

HUMAN C-MYC CELL-BASED KITS

Part # 63ADK053PEG & 63ADK053PEH

Test size#: 500 tests (63ADK053PEG), 10,000 tests (63ADK053PEH) - assay volume: 20 µL

Revision: #05 of September 2023

Store at: ≤- 60°C (63ADK053PEG); ≤- 60°C (63ADK053PEH)

For research use only. Not for use in diagnostic procedures.

ASSAY PRINCIPLE

Revvity Human c-Myc cell-based assay is only intended for quantitative measurement of Human c-Myc in cells using HTRF® technology.

Human c-Myc is detected in a sandwich assay format using 2 different specific antibodies, labeled with Europium Cryptate (donor) and with d2 (acceptor).

The principle of detection is based on HTRF® technology. When the dyes are in close proximity, the excitation of the donor with a light source (laser or flash lamp) triggers a Fluorescence Resonance Energy Transfer (FRET) towards the acceptor, which in turn fluoresces at a specific wavelength (665 nm). The donor & acceptor labeled antibodies bind to the Human c-Myc present in the sample, thereby generating FRET. Signal intensity is proportional to the number of antigen-antibody complexes formed and therefore to the Human c-Myc concentration (Fig. 1).

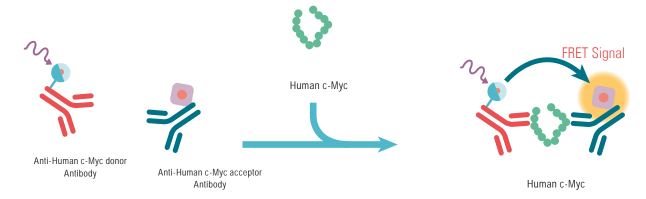
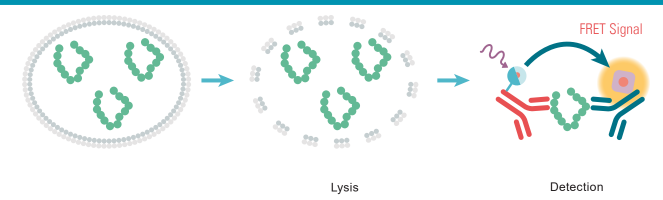


Figure 1: Principle of HTRF Human c-Myc sandwich assay.

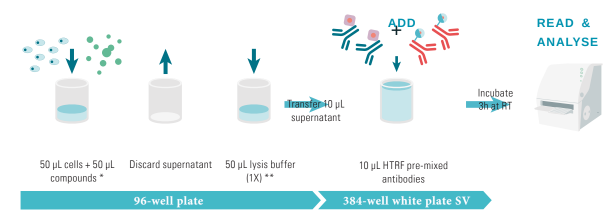
The assay is run under a two-plate assay manual, where cells are plated, stimulated and lysed in the same culture plate. Lysates are then transferred to the assay plate for the detection of Human c-Myc by HTRF® reagents. This manual gives the cells viability and confluence to be monitored.

Technical support team can help you to set-up this manual or another one. Please contact us at www.revvity.com

MANUAL AT A GLANCE



TWO-PLATE ASSAY MANUAL (FOR ADHERENT CELLS):



- * Note that concentration above 0.5% DMSO will impair assay performances.
- ** Depending on cell lines used, volume of lysis should be optimized, it can also be necessary to dilute the cell lysate to ensure samples are within the assay linear range.

MATERIALS PROVIDED:

Kit components	500 tests Cat # 63ADK053PEG	10,000 tests Cat # 63ADK053PEH
Control lysate Frozen/ready-to-use	1 vial - 150 μL	2 vials - 150µL
Anti-Human c-Myc-Eu Cryptate Antibody	1 vial - 50 μL Frozen - 50 X	1 vial - 1 mL Frozen - 50 X
Anti-Human c-Myc-d2 Antibody	1 vial - 50 μL Frozen - 50 X	1 vial - 1 mL Frozen - 50 X
Lysis buffer #4 * stock solution 4X	1 vial - 16 mL Frozen	1 vial - 130 mL Frozen
Detection Buffer #3 ** ready-to-use	1 vial - 7 mL Frozen	1 vial - 105 mL Frozen

^{*} Amounts of reagents provided are sufficient for generating 50 µL of cell lysate per well.

PURCHASE SEPARATELY:

- HTRF®-Certified Reader**. Make sure the setup for Eu Cryptate is used
- For a list of HTRF-compatible readers and set-up recommendations, please visit www.revvity.com
- Small volume (SV) detection microplates Use white plate only.
- For more information about microplate recommendations, please visit our website at: www.revvity.com

^{**} The Detection Buffer is used to prepare working solutions of acceptor and donor reagents.

STORAGE AND STABILITY

Antibodies, control lysate and buffers should be stored frozen until use.

Thawed detection buffer can be stored at 2-8°C in your premises. Thawed antibodies are stable 48 hours at 2-8°C; they can be refrozen (at -20°C or below) and thawed at least one more time. Control lysate must be stored frozen at -60°C or below. Thawed control lysate can be refrozen (at -60°C or below) and thawed one more time.

REAGENT PREPARATION

Allow all reagents to thaw before use.

We recommend centrifuging the vials gently after thawing, before pipeting the stock solutions.

Prepare the working solutions from stock solutions by following the instructions below.

POSITIVE CONTROL SOLUTION: READY-TO-USE

The control cell lysate is only provided as an internal assay control to check the quality of the results obtained. The window between control lysate and negative control should be greater than 2.

TO PREPARE WORKING ANTIBODY SOLUTIONS:

HTRF® reagent concentrations have been set for optimal assay performances. Note that any dilution or improper use of the d2 and Cryptate-antibodies will impair the assay's quality. Be careful, as working solution preparation for antibodies may differ between the 500 and 10,000 tests data point kit.

Antibody working solutions are stable for 2 days at 4°C. Dilute the antibodies with detection buffer #3.

500 TESTS KIT - 63ADK053PEG 10,000 TESTS KIT - 63ADK053PEH Anti-Human c-Myc- Cryptate antibody Dilute 50-fold the frozen stock solution with detection 1 vol 49 vol Dilute 50-fold the frozen stock solution with Detection 49 vol buffer #3: e.g. add 2.45 mL of detection buffer to the 1 vol buffer #3: e.g. add 49 mL of detection buffer to the 0.05 mL of Cryptate-antibody stock solution. 1 mL of Cryptateantibody stock solution. Anti-Human c-Myc-d2 antibody Dilute 50-fold the frozen stock solution with Detection Dilute 50-fold the frozen stock solution with Detection 1 vol 49 vol buffer #3: e.g. add 2.45 mL of detection buffer to the 1 vol 49 vol buffer #3: e.g. add 49 mL of detection buffer to the 0.05 mL of d2- antibody stock solution. 1 mL of d2- antibody stock solution. Antibody mix It is possible to pre-mix the two ready-to-use antibody It is possible to pre-mix the two ready-to-use antibody solutions just prior to dispensing the reagents by solutions just prior to dispensing the reagents by adding 1 volume of d2-antibody solution to 1 volume of adding 1 volume of d2-antibody solution to 1 volume of Cryptate-antibody solution. Cryptate-antibody solution.

TO PREPARE LYSIS BUFFER:

Make sure that the lysate has been generated by using the kit reagents.

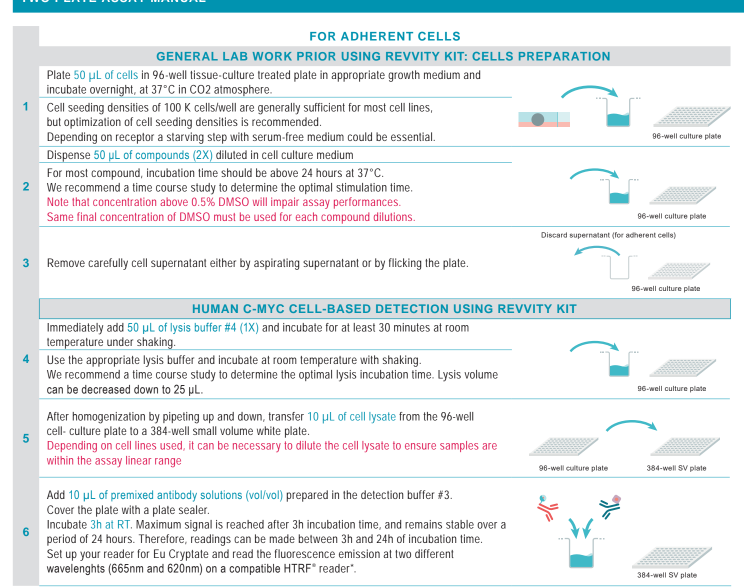
Prepare the required amount of lysis buffer before running the assay, working solutions are stable for 2 days at 2-8°C.

Lysis buffer 1X:

Determine the amount of lysis buffer needed for the experiment. Each well requires generally 50 µL of lysis buffer. Prepare a lysis buffer solution 1X by diluting 4-fold the lysis buffer 4X with distilled water.

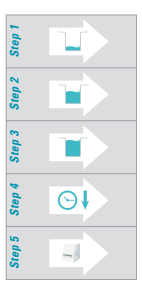
Preparation of lysis buffer 1X Dilute the "lysis buffer 4X" 4-fold with distilled water to prepare lysis buffer 1X. e.g. take 1.25 mL of lysis buffer 4X and add it to 3.75 mL of distilled water. Mix gently. Dilute the "lysis buffer 4X" 4-fold with distilled water to prepare lysis buffer 1X. e.g. take 1.25 mL of lysis buffer 4X and add it to 3.75 mL of distilled water. Mix gently.

TWO PLATE ASSAY MANUAL



^{*} For more information about HTRF® compatible readers and for set-up recommendations, please visit our website at: www.revvity.com

Standard manual for two-plate assay manual in 20 µL final volume (after lysis step)



Non treated cell lysate	Treated cell lysate	Positive control	Negative control	Blank control
Dispense 10 µL of non treated cell lysate	Dispense 10 µL of treated cell lysate	Dispense 10 µL of control lysate	Dispense 10 µL of lysis buffer 1X	Dispense 10 µL of non treated cell lysate
Add 5 μL of Anti-Human c-Myc-d2 Antibody working solution to all wells				Add 5 µL of detection buffer
Add	5 μL of Anti Human c-M	lyc-Eu Cryptate Antibody	y working solution to all	wells
		h at room temperature. I 24 hours. Therefore, rea incubation time.		
	Remove the plate sea	aler and read on an HTR	F® compatible reader	

The blank control is used to check the Cryptate signal at 620 nm.

The Negative control is used to check the non-specific signal. The ratio between control lysate signal / non-specific signal should be greater than 2.

DATA REDUCTION & INTERPRETATION

1. Calculate the ratio of the acceptor and donor emission signals for each individual well.

Ratio =
$$\frac{\text{Signal 665 nm}}{\text{Signal 620 nm}} \times 10^4$$

2. Calculate the % CVs. The mean and standard deviation can then be worked out from ratio replicates.

For more information about data reduction, please visit www.revvity.com

RESULTS

These data should be considered only as an example. Results may vary from one HTRF® compatible reader to another.

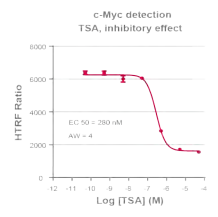
The curves are drawn up by plotting HTRF® Ratio versus the log [compound] concentrations.

Results were obtained on human Hela cells (40,000 cells/well) using the two-plate assay manual for adherent cells.

Human Hela cells were treated for 24 hours at 37°C with increasing concentrations of TSA,a c-Myc inhibitor.

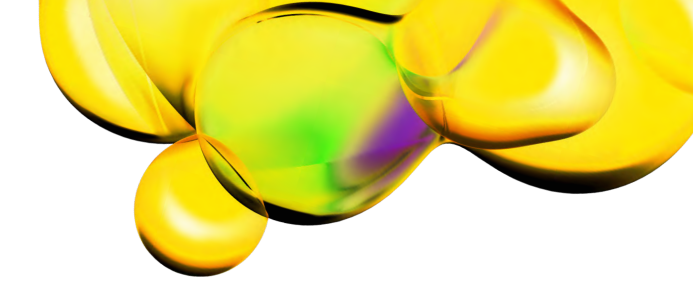
PHERAstarFS with flash lamp (BMG) was used for reading.

	c-Myc - Results		
Log [TSA] (M)	Ratio (1)	CV% (2)	
-4.3	1555	3.2%	
-5.3	1718	1.9%	
-6.3	2845	2.2%	
-7.3	6056	1.0%	
-8.3	6006	5.6%	
-9.3	6364	3.4%	
untreated cells	6382	3.0%	



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