

# Triglyceride Assay Kit 100 point assay kit

Cat# TG-1-NC

STORAGE CONDITIONS	
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Reagents A & B, Buffers:

Store at 2 - 8°C. Use reconstituted Glycerol Reagent A within 7 days.

Glycerol Standard

- -20°C

ALL PRODUCTS ARE FOR RESEARCH USE ONLY. NOT APPROVED FOR HUMAN OR VETERINARY USE OR FOR USE IN DIAGNOSTIC OR CLINICAL PROCEDURES





The contents of this kit are sufficient for the assay of 100 assay points in a 96 well plate format. The only reagents sold separately are Glycerol Reagent A (cat# RGTA-10) and the glycerol standard for the Triglyceride Assay kit (cat# TG-GLYSTAN). For additional reagents or formats, please order our bulk 500 point assay kit (cat# TG-5RB).

## ITEMS INCLUDED IN THE KIT \_\_\_\_\_

ITEM	DESCRIPTION	UNIT	QTY	STORAGE
Wash buffer		BOTTLE	50ml	4°C
Lysis buffer		BOTTLE	15ml	4°C
Glycerol Reagent	Reconstitute w/ 11.0 ml deionized water	BOTTLE	11ml	4°C
Α	prior to use. Store at 4°C. Use reconstituted			
	Reagent A within 7 days.			
Reagent B	Reconstitute w/ 2.5 ml deionized water prior to use	BOTTLE	2.5 ml	4°C
TG Glycerol	Glycerol @ 1mM [Reconstitute with	VIAL	100 μl	-20°C
standard	400 μl Standards Diluent to make the			
	200 μM glycerol standard; see page 4			
	for recommended dilution scheme]			
Standards		BOTTLE	2.5 ml	4°C
Diluent				
Tray	Clear polyvinyl tray for multi-channel	EACH	3	
	pipetters			
Assay Plates,	96-well assay plate, blank	PLATE	2	
blank				

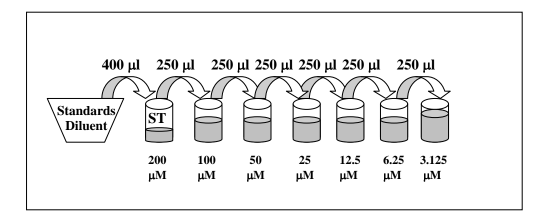
#### Other equipment/reagents required but not provided with the kit:

- Single-channel pipetter
- Multi-channel pipetter
- Plate reader with a filter of 540 nm
- Tubes to dilute glycerol standards
- Large gauge needle

### **ASSAY PROCEDURE**

- 1. Warm the Wash buffer and Lysis buffer in a 37°C water bath.
- 2. Prepare the Reagent B by adding 2.5ml deionized water per bottle and gently invert. DO NOT VORTEX! Use a pipet to ensure that the powder is completely dissolved. Keep at room temperature. Store in a light protected bottle. Reconstituted Reagent B is stable for 60 days refrigerated (2-8°C); store any remaining solution refrigerated (2-8°C). Bring Reagent B to room temperature.
- 3. Remove all media from mature human adipocytes. Using about 15 ml of the wash buffer, wash the cells one time with 150 µl wash buffer. Label the disposable tray "wash buffer" and retain for later use.
- 4. Remove all Wash buffer. Using a new tray, add 15  $\mu$ l Lysis buffer to each well. Incubate at 37°C 50°C for 20 minutes.
- 5. After the incubation is complete, visually confirm cell lysis by checking the wells under a microscope. If cells are not fully lysed, incubate another 10 minutes.
- 6. Add 135 µl warm Wash Buffer to each well.
- 7. Add 20  $\mu$ l Reagent B to each well. It is not necessary to mix at this time, however, gently tap the plate to help mix reagents. Incubate the plate at 37°C for 2 hours.
- 8. One hour prior to the assay, bring the glycerol standard and Reagent A to room temperature. Warm the Standards Diluent to 37°C. Prepare the standard curve as follows:

Pipette 400  $\mu$ l of the Standards Diluent into the 1 mM glycerol standard tube provided and mix well by vortexing. This produces a diluted stock glycerol standard of 200  $\mu$ M. Pipette 250  $\mu$ l of diluent into 6 tubes (not provided). Using the newly diluted stock glycerol solution, prepare a dilution series as depicted below. Mix each new dilution thoroughly before proceeding to the next. The 200  $\mu$ M stock dilution serves as the highest standard, and the diluent serves as the zero standard.



Note: The above dilution series generates enough volume to perform the standard curve in duplicate. If you wish to perform the standard curve in duplicate, please note that eight fewer data points can be assayed with this kit.

- 9. Also at this time prepare the Reagent A by adding 11.0 ml deionized water per bottle and gently invert. DO NOT VORTEX! Use a pipet to ensure that the powder is completely dissolved. Keep at room temperature. If using a Reagent A solution previously prepared and stored at 2-8°C, also bring to room temperature. Make sure there is enough Reagent A from one solution to treat all the points in the assay. It may be necessary to combine solutions. Store in a light protected bottle. Reconstituted Glycerol Reagent A is stable for 7 days refrigerated (2-8°C); store any remaining solution refrigerated (2-8°C).
- 10. To a blank 96 well plate, add 80  $\mu$ l wash buffer to each well needed for the assay (NOTE: do not add Wash Buffer to the wells used for the standard curve).
- 11. Working with one row or column at a time, mix the lysates very well using a multi-channel pipet. Immediately transfer 20 µl per well of the lysates to the corresponding well of the plate containing the wash buffer. **This results in a Dilution Factor of 5.**
- 12. Prepare the standard curve. Pipet 100  $\mu$ l of each standard into a well. (NOTE: Eight wells are necessary for the curve or sixteen if performing the curve in duplicate. If there are remaining wells on the assay plate, you can utilize the remaining wells. If not, a second plate is included in this kit).
- 13. Using the third tray, add 100  $\mu$ l Reagent A to samples and standards. Mix by pipetting up and down one time. Incubate at room temperature for 15 minutes.
- 14. Read at 540 nm using a microtiter plate reader.

## **GLYCEROL STANDARD CURVE**

This kit is designed to show relative lipid accumulation of experimental treatments compared to controls. The assay is based on the equation

1 M Triglyceride yields 1M glycerol + Free Fatty Acids

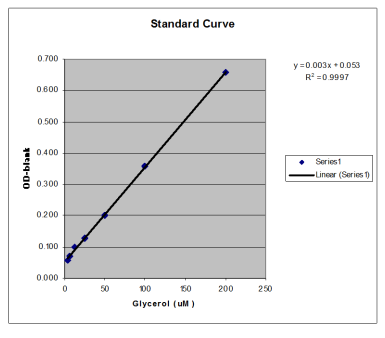
The reagent measures the concentration of glycerol released after lysing the cells and hydrolyzing the triglyceride molecules. The triglyceride concentration can then be determined from the glycerol values.

Generate standard curve: see example below [DO NOT use this standard curve to generate your data. This is an example.]

Subtract the OD value of the 0μM standard from all OD values including the standard curve.

					Avg
Glycerol			OD-	OD-	OD-
(uM)	OD	OD	blank	blank	blank
0	0.048	0.048			0.048
3.125	0.059	0.058	0.011	0.01	0.059
6.25	0.07	0.07	0.022	0.022	0.070
12.5	0.098	0.098	0.05	0.05	0.098
25	0.127	0.13	0.079	0.082	0.129
50	0.2	0.205	0.152	0.157	0.203
100	0.353	0.362	0.305	0.314	0.358
200	0.649	0.667	0.601	0.619	0.658

slope	0.003
intercept	0.053
$R^2$	0.9997



y = observed O.D. minus the blank

 $x = concentration of glycerol in \mu M$ 

To calculate x for each y, (i.e. to change the observed O.D. into glycerol concentration) use the following equation:

y=(slope) times (x) plus intercept

y=mx+b so x=(y-b)/m

x=(y-0.0006)/0.0014 where 0.0014= slope of the line and 0.0006= y intercept.

Be careful to enter the proper sign for the y intercept value as it may be a negative number.

Any OD values greater than the highest standard (200  $\mu$ M) should be suspect. The compound should be re-assayed using a lower dose of the compound at treatment OR a dilute solution of the condition medium at the time of the assay.

The R<sup>2</sup> value should be equal or greater then 0.98 for the standard curve to be valid. Any R<sup>2</sup> values below 0.98, must have the standard curve run again.

Solve for the Total Glycerol concentration (i.e. total triglyceride concentration) for each OD. Remember to include the Dilution Factor in the equation.

Data is expressed as µM Glycerol.

NOTE: Any OD values that are negative after the blank is subtracted should be considered to be 0 for the OD value.

#### REFERENCES

- 1. Green, H. and Kehinde, O. (1974) Sublines of mouse 3T3 cells that accumulate lipid. *Cell* 1, 113-116.
- 2. Hauner, H., et al., (1989) J. Clin. Invest. (84), 1663-1670.
- 3. Kuri-Harcuch W, Wise LS, Green H. (1978) Interruption of the adipose conversion of 3T3 cells by biotin deficiency: differentiation without triglyceride accumulation. *Cell* **14:**53-58.

#### **TROUBLESHOOTING**

Problem	Suggestions
High background or the triglyceride reagent turns a darker color before the assay begins.	<ul><li>Use clean tray and tips</li><li>Change pipet tips frequently</li></ul>
Edge effects	Ensure a saturated humidity in the incubator to prevent evaporation from the outside wells
Inconsistent OD reading	<ul> <li>Be careful when pipetting to avoid bubbles. If bubbles persist, burst the bubbles using a large gauge needle prior to reading and read the plate again.</li> <li>Mix the lysates well before transferring the 20µl to the Wash buffer plate.</li> </ul>

## FREQUENTLY ASKED QUESTIONS

- 1. Can I buy the reagents separately? The only reagents sold separately are Glycerol Reagent A (cat# RGTA-10) and the glycerol standard for the Triglyceride Assay kit (cat# TG-GLYSTAN).
- 2. Can I use another plate format besides 96 well? This kit is designed for the assay of A 96 well plate (100 assay points). We do not have a protocol for other formats.
- 3. Can I use this kit to measure total triglyceride in other cell lines and other human and non-human cells? Yes. The assay is not species specific. As long as the sample concentration is in the linear range, this kit should be able to detect it.
- 4. **My cells did not lyse.** What can I do? If cells are not fully lysed, incubate for another 10 minutes at 37°C 50°C. Sometimes mixing by pipetting up and down several times is necessary for full lysis.
- 5. I do not have time to complete the assay. Can I freeze the samples? Yes. The cell lysates can be stored at -80°C for a maximum of 7 days. Mix the thawed lysates in the plate by pipetting up and down several times. Allow all reagents and samples to reach room temperature <a href="BEFORE">BEFORE</a> adding the Wash Buffer and Glycerol Reagent A to complete the assay.

## APPENDIX A: PLATE LAYOUT \_\_\_\_\_

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## APPENDIX B: PROCEDURE FLOWCHART

Remove all media from wells.

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Wash with 150  $\mu$ l Wash Buffer Remove all Wash Buffer from wells and add 15  $\mu$ l Lysis Buffer/well.

Incubate 20 minutes at 37 – 50°C

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Verify lysis and add 135 µl warm Wash Buffer. Mix by pipetting up and down 3 times.



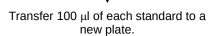
Add 20 μl Reagent B/well and incubate 2 hours at 37 °C.



One hour prior to assay,

reconstitute Glycerol Reagent A and prepare standards. Keep all at room temperature.

Add 80 µl Wash Buffer to a new plate.
Mix lysates and transfer 20 µl lysates to the wells containing Wash Buffer.





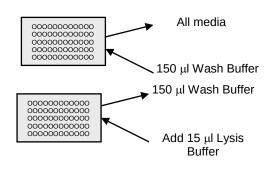
Add 100 µl Reagent A to samples and standards.

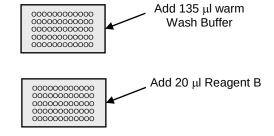


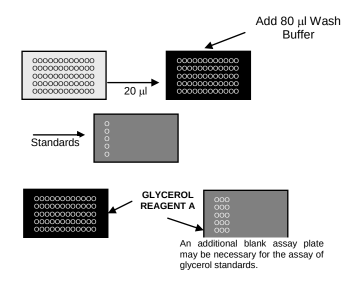
Incubate 15 minutes at 25°C (room temperature). Pop the bubbles in each well.



Measure the optical density of each well at 540 nm using a spectrophotometer plate reader.







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