine, collagen, fibronectin, or other monoclonal antibodies supported the growth of neural processes, but were less effective than antibody to Thy-1.

Detailed studies of differentiated mammalian neurons would be aided by examining identified cells in vitro. This approach requires that viable cells be isolated, unequivocally identified, and cultured. Like other neurons of the central nervous system (CNS), mammalian retinal ganglion cells normally do not regenerate their axons after transection, and indeed many of the cells degenerate (1). It is important, therefore, to know whether differentiated retinal ganglion cells can survive and regenerate processes in culture.

Since ganglion cells are the only retinal cells that project to other areas of the CNS, they can be labeled by retrograde transport of markers injected into their projection sites, such as the superior colliculus and lateral geniculate nucleus (2). In histological sections Thy-1 anti-

gen is located on cells and processes in the inner retina and can be used to specifically identify the ganglion cells in vitro (2, 3). Thy-1 is also expressed on the surface of many neurons not found in the retina as well as on T lymphocytes and embryonic muscle (4). Because of sequence homology with immunoglobulins, it has been suggested that molecules displaying the Thy-1 antigen play a role in cellular recognition and morphogenesis in the nervous system (4).

In this report we describe aspects of the identification and growth of solitary rat retinal ganglion cells in culture. Mac-Leish et al. (5) observed that salamander neurons could be grown and maintained on antibody-coated glass cover slips; we tried this method with several antibodies. Comparing growth on different substrates, we found that, by the second day