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# A DIRECT COMPARISON OF THUNDER™ ASSAY PERFORMANCE WITH TWO EXISTING TR-FRET PLATFORMS

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### **KEY POINTS**

We compared head-to-head the performance of THUNDER™ with two existing TR-FRET assay technologies. Six endogenous phosphorylated proteins were measured in lysates from cells treated with pathway-specific modulators. THUNDER<sup>™</sup> phospho-protein assays exhibit superior or comparable performance at a lower cost per well.

# INTRODUCTION

Phosphorylation is one of the most important and common posttranslational modifications of proteins. Protein phosphorylation plays a critical role in cell signaling in response to extracellular stimulus and is of fundamental importance in biological regulation and human diseases [1]. Accordingly, the availability of assays capable of measuring, target-specific protein phosphorylation in a physiologically relevant cellular context is important for both basic research and drug discovery.

A number of technologies are currently available for the microplatebased measurement of intracellular phosphorylated protein levels in crude cell lysates. The enzyme-linked immunosorbent assay (ELISA) was developed in the 1970s and remains the mainstay for this kind of analysis. However, ELISA has several well-known limitations. Conventional ELISAs use large volumes of sample, are time-consuming and error-prone due to a laborious protocol with multiple washing and incubation steps, have a short period for signal detection (usually within 30 minutes) and are typically performed in 96-well plates, being difficult to miniaturize to higher density plate formats.

In recent years, time-resolved Förster resonance energy transfer TR-FRET technology has emerged as a superior alternative to

ELISA [2]. TR-FRET combines the low background and sensitivity of time-resolved fluorometry with the homogeneous assay format of FRET. TR-FRET assays are performed using a simple no-wash, "add-incubate-measure" protocol that substantially reduces assay complexity and hands-on time, while improving throughput and suitability for miniaturization and automation. In addition, TR-FRET assays have reduced sample volume requirements, higher reproducibility and long signal stability.

Bioauxilium has recently introduced THUNDER<sup>TM</sup>, a new immunoassay platform based on an enhanced TR-FRET technology (see Application Note THUNDER-APP001). The THUNDER<sup>TM</sup> Cell Signaling Assay Kits are designed to measure endogenous levels of specific intracellular phosphorylated proteins with high sensitivity, specificity, robustness and cost effectiveness. The aim of this Application Note was to compare head-to-head the performance of THUNDER<sup>TM</sup> in 384-well plate format with two existing TR-FRET technologies (Companies A and B) measuring a panel of six phosphorylated proteins. The three TR-FRET assay technology platforms were evaluated for their capacity to measure relative levels of phosphorylated 4EBP1 (T37/T46), AKTpan (S473), ERK1/2 (T202/Y204) p38 $\alpha\beta\gamma$  (T180/Y182), SLP-76 (S376) and STAT3 (Y705) in whole-cell lysates from cells treated with pathway-specific modulators.

## THUNDER™ TR-FRET ASSAY PRINCIPLE

The three TR-FRET technologies use different donor and acceptor fluorophores (shown in Table 1). However, all TR-FRET assay platforms are based on the same sandwich immunoassay principle. The assay principle of THUNDER™ is shown in Figure 1. Following treatment, cells are first lysed with the specific Lysis Buffer included in the kit. The target phosphorylated protein in the whole cell lysate is then detected in a single addition step with a pair of fluorophore-labeled specific antibodies that recognize distinct epitopes on the protein. One antibody is labeled with a Europium chelate donor (Eu-Abl), whereas the second antibody is labeled with a far-red small fluorophore acceptor (FR-Ab2). The binding of the two labeled antibodies to the target protein takes place in solution and brings the two dyes into close proximity. Excitation of the donor Europium chelate molecules at 320 or 340 nm triggers a FRET from the donor to the acceptor molecules, which in turn emit a signal at 665 nm. This signal is proportional to the concentration of phosphorylated protein in the cell lysate. In the absence of the specific target protein, the donor and acceptor fluorophores are too distant from each other for FRET to occur.

## MATERIALS AND METHODS

The THUNDER™ assay kits used for this evaluation are listed in Table 2. Comparative kits were purchased from established vendors. Reagents provided with each THUNDER™ assay kit are listed in Table 3. All reagents were prepared according to each manufacturer's recommendations. Milli-Q® water was used for diluting all Lysis and Detection buffers. Other reagents and materials used in the study are shown in Table 4.

#### Table 2 THUNDER™ kits used in the study.

Assay Kit	Catalog number
Phospho-4EBP1 (T37/T46)	KIT-4EBP1P-100
Phospho-AKTpan* (S473)	KIT-AKTS473P-100
Phospho-ERK1/2 (T202/Y204)	KIT-ERKP-100
Phospho-SLP-76 (S376)	KIT-SLP76P-100
Phospho-STAT3 (Y705)	KIT-STAT3P-100
Phospho-p38αβγ (T180/Y182)	KIT-P38P-100

\* AKTpan is AKT 1/2/3

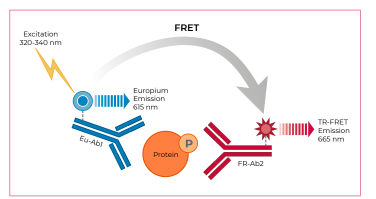
 Table 3
 Reagents supplied with each THUNDER™ Cell Signaling

 Assay Kit.

Kit contents
Europium chelate-labeled phospho-protein antibody (Eu-Ab1)
Acceptor-labeled phospho-protein antibody (FR-Ab2)
Lysis Buffer (5X)
Detection Buffer (10X)
Positive control cell lysate
Phosphatase Inhibitor Cocktail (100X)

Table 1Donor and acceptor fluorophores used by eachTR-FRET technology.

	TR-FRET Technology			
FRET moiety	THUNDER™	COMPANY A	COMPANY B	
Donor	Eu chelate	Eu cryptate	Eu chelate	
Acceptor	Far-red dye	d2 dye	ULight dye	





#### Table 4 Reagents and materials used in the study.

Reagents and materials	Source (catalog #)
Cell lines	
A431	ATCC (CRL-1555)
HEK293	ATCC (CRL-1573)
HeLa	ATCC (CCL-2)
Jurkat	ATCC (TIB-152)
MCF7	ATCC (HTB-22)
Reagents	
Anisomycin	Cayman (11308)
DMEM	Wisent (320-005-CL)
EGF	PeproTech (AF-100-15)
EMEM	Wisent (320-005-CL)
FBS	Wisent (098-050)
H <sub>2</sub> O <sub>2</sub>	Sigma (216763)
ΙΝFα	ProSpec (CYT-460)
Insulin	Sigma (19278)
PP242	Cayman (0469889-17)
RPMI	Wisent (350-007-CL)
Ultra pure water	MilliQ® water
Materials	
96-well culture microplate	Costar (3595)
384-well white microplate	PerkinElmer (6007290)
Microplate seal	PerkinElmer (6050185)

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THUNDER™ TR-FRET Technology

The overall assay workflow is the same for all TR-FRET assays and is outlined in Figure 2. All assays were conducted manually using the standard two-plate transfer protocol for each kit, whereby the cells are seeded, treated and lysed in a 96-well culture plate, and lysates are then transferred to a low-volume 384-well assay plate for protein detection.

STEP 1	STEP 2	STEP 3
Cell treatment	Cell lysis	Protein detection
• Seed cells in culture plate • Add media	• Adherent cells: remove media and add 1X Lysis Buffer	• Transfer lysate (15 µL) to detection plate
+/- compound • Incubate for	• Suspension cells: add 5X Lysis Buffer	• Add <b>4X Antibody</b> <b>Mix</b> (5 µL)
optimized time	Incubate for <b>30 min</b>	Incubate for 1-18 h     Read TR-FRET signal

Figure 2 THUNDER™ assay workflow using the two-plate transfer protocol.

Cell lines, culture conditions and treatment protocols used for this study are listed in Table 5. For each comparison, assays were conducted side-by-side using the same batch of treated cells. Adherent cells were seeded in 96-well culture plates and treated the following day. Suspension cells were treated immediately following plating. Cells were treated with varying concentrations of pathway-specific modulators according to pre-optimized conditions. All compounds were dissolved in serum-free culture media.

All assays were conducted according to each manufacturer's protocols. The two-plate protocol used for the THUNDER™ assay platform is summarized in Figure 3. Following treatment, adherent cells were lysed with the corresponding 1X Lysis Buffer, whereas suspension cells were lysed with the corresponding concentrated Lysis Buffer. The different Lysis Buffers contained phosphatase inhibitors. Following a 30-min incubation at room temperature (RT) on an orbital shaker (400 rpm), aliquots of lysates were then transferred in triplicate to the same white 384-well plate microplate followed by the addition of the corresponding antibody detection mix (prepared in the corresponding 1X detection buffer). All assays were run in a total assay volume of 20 µL. Plates were covered with an adhesive plate sealer to reduce evaporation during incubation. The plate sealer was removed before TR-FRET reading. Plates were read at multiple incubation times on an EnVision® 2104 Multilabel Plate Reader in TR-FRET mode (lamp excitation) using an excitation filter at 320 nm and emission filters at 615 nm (620 nm for Company A) and 665 nm. TR-FRET data were expressed as emission ratios of acceptor/donor (665 nm/615 nm) signals.

Data in the figures are shown as mean  $\pm$  standard deviation (three wells per assay point). Compound concentration response data were analyzed using nonlinear regression and fitted to a sigmoidal four-parameter equation with  $1/Y^2$  weighting (GraphPad Prism software). Performance metrics were S/B ratio, IC<sub>50</sub> and EC<sub>50</sub>, intraassay variability (%CV), and stability of S/B ratio and pharmacology following overnight incubation [3].

#### Adherent cells

Plate 50 $\mu$ L of cells in 96-well culture plate
Incubate overnight
Add 50 $\mu$ L of 2X compound in culture medium
Incubate for optimized time
Remove medium. Add 50 $\mu$ L of 1X Supplemented Lysis Buffer
Incubate 30 min at RT with shaking
Transfer 15 $\mu$ L of lysate to 384-well plate
+
Add 5 µL of 4X Antibody Mix (Eu-Ab1 + FR-Ab2)

Incubate 1 h to overnight at RT

Read TR-FRET

#### **Suspension cells**

Plate 20  $\mu L$  of cells in 96-well culture plate

Incubate overnight

Add 20 µL 2X compound in culture medium

Incubate for optimized time

Add 10 µL of 5X Supplemented Lysis Buffer

Incubate 30 min at RT with shaking

Transfer 15 µL of lysate to 384-well plate

Add 5 µL of 4X Antibody Mix (Eu-Ab1 + FR-Ab2)

Incubate 1 h to overnight at RT

**Read TR-FRET** 

Figure 3 THUNDER™ assay protocols for adherent and suspension cells.

 Table 5
 Cell lines and treatment conditions used for testing the TR-FRET assay kits.

	Phospho-protein						
	<b>4EBP1 ΑΚΤραη ΕRΚ1/2 SLP-76 STAT3 p38</b> αβγ						
Cell line	A431	MCF7	HEK293	Jurkat	HeLa	HeLa	
Cell seeding density	25,000 cells/well	60,000 cells/well	50,000 cells/well	400,000 cells/well	40,000 cells/well	50,000 cells/well	
Cell culture conditions	DMEM +10% FBS	DMEM +10% FBS	EMEM +10% FBS	RPMI	DMEM +10% FBS	DMEM +10% FBS	
Treatment conditions	PP242 3 hours at 37°C	Insulin 10 min at RT	EGF 10 min at RT	H <sub>2</sub> O <sub>2</sub> 15 min at RT	IFNα2b 20 min at RT	Anisomycin 1 hour at 37°C	

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THUNDER™ TR-FRET Technology

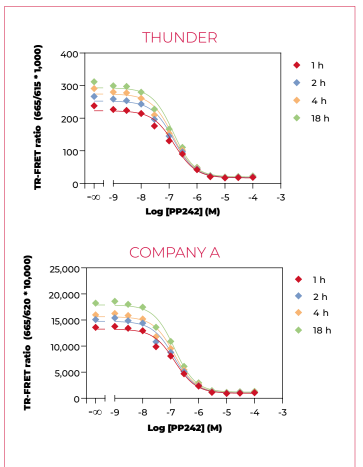
### **RESULTS AND DISCUSSION**

In order to adequately benchmark the performance of the new THUNDER<sup>™</sup> TR-FRET platform against two competitive TR-FRET technologies, comparisons were conducted using real cell lysate samples, rather than recombinant proteins spiked into the corresponding lysis buffers. Indeed, the assay performance obtained with recombinant proteins is not always predictive of the performance with real cell extracts, since antibody pairs might recognize the target proteins differently in each sample matrix. In addition, it is essential to evaluate the effectiveness of each kit's lysis buffer for releasing the target protein. To better assess the sensitivity and dynamic range of each assay kit, we conducted concentrationresponses curves for each target protein tested.

The results of the head-to-head comparisons are summarized in Figures 4 to 9. Overall, all three TR-FRET technologies showed an increase in the specific signal (treated cells) and S/B ratios (treated versus untreated cells) as a function of incubation time. However, there were differences in terms of the maximal S/B ratios reached by each platform. THUNDER<sup>™</sup> assays showed the highest S/B ratios for phospho-AKTpan, phospho-ERK1/2 and phospho-p38aβγ and S/B ratios comparable with those of Company A for phospho-4EBP1. Companies A and B showed the highest S/B ratios generated by THUNDER<sup>™</sup> for these two phospho-proteins were only slightly lower (36% and 25%, respectively). Company B exhibited the lowest S/B ratios for phospho-AKTpan, phospho-ERK1/2 and phospho-STAT3.

Table 6 summarizes the S/B ratios obtained with the three TR-FRET assay platforms for each phospho-protein tested at the detection time point recommended by each manufacturer. Whereas assays from Bioauxilium and Company A exhibited acceptable S/B ratios ( $\geq$  3) at all recommended detection times, two assays from Company B (phospho-AKTI/2/3 and phospho-STAT3) showed sub-optimal S/B ratios (< 3).

The three TR-FRET technologies exhibited comparable inter-well variability (typical %CV <8%) and sensitivity (IC<sub>50</sub> and EC<sub>50</sub> values) for all six phosphorylated proteins tested (Figures 4 to 9). In addition, all technologies tolerated plate reading after overnight incubation, which allows off-line readings for increased flexibility and productivity. Of note, all THUNDER<sup>TM</sup> assays except phospho-STAT3 allowed plate reading after only 1 hour of incubation. This was not the case for AKTpan and STAT3 from Companies A and B.



	тни	NDER™	Company A		
Incubation time	S/B EC <sub>50</sub> (nM)		S/B	EC <sub>50</sub> (nM)	
lh	12.9	154	12.7	139	
2 h	13.4	144	13.8	132	
4 h	14.0	136	13.6	139	
18 h	13.7	141	13.9	138	

Figure 4 Head-to-head assessment of the Phospho-4EBP1 (T37/T46) assays.

Table 6 Summary of S/B ratios obtained with each TR-FRET technology at the recommended detection time points.

Assay	S/B ratio (at manufacturer's recommended detection time)			Conclusion
	THUNDER™	Company A	Company B	
Phospho-4EBP1	14.0 (4 h)	13.8 (2 h)	Kit not available	Comparable performance between THUNDER™ and Company A kits
Phospho-AKTpan	5.7 (4 h)	3.0 (4 h)	1.9 (4 h)	Highest S/B ratio with THUNDER™ kit
Phospho-ERK1/2	11.2 (4 h)	10.8 (4 h)	5.9 (4 h)	Comparable performance between THUNDER™ and Company A kits (but highest S/B ratio with THUNDER™ kit after overnight incubation)
Phospho-SLP-76	4.7 (4 h)	4.7 (4 h)	6.3 (4 h)	Highest S/B ratio with Company B kit
Phospho-STAT3	4.3 (18h)	6.7 (18 h)	2.8 (4 h)	Highest S/B ratio with Company A kit
Phospho-p38αβγ	19.2 (4 h)	13.5 (4 h)	Kit not available	Highest S/B ratio with THUNDER™ kit

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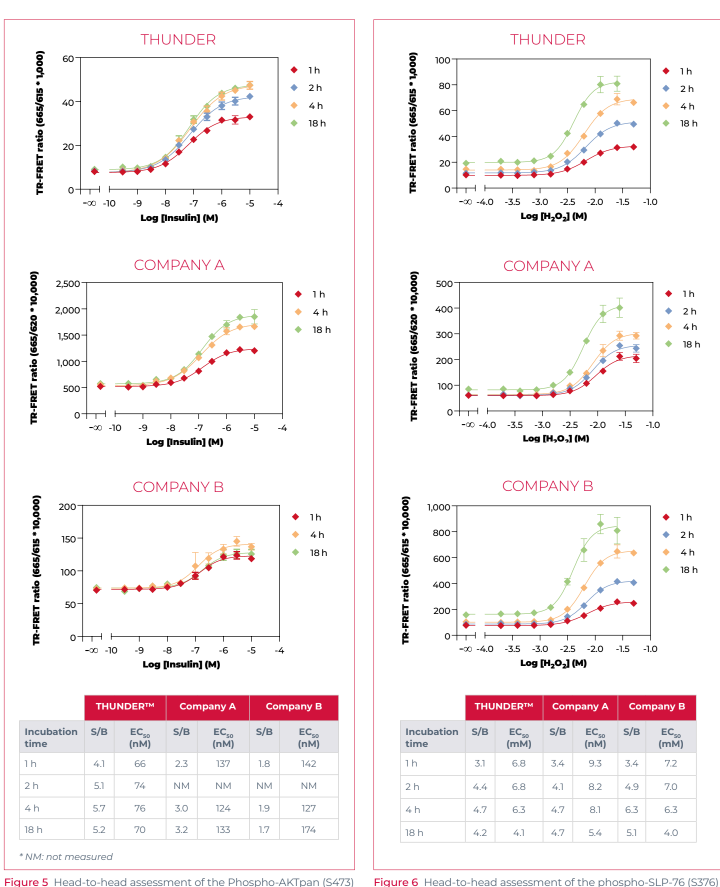


Figure 5 Head-to-head assessment of the Phospho-AKTpan (S473) assays.

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assays.

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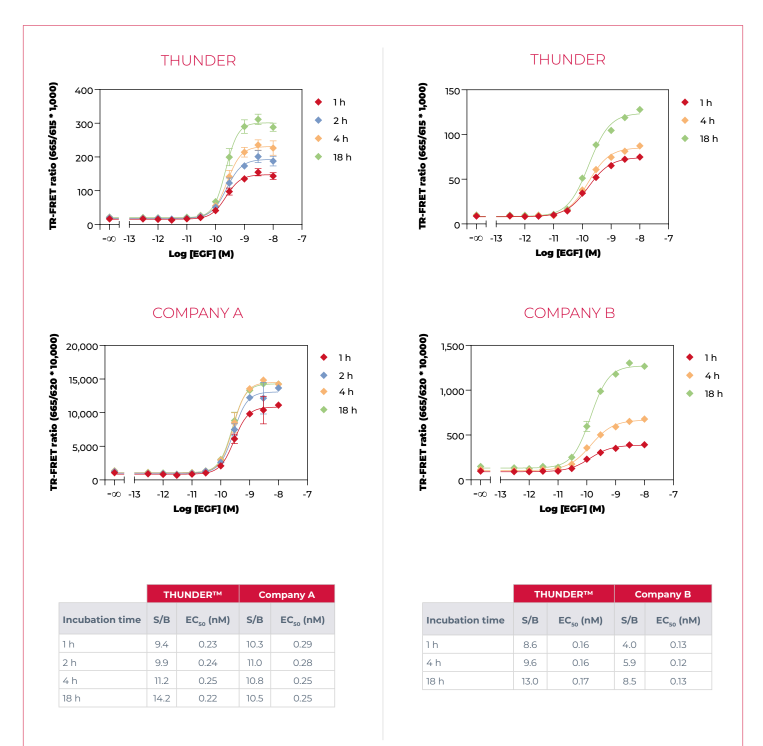
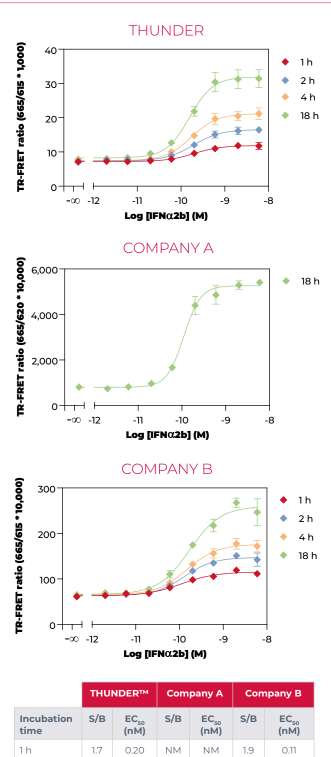


Figure 7 Head-to-head assessment of the phospho-ERK1/2 (T202/Y204) assays.

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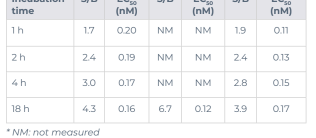
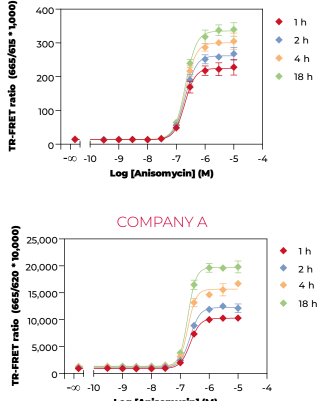


Figure 8 Head-to-head assessment of the phospho-STAT3 (Y705) assays.



THUNDER

400

Log [Anisomycin] (M)

	тни	NDER™	Company A	
Incubation time	S/B EC <sub>so</sub> (nM)		S/B	EC <sub>50</sub> (nM)
lh	16.1	196	11.1	220
2 h	18.4	202	11.6	215
4 h	19.2	207	13.5	182
18 h	23.2	207	15.0	182

Figure 9 Head-to-head assessment of the phospho-p38 $\alpha\beta\gamma$ (T180/Y182) assays.

#### THUNDER™ TR-FRET Technology

To serve both academic and industrial researchers, assay technologies need to deliver quality data at an acceptable cost per well. As shown in Table 7, THUNDER™ assay kits are more cost-effective than the other two commercial TR-FRET technologies.

#### Table 7 Assay cost analysis.

	Cost per well (USD) <sup>1</sup>			
Kit size	THUNDER™	Company B		
96 tests	\$3.35	\$5.66 <sup>2</sup>	Not available	
500 tests	\$2.11	\$3.40	\$3.87 <sup>3</sup>	
2 500 tests	\$1.80	Not available	Not available	
5 000 tests	\$1.12	Not available	Not available	
10 000 tests	\$0.70	\$0.99	\$0.73 <sup>3</sup>	

<sup>1</sup>Assay cost per well is calculated by dividing the commercial cost of the assay by the number of samples that can be analyzed. <sup>2</sup> Few kits available in the 96-test format.

<sup>3</sup>When purchasing the positive control lysate (not included in the kit).

### CONCLUSIONS

This study aimed at benchmarking the performance of three homogeneous TR-FRET assay platforms on the market today for measuring the relative levels of six different endogenous phosphorylated proteins in whole cell lysates. All three platforms were studied for their capacity to deliver quality data using the two-plate assay protocol with target protein detection in 384-well microplates.

Collectively, the results of these head-to-head comparisons showed that all THUNDER<sup>TM</sup> assays tested provide either superior or comparable S/B ratios, and comparable assay pharmacology (EC<sub>50</sub> and IC<sub>50</sub> values). In addition, THUNDER<sup>TM</sup> is more cost effective than the other two TR-FRET technologies.

These key advantages, combined with rigorous assay validation using cell lysates from stimulated/inhibited cells, and higher flexibility in terms of kit sizes and formats (detection of either phosphorylated, total or phosphorylated plus total proteins with the same kit), make the THUNDER<sup>™</sup> assay platform an alternative of choice for monitoring cellular protein phosphorylation.

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