

## Produktinformation



Forschungsprodukte & Biochemikalien



Zellkultur & Verbrauchsmaterial



Diagnostik & molekulare Diagnostik



Laborgeräte & Service

Weitere Information auf den folgenden Seiten! See the following pages for more information!



## Lieferung & Zahlungsart

siehe unsere Liefer- und Versandbedingungen

## Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

### SZABO-SCANDIC HandelsgmbH

Quellenstraße 110, A-1100 Wien

T. +43(0)1 489 3961-0

F. +43(0)1 489 3961-7

mail@szabo-scandic.com

www.szabo-scandic.com

linkedin.com/company/szaboscandic in





Fax: 1.858.481.8694 Email: info@bpsbioscience.com

# Data Sheet PDE1B TR-FRET Assay Kit Catalog # 60704

**DESCRIPTION:** Phosphodiesterases (PDEs) play an important role in dynamic regulation of cAMP and cGMP signaling. PDE1B is a calcium/calmodulin-dependent cyclic nucleotide phosphodiesterase that is highly expressed in the striatum. It plays a physiological role in the central nervous system, and PDE1B activity has been linked to impaired cognition and spatial learning.

The *PDE1B TR-FRET Assay Kit* is designed for identification of inhibitors of PDE1B using TR-FRET (Time Resolved Fluorescence Resonance Energy Transfer) technology. The assay is based on the generation of FAM-labeled nucleotide monophosphates by the PDE1B. These phosphate groups bind to terbium-labeled nanoparticles, resulting in energy transfer from the terbium to the FAM, which emits a fluorescent signal at 520 nm. The change in fluorescent intensity can be easily measured using a fluorescence plate reader.

The PDE1B TR-FRET Assay Kit comes in a convenient 96-well format, with purified PDE1B enzyme, fluorescently labeled PDE1B substrate (cAMP), binding agent, and PDE assay buffer for 100 enzyme reactions. Using this kit, only two simple steps on a microtiter plate are required for the PDE1B activity assay. First, the fluorescent-labeled cAMP is incubated with a sample containing PDE1B for 1 hour. Second, a binding agent and a terbium donor are added to the reaction mix and incubated for 30 minutes. Then, fluorescence intensity can be measured using a fluorescence reader.

Fax: 1.858.481.8694
Email: info@bpsbioscience.com

#### **COMPONENTS:**

Catalog #	Component	Amount	Storage	
60011	PDE1B recombinant enzyme	1 µg	-80°C	
60200	FAM-Cyclic-3', 5'-AMP: 20 µM	20 µl	-80°C	
60393	PDE assay buffer	25 ml	-20°C	
	Tb donor	30 µl	-80°C	(Avoid
60390	Binding Agent	200 µl	+4°C	freeze/ thaw
	Binding Buffer A	20 ml	+4°C	cycles!)
	Binding Buffer B	20 ml	+4°C	
VWR	Black, low binding NUNC microtiter	1	Room	
62408-936	plate		temp.	

#### MATERIALS OR INSTRUMENTS REQUIRED BUT NOT SUPPLIED:

Fluorescent microplate reader capable of measuring Time Resolved Fluorescence Resonance Energy Transfer (TR-FRET)

**APPLICATIONS:** Great for screening small molecular inhibitors for drug discovery and HTS applications.

**STABILITY:** At least 6 months from date of receipt when stored as directed.

REFERENCE(S): Siuciak JA, et al. (2007) Neuropharmacology 53(1):113-24.

#### **ASSAY PROTOCOL:**

All samples and controls should be tested in duplicate.

#### **Protocol for PDE1B assay**

#### Step 1:

- 1) Dilute 20 μM FAM-Cyclic-3',5'-AMP substrate stock solution 100-fold with PDE assay buffer to make a 200 nM solution. Make only a sufficient quantity needed for the assay; store remaining stock solution in aliquots at -20°C.
- Add 25 μI of FAM-Cyclic-3',5'-AMP (200 nM) to each well designated "Substrate Control", "Positive Control", and "Test Inhibitor". Add 25 μI of PDE assay buffer to each well designated "Tb-only Control".
- 3) Add 5 µl of inhibitor solution to each well designated "Test Inhibitor". Add 5 µl of the same solution without inhibitor (inhibitor buffer) to the "Tb-only Control", "Substrate Control" and "Positive Control".
- 4) Thaw PDE1B on ice. Upon first thaw, briefly spin tube containing enzyme to recover the full contents of the tube. Aliquot PDE1B enzyme into single-use



Fax: 1.858.481.8694
Email: info@bpsbioscience.com

aliquots. Store remaining undiluted enzyme in aliquots at -80°C immediately. Note: PDE1B is very sensitive to freeze/thaw cycles. Do not re-use thawed aliquots or diluted enzyme.

5) Dilute PDE1B in PDE buffer to 0.1 ng/µl (2 ng/reaction) in PDE buffer\*. Add 20 µl of PDE assay buffer to the wells designated as the "Tb-only Control " and "Substrate Control". Initiate reaction by adding 20 µl of PDE1B (0.1 ng/µl) to the wells designated for the "Positive Control" and "Test Inhibitor". Discard any remaining diluted enzyme after use. \*Note: optimal enzyme concentration may vary with the specific activity of the enzyme.

	Tb Only Control	Substrate Control	Positive Control	Test Inhibitor
FAM-Cyclic-3',5'-AMP (200 nM)	_	25 µl	25 µl	25 µl
PDE assay buffer	45 µl	20 µl	_	_
Test Inhibitor	_	_	_	5 µl
Inhibitor Buffer (no inhibitor)	5 µl	5 µl	5 µl	_
PDE1B (0.1 ng/µl)	-	_	20 µl	20 µl
Total	50 µl	50 µl	50 μl	50 μl

6) Incubate at room temperature for 1 hour.

#### Step 2:

- Make binding dilution buffer by mixing equal volume Binding Buffer A and Binding Buffer B. For example, mix 1 ml Binding Buffer A with 1 ml Binding Buffer B.
- 2) Mix **binding agent** thoroughly and dilute **binding agent** 1:50 with binding dilution buffer made in Step 1.ss
- 3) Add Tb donor (1:1,000 dilution) to the mixture in Step 2.
- 4) Add 100  $\mu$ I to each well. Incubate at room temperature for 1 hour with slow shaking.
- 5) Read the fluorescent intensity in a microtiter-plate reader capable of TR-FRET.



6044 Cornerstone Court W, Ste E San Diego, CA 92121

**Tel:** 1.858.829.3082 **Fax:** 1.858.481.8694

Email: info@bpsbioscience.com

#### **Instrument Settings**

Reading Mode	Time Resolved		
Excitation Wavelength	330±20		
Emission Wavelength	490±10		
Lag Time	50 μs		
Integration Time	50 μs		
Excitation Wavelength	330±20		
Emission Wavelength	520±10		
Lag Time	50 µs		
Integration Time	50 μs		

#### **CALCULATING RESULTS:**

$$FRET = \frac{S_{520} - \left(\frac{Tb_{520}}{Tb_{490}} \times S_{490}\right)}{S_{490}} \times 1000$$

Where  $S_{520}$  = Sample 520 nm reading,  $S_{490}$  = Sample 490 nm reading,  $Tb_{520}$  = Tb only 520 nm reading,  $Tb_{490}$  = Tb only 490 nm reading. When percentage activity is calculated, the FRET value from substrate only control can be set as zero percent activity and the FRET value from positive control can be set as one hundred percent activity.

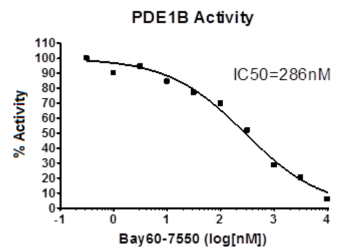
$$\% \ Activity = \frac{FRET_S - FRET_{Sub}}{FRET_P - FRET_{Sub}} \times 100\%$$

Where  $FRET_s = Sample FRET$ ,  $FRET_{Sub} = Substrate only control FRET$ , and  $FRET_P = Positive control FRET$ .



Fax: 1.858.481.8694
Email: info@bpsbioscience.com

#### **EXAMPLE OF ASSAY RESULTS:**



Inhibition of PDE1B by Bay60-7550, measured using the *PDE1B TR-FRET Assay Kit*, BPS Bioscience # 60704. *Data shown is lot-specific. For lot-specific information, please contact BPS Bioscience, Inc. at info@bpsbioscience.com* 

#### **RELATED PRODUCTS:**

<u>Product</u>	Catalog #	<u>Size</u>
PDE1A1	60010	10 µg
PDE1B	60011	10 µg
PDE1B (rat)	60009	10 µg
PDE1C	60013	10 µg
PDE1C (mouse)	60012	10 µg
PDE Assay Kit	60300	96 rxns.
PDE1C TR-FRET Assay Kit	60705	96 rxns.
PDE4D TR-FRET Assay Kit	60700	96 rxns.
PDE1A Assay Kit	60310	96 rxns.
PDE1B Assay Kit	60311	96 rxns.
PDE1C Assay Kit	60312	96 rxns.
PDE2A Assay Kit	60320	96 rxns.
PDE4A Assay Kit	60340	96 rxns.
PDE5A Assay Kit	60350	96 rxns.
PDE7A Assay Kit	60370	96 rxns.
PDE7B Assay Kit	60371	96 rxns.
PDE10A Assay Kit	60400	96 rxns.
PDE11A Assay Kit	60411	96 rxns.