

Fat Turnover Assay in Drosophila

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[Abstract] Like all animals, *Drosophila* shows robust fat (triglyceride) turnover, *i.e.*, they synthesize, store and utilize triglyceride for their daily metabolic needs. The protocol describes a simple assay to measure this turnover of triglycerides in *Drosophila*.

[Background] Almost all animals store energy reserves in the form of glycogen and triglycerides. Many physiological, pathological and environmental conditions cause changes in the total level of these energy reserves, especially triglycerides. However, it's not always clear whether the resulting changes in triglycerides are due to reduced breakdown, increased synthesis or vice versa. With this protocol, it is possible to determine both the rate of synthesis and degradation of the newly synthesized triglycerides in flies.

Materials and Reagents

- 1. 1.5 ml Eppendorf tubes
- 2. Hamilton glass syringes (Hamilton, catalog number: Gastight 1700)
- 3. Metallic needle
- 4. Razor blades (VWR, catalog number: 55411)
- 5. TLC silica gel 60 plates (Figure S1) (EMD Millipore, catalog number: 105626)
- 6. Drosophila vial (Genesee Scientific, Flystuff, catalog number: 32-109)
- 7. Whatman® chromatography paper (Sigma-Aldrich, catalog number: WHA3030861)
- 8. Drosophila melanogaster
- 9. D-[14C(U)]-glucose (PerkinElmer, catalog number: NEC042V250UC)
- 10. Yeast extract
- 11. Sugar
- 12. Liquid nitrogen and nitrogen gas
- 13. Chloroform (Sigma-Aldrich, catalog number: 366927)
- 14. Methanol (EMD Millipore, catalog number: MX0475-1)
- 15. Cupric sulfate acid (EMD Millipore, catalog number: 102790)
- O-phosphoric acid (85%) (Thermo Fisher Scientific, Fisher Scientific, catalog number: A242-212)
- 17. Lipid standards:
 - a. Triolein (Sigma-Aldrich, catalog number: T7140)



- b. Phosphatidylcholine (Sigma-Aldrich, catalog number: P3556)
- c. Phosphatidylinositol (Sigma-Aldrich, catalog number: P6636)
- d. Cholesterol (Sigma-Aldrich, catalog number: C8667)
- e. Lauric acid (Sigma-Aldrich, catalog number: L556)
- f. Myristic acid (Sigma-Aldrich, catalog number: M3128)
- g. Palmitic acid (Sigma-Aldrich, catalog number: P0500)
- 18. Hexane (Sigma-Aldrich, catalog number: 32293)
- 19. Diethyl ether (Sigma-Aldrich, catalog number: 309966)
- 20. Acetic acid (AMRESCO, catalog number: 0714)
- 21. Sodium chloride (NaCl) (Sigma-Aldrich, catalog number: S9888)
- 22. Scintillation fluid (PerkinElmer, catalog number: Ultima GoldTM/6013329)
- 23. Drosophila food recipes (see Recipes)
- 24. Solvent mixture (see Recipes)
- 25. Cupric sulfate/phosphoric acid solution (see Recipes)
- 26. 0.9% NaCl solution (see Recipes)

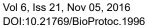
Equipment

- 1. TLC chamber (Clarkson Laboratory and Supply, model: Latch-Lid ChromatoTank 80-30)
- 2. Kontes microcentrifuge motor and pestles (Sigma-Aldrich, catalog number: Z359971-1EA)
- 3. Centrifuge (Eppendorf, model: Centrifuge 5810R)
- 4. Benchtop vacuum oven (VWR, model: 97027)
- 5. Scintillation counter (Beckman Coulter, model: LS6500)
- 6. Scintillation vials (PerkinElmer, catalog number: 6000292)
- 7. Nalgene PPCO wash bottles (Thermo Fisher Scientific, Thermo Scientific[™], catalog number: 2405-0500)

Procedure

1. The assay describes fat turnover in adult mated *Drosophila* females maintained on two different diets. The diets are HY (High yeast containing 5% yeast extract and 5% sugar) and LY (low yeast, containing 0.5% yeast extract and 5% sugar, see *Katewa et al.*, *2016*). On the tenth day about 300 flies (12 batches of 25 flies each) were transferred to fresh food vials with 2 μCi of ¹⁴C labeled glucose. Vials were prepared by adding 30 μl of 5% sugar/glucose solution on top and allowed to settle for 4 h. For preparing the sugar/glucose solution 100 μl of D-[¹⁴C(U)]- glucose was added to 900 μl of 5% sugar solution.

Note: When you add 30 µl of sugar/glucose solution on top of the food, it's best to use the sugar concentration that is closer to the concentration of sugar present in the regular lab fly food. For





- example, if your lab uses 10% sugar in the fly food, use 10% sugar solution to make the sugar/glucose solution.
- 2. The flies were maintained on the labeled food for 24 h, after which half of the flies were snapfrozen in liquid nitrogen. This is the 0 h sample. The other half was transferred to a fresh nonradioactive food vial and was kept on this food for the next 60 h, and was then immediately frozen. This is the 60 h sample.
- 3. For lipid extraction, weigh about 20 mg of flies/replicate. The weight is important for normalization and extraction of lipids. The number of flies will differ depending on the diets used in the experiment. For example, to get about 20 mg of total weight from the HY group, I need about 15-18 female flies, whereas for the LY group, I need about 22-25 flies. Homogenize the flies in about 100 µl of 0.9% NaCl in a 1.5 ml Eppendorf tube with a Kontes microcentrifuge motor and pestle. Transfer the homogenate to a 2 ml glass vial. Use additional 100 μl of 0.9% NaCl solution to rinse the Eppendorf tube and transfer to the glass vial. Add 800 µl of chloroform:methanol (2:1, v/v) to the glass vial with the homogenate, vortex for 15 sec and let it stand at room temperature for 20 min. Vortex again for 15 sec and centrifuge at 1,640 x g for 10 min to separate the two phases. By using a glass Hamilton syringe, carefully remove the lower phase containing the lipid fractions (Folch et al., 1957).
- 4. The lipid fraction is transferred to a glass vial and dried under a continuous stream of nitrogen gas (always in a chemical fume hood that is designated for radioactive work). A glass pipette or a metallic needle could be attached at the end of gas supply and suspended in the glass vials (about one inch away from the solvent surface). Once completely dried, add 100 µl of chloroform to the tube.
 - Note: There are two ways to separate the lipids into different fractions. One way is by using the solid phase extraction (SPE) tubes (for details see Katewa et al., 2012); another way is to use thin layer chromatography (TLC) plates (Chatterjee et al., 2014; Katewa et al., 2016). Here I am describing the TLC protocol as it is cost effective, simpler and also provides a visual separation of lipids.
- 5. A glass Hamilton syringe is used to spot the resuspended lipid on TLC plates. 25 µg triolein was loaded to identify the triglyceride (TG) migration band. Plates are allowed to air dry in a chemical hood for about 20 min. Plates are developed in a solvent mixture (see Recipes section, Kishimoto et al., 2001) in a pre-conditioned TLC chamber. Pre-condition of the chamber involves few additional steps. The chamber is prepared by adding all the components of the solvent mix (Hexane/diethyl ether/acetic acid [70/30/1, v/v]) in the chamber, covering the chamber and mixing vigorously. Additionally, for proper migration of the solvent on the TLC, the inner side of the TLC chamber is lined with two overlapping sheets of Whatman® chromatography paper such that it lines the back and side walls of the chamber (Figure 1). Next, cover the chamber and let it equilibrate for 30 min, before you put the first TLC in the chamber.

Note: Spots should be at least a centimeter above the solvent level.



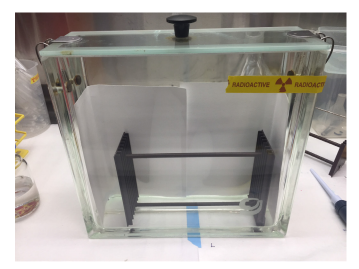


Figure 1. TLC chamber with lid. TLC chamber is lined with two overlapping sheets of Whatman[®] chromatography paper for saturation of the chamber. The inner Teflon[®] coated rack allows to simultaneous run several TLC plates.

- 6. Load the dried TLC plate in the chamber and let it run for 10-15 min (depending on the rate of migration of the liquid front). Take the plates out and air dry in a chemical hood.
- 7. For visual identification of TG spots, the TLC plate is sprayed with a solution of cupric sulfate/phosphoric acid (see Recipes section) and allowed to air dry for 30 min. The plates are heated in an oven at 125 °C for 30-45 min (until the TG band is visible). Remove the plate and let it cool. Figure 2 shows the efficiency of TLC to separate different lipid classes.



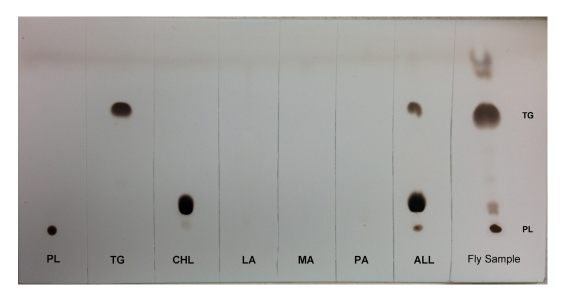


Figure 2. TLC separation of different lipids from adult female *Drosophila* melanogaster.

Total lipid was extracted from 15-20 adult female flies, dried and resuspended in 100 μ l of chloroform and loaded on TLC plates. The plates were developed further to visualize migration of different lipids. Lane arrangements are as follows: Lane 1, Phospholipid (PL, a 1:1 mixture of 10 μ g phosphatidylcholine and phosphatidylinositol). Lane 2, Triglyceride (TG, 25 μ g triolein). Lane 3, Cholesterol (CHL, 10 μ g). Lane 4, Lauric acid (LA, 10 μ g). Lane 5, Myristic acid (MA, 10 μ g). Lane 6, Palmitic acid (PA, 10 μ g). Lane 7, ALL (All standards mixed in equal proportions) and Lane 8, 100 μ l Fly sample.

8. The TG bands from the plates are scraped with a razor blade and transferred to scintillation vials. Add about 0.5 ml of hexane to extract the TG from the silica. Add 3 ml of the scintillation fluid, mix and count the ¹⁴C radioactivity. Radioactivity count in the 0 h samples indicate the amount of incorporation of glucose into triglycerides and radioactivity count in 60 h samples indicate the amount of label TG still retained. The difference between 0 h and 60 h samples indicate the breakdown of the labeled triglycerides.

Data analysis

The amount of ¹⁴C present in 0 h sample denotes the *de novo* synthesis of TG and is expressed as CPM/mg fly weight. One experiment containing 5-6 independent samples of 15-25 flies is used to obtain mean ± SEM values. Additional experimental repeats should be carried out to confirm the observed results. Based on the design of the study, one can use Student's *t*-test or ANOVA for obtaining the significance. Here, two different diets have been used, and the results (Figure 3) suggest that female flies on low yeast (LY) diet show increased *de novo* synthesis and a faster breakdown of TG when compared to the flies fed on high yeast (HY) diet.

Note: It's best to use independent bottles to obtain flies for replicates in one experiment.



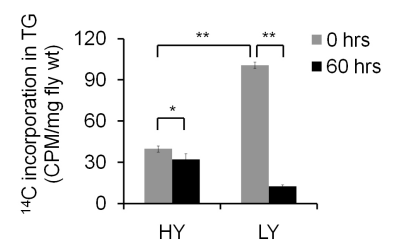


Figure 3. Fat turnover in adult female flies fed on high yeast (HY) and low yeast (LY) diets.

The amount of 14 C incorporation in 0 h sample indicates the *de novo* synthesis of TGs. The 60 h sample indicates the breakdown. Student's *t*-test was used to measure statistical significance, and error bars denote SEM of five independent preparations (*indicates P < 0.05 and **indicates P < 0.001)

Recipes

1. Drosophila food recipes

Detailed fly media recipes can be found in the Supplemental Experimental Procedures section of Katewa *et al.* (2016).

2. Solvent mixture

Hexane/diethyl ether/acetic acid (70/30/1, v/v). Add 105 ml of hexane, 45 ml of diethyl ether and 1.5 ml of acetic acid in the TLC chamber and mix vigorously.

- 3. Cupric sulfate/phosphoric acid solution8% cupric sulfate in 10% aqueous phosphoric acid
- 4. 0.9% NaCl solution
 - 9 g NaCl in 1 L of double-distilled water

Acknowledgments

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References

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