Many insights into the molecular mechanisms underlying learning and memory have been elucidated through the use of simple behavioral assays in model organisms such as the fruit fly, *Drosophila melanogaster*. *Drosophila* is useful for understanding the basic neurobiology underlying cognitive deficits resulting from mutations in genes associated with human cognitive disorders, such as intellectual disability (ID) and autism. This work describes a methodology for testing learning and memory using a classic paradigm in *Drosophila* known as courtship conditioning. Male flies court females using a distinct pattern of easily recognizable behaviors. Premated females are not receptive to mating and will reject the male's copulation attempts. In response to this rejection, male flies reduce their courtship behavior. This learned reduction in courtship behavior is measured over time, serving as an indicator of learning and memory. The basic numerical output of this assay is the courtship index (CI), which is defined as the percentage of time that a male spends courting during a 10 min interval. The learning index (LI) is the relative reduction of CI in flies that have been exposed to a premated female compared to naïve flies with no previous social encounters. For the statistical comparison of LIs between genotypes, a randomization test with bootstrapping is used. To illustrate how the assay can be used to address the role of a gene relating to learning and memory, the pan-neuronal knockdown of *Dhap-at* was characterized here. The human ortholog of *Dhap-at*, glyceronephosphate O-acyltransferase (GNPT), is involved in rhizomelic chondrodysplasia punctata type 2, an autosomal-recessive syndrome characterized by severe ID. Using the courtship conditioning assay, it was determined that Dhap-at is required for long-term memory, but not for short-term memory. This result serves as a basis for further investigation of the underlying molecular mechanisms.
Protocol

NOTE: In the protocol outlined below, one replicate of collection, training, and testing is described. In order to test the reproducibility of the results, these steps should be repeated in parallel, on multiple days, and with separate groups of flies (Table 1). The protocol is based on a 10 day life cycle from egg to adult, which is normal when rearing flies under constant conditions of 25 °C, 70% humidity, and a 12 h light/dark cycle. All aspects of this protocol assume that the conditions are kept constant throughout the entire assay. Times are indicated as hours before lights turn on (BLO) or after lights turn on (ALO) in the incubator, as this can be conveniently set depending on the researcher's preferred time of day. Use CO₂ gas only for the initial collection of naïve male flies and for the collection of premated females. This protocol for courtship conditioning is composed of the following steps:

1. Establishment of premated female collection cultures
2. Establishment of cultures for the collection of male test subjects
3. Preparation of housing blocks
4. Establishment of mating vials for the production of standardized premated females
5. Collection of male test subjects
6. Training
7. Testing
8. Video data analysis and statistics

1. Establishment of Premated Female Collection Cultures

1. Prepare powerfood^{16}. Boil 0.8% (w/v) agar, 8% (w/v) yeast, 2% (w/v) yeast extract, 2% (w/v) peptone, 3% (w/v) sucrose, 6% (w/v) glucose, 0.05% (w/v) MgSO₄, and 0.05% (w/v) CaCl₂ in water for 15 min. Allow the solution to cool to 70 °C before adding 0.05% (w/v) methylparabene (CAUTION: toxic) and 0.5% (v/v) propionic acid (CAUTION: toxic). Mix well by stirring while cooling further to 50 °C to obtain a homogeneous solution.

2. Before the food solidifies at room temperature, add ~50 mL of powerfood to each 175 mL plastic vial. Allow the food to cool further. Close the vial with a plug.

NOTE: Powerfood is a specialized food mixture formulated specifically for the production of large numbers of flies, presumably by inducing egg laying. Powerfood is not used to produce male flies that will be used for behavior analysis (step 2) because the atypical diet and potential crowding might influence development.

3. On day -11 (Table 1), start 5-20 wildtype cultures with approximately 60-100 flies (a mix of males and females) in powerfood vials; these will be used in step 4 to produce standardized premated females^{17}. Add a filter paper to each vial to increase the area upon which the larvae can pupate; this will increase the number of flies that can eclose.

4. Periodically repeat steps 1.1-1.3 throughout the experiment to obtain sufficient newly eclosing flies as input for the “establishment of mating vials for the production of standardized premated females” (step 4).

2. Establishment of Cultures for the Collection of Male Test Subjects

1. Prepare normal food^{16}, made with 0.5% (w/v) agar, 2.75% (w/v) yeast, 5.2% (w/v) corn flour, 11% (w/v) sugar, 0.05% (w/v) methylparabene, and 0.5% (v/v) propionic acid in water, as described in steps 1.1 and 1.2. Close the 175-mL plastic vials with a fly vial plug.

2. On day -10 (Table 1), place about 10-20 males with approximately 30-75 virgin females (Materials/Equipment Table) in each 175 mL vial containing normal food. Add a filter paper to increase the surface area for pupation and to maximize productivity.

3. Establish three to six 175 mL vials per genotype to obtain the required number of test subject males.

NOTE: More vials may be needed, depending on the productivity of the desired genotype.

3. Preparation of Housing Blocks (Figure 2A)

1. Melt approximately 50 mL of powerfood per housing block in a microwave, or prepare it fresh.
4. Establishment of Mating Vials for the Production of Standardized Premated Females

1. On day -1, remove and discard all adult wildtype flies from premated female collection cultures at 2-5 h BLO.
2. Collect flies using the aspirator (Figure 2B) from these vials in 2- to 3-h intervals (e.g., at 30 min, 2.5 h, and 5 h ALO) and place them in a new powerfood vial supplemented with a small amount of yeast paste and filter paper.
3. To avoid crowding and to promote an optimal mating atmosphere, do not transfer more than 150-200 flies to each new vial. Ensure the mating of all females by providing at least 25% males. Ensure that sufficient females are present in the mating vials to accommodate the size of the experiment.

NOTE: As this is a crucial step in the protocol, make sure that only freshly eclosed flies and no old flies, larvae, or pupae are transferred to the new mating vial.

4. Incubate these "mating vials" for four days to allow sufficient time for all females to have mated.

5. Collection of Male Test Subjects

1. On day 1 (Table 1) at 2-3 h BLO, use CO₂ to remove all adult flies from the male collection vials (step 2), but let more flies eclose over the next few hours.
2. Over the next 5-6 h, remove newly eclosed flies every 20-30 min using CO₂ and put each male in an individual well of the housing block (step 3) using the aspirator (Figure 2B).
3. Re-seal the well with the adhesive PCR film.

NOTE: This is a crucial step in the protocol. The males should be collected frequently. Collected males should be isolated in the housing block close to the time of eclosion, when they demonstrate pale pigmentation and the presence of the meconium in the translucent abdomen. Gentle use of the aspirator allows the transfer of flies; however, inappropriate use will stress the flies, causing variance in the assay (see the Discussion).

4. Aim to collect up to 48 males per genotype. This provides a small excess to the maximal number of males needed for the analysis of both naive and trained conditions, allowing for some loss during later transfer steps.

6. Training

1. Remove the flies from the mating vial (step 4.2) using CO₂ and separate the premated females from the males.
2. Using the aspirator, add a single anesthetized, premated female to each well in one row of a new housing block.
3. Using the aspirator and without anesthesia, transfer an individual naive male from the housing block set up in step 5.2 to the well containing a premated female. After the male is placed into the well, re-seal immediately with the adhesive film; do not allow the male to escape.

NOTE: Transfer male flies from the aspirator to the housing block by taking advantage of their natural "negative geotaxis" behavior (see the Discussion).

4. Repeat steps 6.2-6.3 until enough male-female pairs are established. Ideally, establish 24 pairs, two full rows of a housing block, per genotype. Leave the remaining naive males in the original housing block set up in step 5.2.
5. Leave the male-female pairs undisturbed during the training period (Table 2, Figure 1B).

NOTE: During this time, the male will court and be rejected by the premated female. For learning and STM, the training period is 1 h and for LTM, the training period is 7-9 h.

6. End the training (Table 2, Figure 1B) by using an aspirator to gently separate the male from the premated female; do not use anesthesia. Place the separated male in a new housing block.
7. Use the aspirator to transfer all naive males gently and without anesthesia from the housing block set up in step 5.2 to a new housing block.

NOTE: This step is optional for STM and learning, but it is very important for LTM because the flies are housed for an additional 24 h to test LTM.

8. For STM and LTM, allow the males to rest for 1 h and ~24 h, respectively (Table 2, Figure 1B) before testing (step 7).
9. For learning, immediately test the trained and naive males (step 7).

7. Testing

1. Collect flies from mating vials (step 4.2) using CO₂ and separate the premated females from the males.
2. Let the females recover from the anesthesia for at least 1 h in a vial containing normal food.
3. Mount the video recorders in advance (Figure 2C), in order to have all equipment ready before the testing starts.
4. Start the testing according to the different timelines for learning, STM, and LTM (Table 2, Figure 1B). Perform testing immediately after training for learning, 1 h after training for STM, and 24 h after training for LTM.
5. Using the aspirator, gently transfer an individual male from the resting housing block or from the training housing block if learning is being tested (step 6.7, trained; step 6.8, naive) to one half of a courtship arena with the divider closed (Figure 2D; see File S1 for a building plan).

NOTE: The use of the natural "negative geotaxis" behavior should be sufficient to transfer the male flies from the aspirator to the courtship arena (Discussion).

6. Quickly but gently move the entry hole to the next arena and repeat step 7.5 until all 18 arenas contain one male.
7. Using the aspirator and without CO₂, add one premated female (collected in step 7.2) to the other half of all 18 arenas.
8. Carefully place the courtship chamber under the camera, with the opening of the wells facing down (Figure 2C).
9. Remove the divider of the arenas to allow direct interaction between the males and premated females.
10. Immediately start recording the behavior for at least 10 min.
   NOTE: When using a two-camera setup, the parallel recording of two courtship plates can be done in overlapping timeframes to maximize efficiency.
11. Empty the courtship arenas using a hand-held vacuum cleaner and allow the courtship chamber to ventilate before re-use.
12. Repeat step 7.4-7.11 until the testing of all genotypes and conditions (i.e., naïve and trained) have been completed.

8. Video Data Analysis and Statistics

1. Calculate the courtship index (CI), defined as the percentage of time that the male courts during the first 10 min of the testing period, for each individual male fly.
   NOTE: This can be done manually by observing stereotypical courtship behavior (Figure 1A) or by using computer software for automated quantification of courtship behavior (Discussion).
   NOTE: It is recommended to analyze 40-60 males per condition over the course of three days in order to achieve sufficient statistical power and to judge the consistency of the CI data.
2. Calculate the learning index (LI), defined as the percent reduction in the mean CI of trained males compared to naïve males (LI = (CI_naive – CI_trained) / CI_naive). Evaluate the LI for each day of testing and compare it to the cumulative LI calculated from all testing days combined.
3. Make a separate two-column tab data file with "Genotype" and "CI" as the headers.
   NOTE: These headings are case sensitive. The name of the genotype for each CI must consist of a description of the genotype followed by an underscore and the training condition (e.g., genotype_N and genotype_LTM, etc., where N = naïve and LTM = long term memory; see Supplemental File S2 for an example). This annotation is essential, as the function analearn will identify trained and naïve flies based on the characters present after the first underscore in the "Genotype" column.
4. Use the analearn R script (Supplemental File S3) to perform a randomization test to judge the statistical significance of differences between the LI values from different genotypes.
   1. Source the script (Supplemental File S3) into R, which defines a function called "analearn."
      NOTE: The function definition is: analearn <- function(nboot = 10,000, naivelevel = 'N', refmutation = NA, datname = NA, header = TRUE, seed = NA, writeoutput = TRUE).
   2. Start the function by entering "analearn()" in the R command line and selecting the data file to be analyzed (produced in step 8.3) via the pop-up window.
   3. Choose the reference mutation, which is the control genotype, by entering the corresponding number and pressing enter.
      NOTE: After selecting the reference genotype, the script takes several seconds to perform 10,000 bootstrap replicates.
   4. Observe the output table (Table 3), which contains the genotype, learning condition (i.e., learning, STM, or LTM), mean CI naïve, mean CI trained, LI, the difference between the LI of the control compared to the experimental condition (LI dif), the lower limit (LL) and upper limit (UL) of the 95% confidence interval of the LI dif, and the p-value indicating the probability that there is no significant difference.
      NOTE: analearn will store an output text file in the directory where the data file is located. However, the output table also appears in the R-Studio console. The default name is constructed based on the name of the data file provided.
   5. There are several arguments in the analearn function that can be used to alter the default settings of the function to adjust the parameters of the bootstrapping.
      NOTE: "nboot" defines the number of bootstrap replicates and is set to 10,000 by default. This value can be changed into any integer number larger than zero. Table 5 enlists several arguments that can be used to alter the default settings of the function. However, it is not recommended to use data that is produced with a low number of bootstrap replicates.
Representative Results

The courtship conditioning assay can be used to measure learning and memory in *Drosophila*. In order to demonstrate this, the results presented here analyze STM and LTM in control flies compared to flies with the neuron-specific knockdown of Dhap-at. Control males express an RNA interference hairpin sequence targeting a *Caenorhabditis elegans*-specific gene, putative zinc finger protein C02F5.12\(^{19}\). This control strain ensures an equal number of upstream activating sequence (UAS) elements between the knockdown and control in the same genetic background, and the control RNAi hairpin accounts for any non-specific effects of the RNAi system. Control males (genotype: *UAS-dcr2/+; P(KK108109)VIE-260B/+; elav-Gal4/+\(^{21}\)) show a significant reduction in CI in trained versus naive for both STM (Figure 3A, \(p = 1.2 \times 10^{-4}\), Mann-Whitney U-test) and LTM (Figure 3B, \(p = 1.2 \times 10^{-4}\), Mann-Whitney U-test). This result reflects the normal capacity for learning and memory in these flies. Dhap-at knockdown flies (genotype: *UAS-dcr2/+; P(KK101437)VIE-260B/+; elav-Gal4/+\(^{21}\)) also show a significant reduction of CI in trained versus naive flies for STM (Figure 3A, \(p = 1.2 \times 10^{-4}\), Mann-Whitney U-test). However, they do not show a significant reduction in CI after LTM training (Figure 3B, \(p = 0.33\), Mann-Whitney U-test). The Mann-Whitney U-test was used to compare the mean CI of naïve and trained flies due to the non-parametric distribution of the CI data for some conditions (Figure 3A and 3B). The differences observed through the analysis of CIs are reflected by the LI for each genotype, which measures the percent reduction in CI in naïve versus trained flies (Figure 3C). There is no significant difference in LI between controls and Dhap-at knockdown flies for STM (\(p = 0.115\), 10,000 bootstrap replicates, Figure 3C, Table 3), whereas the LI for LTM is significantly lower in Dhap-at knockdown flies (\(p = 0.0034\), 10,000 bootstrap replicates, Figure 3C, Table 3). For the comparison of LI between genotypes, a randomization test\(^{20}\) adapted from a method first recommended by Kamyshev et al.\(^{7}\) was used. Since the LI is a single value derived from population data (i.e., mean CI-naïve and mean CI-trained), standard statistical methods that rely on experimentally derived distributions do not apply. The randomization test is distribution-free and uses bootstrapping (i.e., random sampling with replacement) to generate hypothetical data sets that are derived from the actual data. The anealern script (File S3) generates a set of hypothetical LIs for each genotype and calculates the difference between the control and the test genotype (LIdiff). This process is repeated 10,000 times, and the resulting values are used to determine the 95% confidence interval of LIdiff (Table 3). This data is used to generate a \(p\)-value indicating the probability that the LI of the two genotypes is different. The results shown here demonstrate that Dhap-at is required in neurons for LTM but not for STM.

In order to control for day-to-day variability, CIs and LIs are compared between replicate days (Table 4). Although some fluctuation in LI is observed from day to day, the results are generally reproducible. It is important to note that CI data can vary greatly depending on the control strain used and the environmental conditions of testing. The CI data shown here is typical for this control genotype, but other genotypes may exhibit a higher or lower mean CI and distribution.
Figure 2: Equipment used for the Drosophila Courtship Conditioning Assay. (A) The housing block is a flat, bottom block with 500 µL of powerfood per well. It is sealed with a qPCR adhesive film with at least 4 holes per well that were created using a syringe needle with a 0.8 mm diameter. Individual rows are cut lengthwise into strips using a razor blade to allow opening and closing. (B) The aspirator is required for the gentle transfer of male and female flies without the use of anesthesia. The inset shows the tip of the aspirator, closed with a piece of cotton to keep the flies within the tip. (C) Setup of a two-camera system for the simultaneous recording of two courtship chambers. (D) A courtship chamber with 18 arenas. Sliding entry holes are used to place the flies in the arenas. The white dividers can be simultaneously opened to initiate interaction between the males and females. Please click here to view a larger version of this figure.
Figure 3: Analysis of STM and LTM in Control and Dhap-at Knockdown Flies. (A-B) Boxplots showing the distribution of CI values for naïve (N) and trained (T) flies of the control (gray) and Dhap-at knockdown (white) genotypes tested for STM (A) and LTM (B). (C) Corresponding LI values for control and Dhap-at knockdown flies tested for STM and LTM. Differences in LI between control and knockdown genotypes were compared using a randomization test (10,000 bootstrap replicates). Error bars indicate the standard error of the mean derived from the LI values calculated on different test days. The genotypes are: w+, UAS-dcr2/+; P{KK108109}VIE-260B/+; and elav-Gal4/+ (Control) and w+, UAS-dcr2/+; P{KK101437}VIE-260B/+; and elav-Gal4/+ (Dhap-at-RNAi). Please click here to view a larger version of this figure.
<table>
<thead>
<tr>
<th>General</th>
<th>Collect</th>
<th>Train</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>day -11</td>
<td>Start premated female collection cultures (step 1.3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>day -10</td>
<td>Start cultures for the collection of male test subjects (step 2.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>day 1</td>
<td>rep. 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>day 2</td>
<td>rep. 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>day 3</td>
<td>rep. 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>day 4</td>
<td>rep. 4</td>
<td>rep. 1</td>
<td></td>
</tr>
<tr>
<td>day 5</td>
<td>rep. 2</td>
<td>rep. 1</td>
<td></td>
</tr>
<tr>
<td>day 6</td>
<td>rep. 3</td>
<td>rep. 2</td>
<td></td>
</tr>
<tr>
<td>day 7</td>
<td>rep. 4</td>
<td>rep. 3</td>
<td></td>
</tr>
<tr>
<td>day 8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>day 9</td>
<td>Video data analysis and statistics (step 8)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

rep = repeat

**Table 1: Example Timeline for Testing LTM over Three Replicates on Individual Days.**

<table>
<thead>
<tr>
<th>Learning</th>
<th>STM</th>
<th>LTM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Training time</td>
<td>1 h.</td>
<td>1 h.</td>
</tr>
<tr>
<td>Resting time</td>
<td>0 h.</td>
<td>1 h.</td>
</tr>
<tr>
<td>start training</td>
<td>0 h. ALO</td>
<td>0 h. ALO</td>
</tr>
<tr>
<td>stop training</td>
<td>1 h. ALO</td>
<td>1 h. ALO</td>
</tr>
<tr>
<td>start test</td>
<td>1 h. ALO</td>
<td>2 h. ALO</td>
</tr>
</tbody>
</table>

ALO = after lights turn on, BLO = before lights turn on, STM = short term memory, LTM = long term memory

**Table 2: Training Duration, Training Times, and Testing Times for Learning, STM, and LTM.**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Learning condition</th>
<th>CI naïve</th>
<th>CI trained</th>
<th>LI</th>
<th>LI difference</th>
<th>Lower limit (95% confidence interval)</th>
<th>Upper limit (95% confidence interval)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>STM</td>
<td>0.467</td>
<td>0.116</td>
<td>0.752</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Dhap-at-RNAi</td>
<td>STM</td>
<td>0.699</td>
<td>0.257</td>
<td>0.633</td>
<td>0.119</td>
<td>-0.030</td>
<td>0.265</td>
<td>0.116</td>
</tr>
<tr>
<td>Control</td>
<td>LTM</td>
<td>0.590</td>
<td>0.384</td>
<td>0.348</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Dhap-at-RNAi</td>
<td>LTM</td>
<td>0.697</td>
<td>0.650</td>
<td>0.068</td>
<td>0.280</td>
<td>0.103</td>
<td>0.446</td>
<td>0.003</td>
</tr>
</tbody>
</table>

**Table 3: Statistical Data Produced from the Analearn Script.**

Statistical data produced from the analearn script. The output file of the bootstrapping R-script containing the genotype, learning condition (i.e., learning, STM, or LTM), mean CI naïve, mean CI trained, LI, difference between LI of the control compared to experimental condition (LI dif), the lower limit (LL) and upper limit (UL) of the 95% confidence interval of LI dif, and the p-value indicating the probability that there is no significant difference.
"writeoutput" can be set to "TRUE" or "FALSE" in order to determine whether an output file will be generated. The default is "TRUE."

When the seed is specified by any integer number larger than zero, the same set of random bootstrap samples is obtained.

By design, a bootstrap analysis will give slightly different results each time it is run, even when using the same data file.

"seed" initializes the random number generator. This is set by default to "NA" and ensures a random number each time the script is used.

The default is "TRUE," but a file with no headers can be used when this argument is changed to "FALSE."

"header" can be used to indicate whether or not the data file contains column headers.

The default is "TRUE," but a file with no headers can be used when this argument is changed to "FALSE."

"datname" refers to the name of the data file and can be specified in this argument instead of the default file selection.

"refmutation" is set to "NA" (not applicable) by default, but can be changed to the name of the control or the genotype in order to perform statistical comparisons.

This will cause the script to automatically select the control genotype.

Table 4: CI and LI Values Obtained on Separate Testing Days.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th></th>
<th>Dhap-at-RNAi</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average CI</td>
<td>Average CI</td>
<td>LI</td>
<td>Average CI</td>
<td>Average CI</td>
<td>LI</td>
</tr>
<tr>
<td></td>
<td>naïve</td>
<td>trained</td>
<td></td>
<td>naïve</td>
<td>trained</td>
<td></td>
</tr>
<tr>
<td>STM</td>
<td>Day 1</td>
<td>0.300</td>
<td>0.125</td>
<td>0.584</td>
<td>0.679</td>
<td>0.239</td>
</tr>
<tr>
<td></td>
<td>Day 2</td>
<td>0.634</td>
<td>0.107</td>
<td>0.831</td>
<td>0.720</td>
<td>0.276</td>
</tr>
<tr>
<td></td>
<td>All Days</td>
<td>0.467</td>
<td>0.116</td>
<td>0.752</td>
<td>0.699</td>
<td>0.257</td>
</tr>
<tr>
<td>LTM</td>
<td>Day 1</td>
<td>0.590</td>
<td>0.441</td>
<td>0.252</td>
<td>0.630</td>
<td>0.646</td>
</tr>
<tr>
<td></td>
<td>Day 2</td>
<td>0.640</td>
<td>0.363</td>
<td>0.432</td>
<td>0.709</td>
<td>0.710</td>
</tr>
<tr>
<td></td>
<td>Day 3</td>
<td>0.547</td>
<td>0.349</td>
<td>0.363</td>
<td>0.738</td>
<td>0.598</td>
</tr>
<tr>
<td></td>
<td>All Days</td>
<td>0.590</td>
<td>0.384</td>
<td>0.348</td>
<td>0.697</td>
<td>0.650</td>
</tr>
</tbody>
</table>

Table 5: Arguments used in the Analearn Function that Can Alter the Default Settings of the Function to Adjust the Parameters of the Bootstrapping

Supplemental File S1: Building plan of a courtship chamber. The file can be opened with any application that allows .stp extensions (CAD-files). Please click here to download this file.

Supplemental File S2: Example of an Input File for the Analearn Script. Please click here to download this file.

Supplemental File S3: The Analearn.R Acript. The file can be opened with R-studio. Please click here to download this file.

Discussion

The courtship conditioning assay is a classic paradigm for the analysis of learning and memory in Drosophila. The protocol presented here follows the general methodology described previously but includes unique aspects such as practical guidelines, specialized equipment, and a data analysis script for randomization tests. Using this protocol, it is possible to analyze large numbers of flies in parallel using 96-well flat-bottom blocks (Figure 2A) to collect and train males. The blocks are sealed with PCR adhesive film, which makes the flies easily accessible when required. Additionally, the unique courtship chambers described here allow for the simultaneous pairing of 18 male-female pairs in a nearly two-dimensional space that is optimal for video analysis. The custom-designed courtship chambers are easy to use, and a building plan is provided (File S1, Figure 2D). This protocol, from the establishment of the cultures used to collect test subjects to the acquisition of video data, takes approximately 20 days (Table 1). Additional time is required for the analysis of video data. In our experience, the STM assay is extremely robust. The LTM assay is also quite robust, but it is more sensitive to confounding environmental variables and therefore can be more difficult to master.

Animal behavior can be quite variable. Therefore, critical steps in the protocol must be performed with care to reduce this variance. First, gentle use of the aspirator (Figure 2B) will reduce the stress that can be imposed by rough handling or by blowing out too strongly. A suggested method of transferring individual flies out of the aspirator is by using negative geotaxis. As flies tend to walk up, one can simply point the tip of the aspirator up; just before the fly reaches the tip, a gentle blow is sufficient to let the fly out. Additionally, to let the males into the courtship chambers before testing, a blow is often not necessary.

Another important step is the collection and generation of male test subjects. All males must be collected when they are very young and socially naïve. This can be achieved by frequent collection during the peak periods of eclosion (step 5.2). If males are not collected in this tight timeframe, they can have early social interactions, which may result in poor learning or high variability in CI. Another factor of male test subjects that should be assessed is the genetic background. Different genetic backgrounds will exhibit different levels of naïve courtship and may also differ in general activity or locomotor ability. When comparing multiple genotypes, care should be taken with regard to genetic background in order to avoid these confounding factors that may influence LI scores. Additionally, the distribution of CI data should be carefully assessed. CI data can be both parametric and non-parametric, depending on the genotype or other environmental factors. In some cases, if the distribution of CI is dramatically skewed away from a normal distribution, it may be better to use the median CI rather than the mean for the calculation of LIs.
However, in our experience, the use of median or mean CI does not make a difference in the statistical interpretation of the data, and the use of the mean CI is the common practice in the literature.

For successful courtship conditioning, the active rejection of male courtship attempts by premated females is crucial during the training period. It is important to ensure that the premated females used in this assay have been efficiently mated and are thus not allowing copulation. This premating is established in the mating vials prepared in step 4, where male and females flies are housed together for 4 days (Table 1). Subsequently, mating can be monitored by regular examination of testing videos and by observing male-female pairs during training. If mating does occur, there are several measures that can be taken during the preparation of premated females. First, premated females should be reared under optimal breeding conditions. Vials can be supplemented with yeast paste and a folded filter paper to increase potential mating surfaces. The incubation of flies under the conditions described here has produced robust premated females in the past, but this may vary in different labs and with the use of different genetic strains. Therefore, it may be necessary to optimize the generation of premated females by varying the incubation time and conditions.

Quantification of courtship behavior is another critical step in this protocol. This can be done manually or automatically using specialized software programs. Automated quantification is fast and, in principle, unbiased. Several programs have been published, however, they are not straightforward to use, often requiring specialized video formats and advanced computational skills. Manual quantification is easy and accurate, but it is highly labor intensive and subject to individual variability and bias. It is important to emphasize that this protocol does not address the requirements for video formatting that are potentially required for the automated quantification of CIs. For manual quantification, use any simple video recording device that has the potential to produce a video of sufficient quality to accurately observe courtship behavior. For automated quantification, there will likely be different requirements depending on the software used, and users should investigate this thoroughly if automated quantification is desired.

In combination with the extensive tools that are available for the genetic manipulation of flies, the courtship conditioning assay provides a robust readout that can be used to dissect molecular mechanisms and neuronal networks involved in learning and memory.

Disclosures

The authors have nothing to disclose.

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