

PolyamineRED <Intracellular Polyamine Detection Reagent>

Catalog NO. FDV-0020

Research use only, not for human or animal therapeutic or diagnostic use.

This product has been commercialized with the support of Biofunctional Synthetic Chemistry Laboratory, Cluster for Pioneering Research, RIKEN

日本語版はこちらから
ダウンロードできます。

①弊社ウェブサイトより
Webページ番号検索にて
【70873】で検索

②QRコードより



Product Background

The polyamine species (Figure 1), including putrescine, spermidine and spermine etc. and its acetyl derivatives, are one of the essential class of metabolites which have linear alkyl structure with two or more amines. Polyamines are found in all living organisms with high concentration, from sub-millimolar to millimolar, in the cells. Polyamines have polycationic properties and shows an enormous number of biological functions. For example, polyamines interact with DNA/RNA in the nuclear and regulate gene expression. Polyamines also interact with negatively charged proteins and control its function. The major source of polyamines is an amino acid ornithine. In the case of mammalian, ornithine is converted to putrescine by ornithine decarboxylase (ODC), followed by synthesizing spermidine and spermine. Because ODC is highly expressed in cancer cells, polyamines are considered as one of the cancer marker. Several detection methods of polyamines are developed to date but most of the methods are commonly low-throughput systems using HPLC with polyamine standard compounds. To clear biological functions of polyamines in the cells, the cell-based assay with easy- and high-throughput-procedures is desired.

PolyamineRED is the world first reagent for detecting intracellular polyamines without any pre-treatment and cell lysis. PolyamineRED is a TAMRA (tetramethylrhodamine)-conjugated derivative of glycine propargyl ester which specifically reacts with linear primary alkylamine but not react with secondary amines, bulky amines including amino acids nor monoamines. PolyamineRED has cell-penetrating properties, specifically reacts with polyamines inside the cells and labelled polyamines with red fluorescent dye TAMRA. (Figure 2).

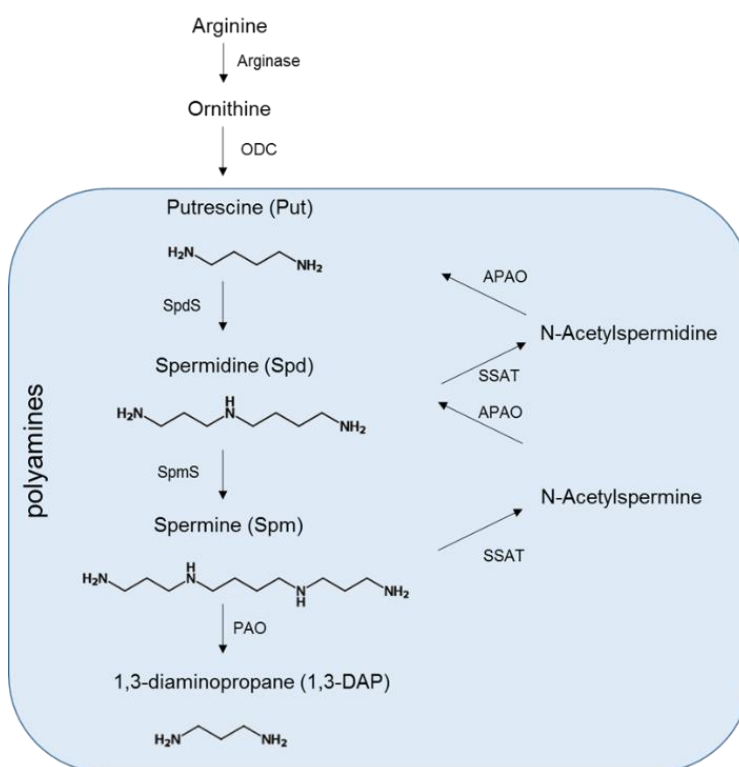


Figure 1. Major polyamine species

Description

Catalog Number: FDV-0020
Size : 0.5 mg
Molecular weight : 611 g/mol
Solubility : Soluble in DMSO
Fluorophore : TAMRA
(red fluorescent dye)
Ex/Em: 560 nm/585 nm
*TAMRA filter sets are available.

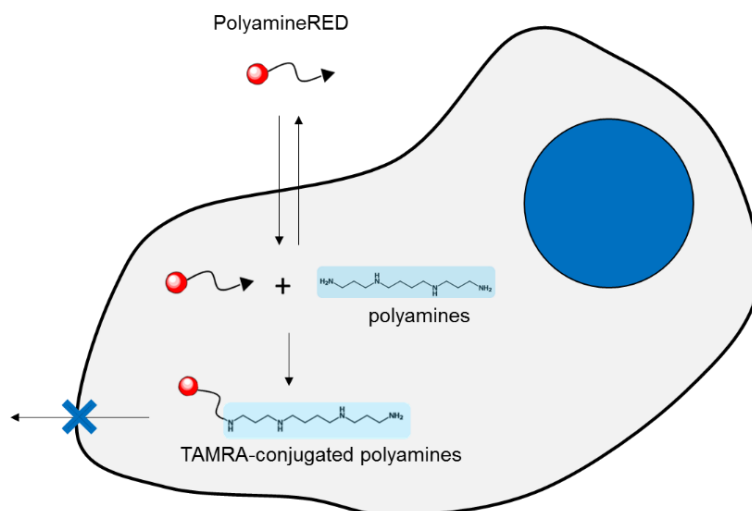


Figure 2. Principle of PolyamineRED

Reconstitution and Storage

Reconstitution : stock solution in 100% DMSO.

Storage (solution) :

Store powder at -20°C.

After reconstitution in DMSO, aliquot and store at -20 °C. Avoid repeated freeze-thaw cycles.

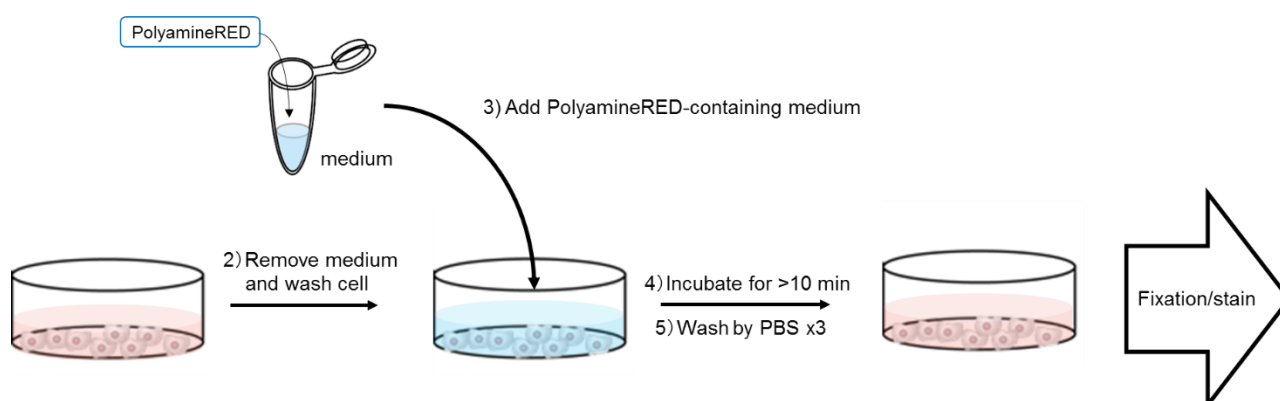
Protect from light.

How to use

General procedure of detection of intracellular polyamines

1. Prepare 10-30 μM PolyamineRED in fresh medium
2. Remove culture medium, wash cells by PBS twice and add PolyamineRED-containing medium to cells
3. Culture cells for at least 10 min
4. Wash cells with PBS 3 times
5. Fixed cells with paraformaldehyde (Option)
Note: MeOH-fixation is not available. Please fix cells by formaldehyde.
6. Additional staining such as DAPI staining or immunocytochemistry with antibodies of interest are available.

1) Preparation of PolyamineRED-containing medium



Reference data

Selectivity of glycine propagyl ester to polyamines

Benzyloxycarbonyl glycine propagyl ester as a model molecule was selectively reacted with polyamines. Reactant of epinephrine, an example of monoamine, and lysine, an example of amino acid, were rarely detected. Reactivity for polyamines depends on the length of polyamine and double linkage products were observed from spermine (4 amino groups) and spermidine (3 amino groups).

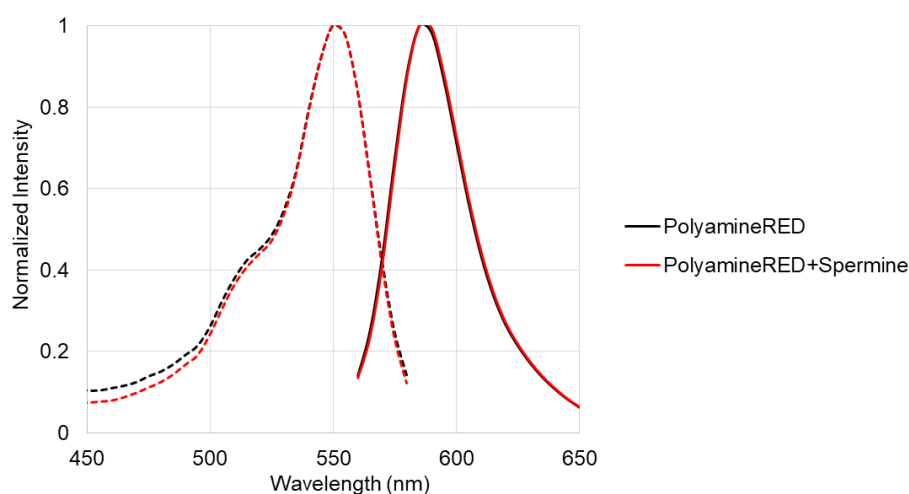
Table Selectivity of glycine propagyl ester to biological amines

amine	Reaction products			Hydrolysis product	Non reacted product
	Total	Single linkage	Double Linkage		
Spermine	82%	59%	23%	17%	1%
Spermidine	78%	67%	11%	21%	1%
Putrescine	66%	66%	<1%	22%	7%
Epinephrine	<1%	<1%	<1%	7%	92%
Lysine	2%	2%	<1%	6%	85%

*This data was cited from Ref.1

Absorption and Fluorescent spectrum of PolyamineRED

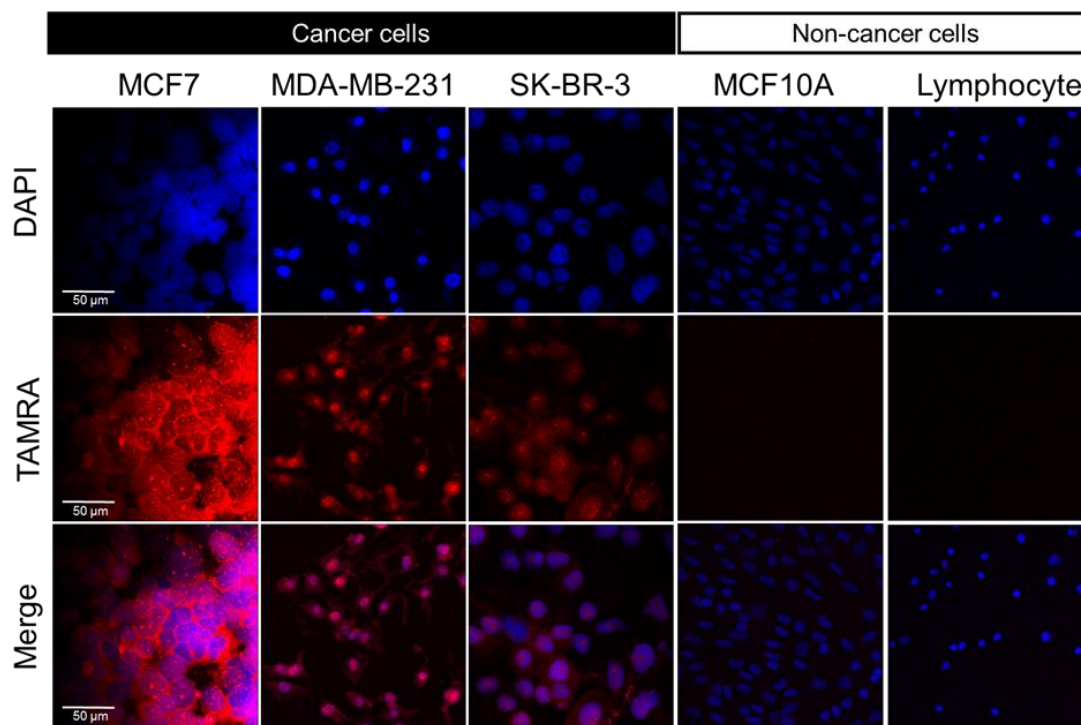
Absorption (dash line) and emission (solid line) of PolyamineRED (Black) and PolyamineRED with spermine (Red) in PBS.



Application data

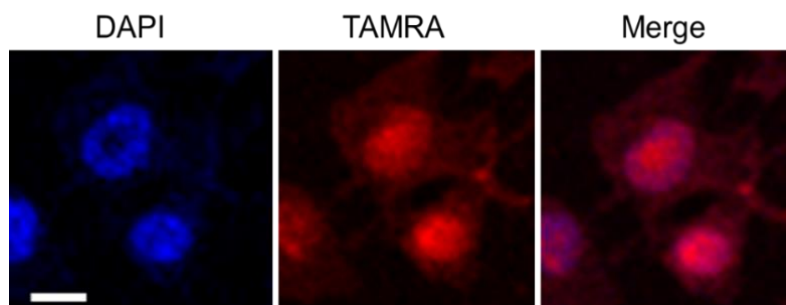
Polyamine imaging in both cancer and non-cancer cells by PolyamineRED

Three cancer cell lines (MCF7, MDA-MB-231 and SK-BR-3) and two non-cancer cells (MCF10A and human lymphocyte) were treated with 30 μ M of PolyamineRED for 10 min. After incubation, cells were washed three times by PBS, followed by DAPI staining and formalin fixation. Images were obtained at Ex/Em=560 nm/585 nm for TAMRA and at Ex/Em=358 nm/461 nm for DAPI. TAMRA fluorescence was detected in cancer cell lines. On the other hand, incubation with non-cancer cell lines showed little fluorescence.



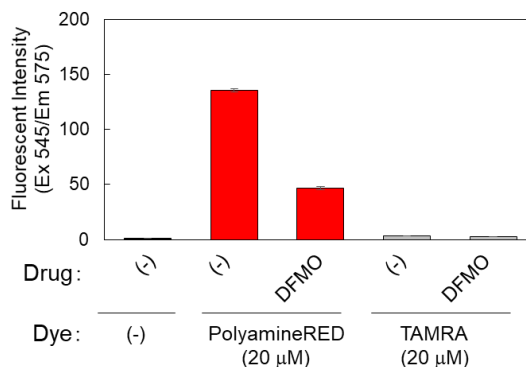
Evaluation of intracellular distribution of polyamines in MDA-MB-231 cancer cell lines

MDA-MB-231 cells were treated with 30 μ M of PolyamineRED for 10 min. After incubation, cells were washed three times by PBS, followed by DAPI-staining and formalin fixation. Images were obtained at Ex/Em=560 nm/585 nm for TAMRA and at Ex/Em=358 nm/461 nm for DAPI. Major TAMRA fluorescent signal was detected from nuclear. This indicates polyamines in MDA-MB-231 are mainly localized in nuclear.



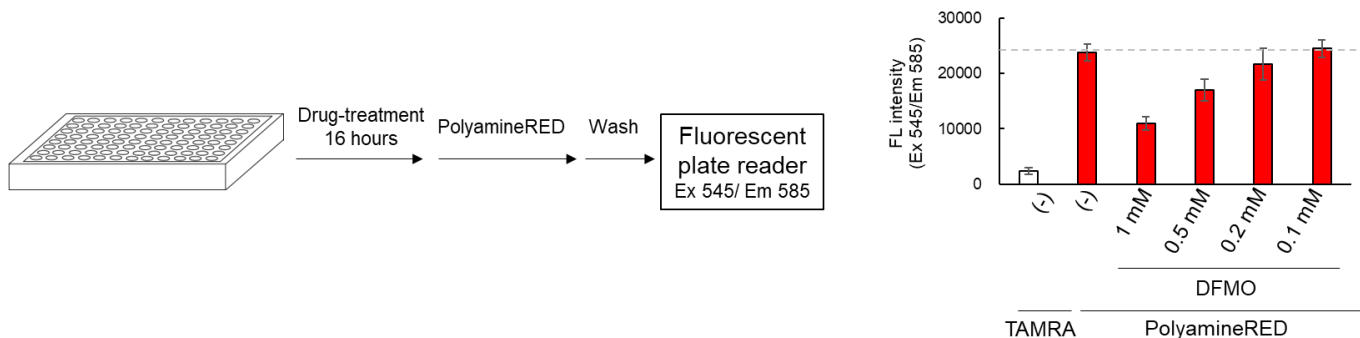
Detection of PolyamineRED signal from cell lysate

786-O cells were seeded in 24 well and culture for 1 day. Confluent cells were treated with DFMO, a ornithine synthase inhibitor; 1 mM, for 16 hours in phenol red-free DMEM (PRf-DMEM) without FBS and further treated with final 20 μ M of PolyamineRED or TAMRA as a negative control for 4 hours. Cells were washed by PBS 3 times and added with 1% SDS-containing PBS. Fluorescent intensity (Ex 540 \pm 10 nm/ Em 585 \pm 20 nm) by the fluorescent spectrophotometer. Fluorescent signals from PolyamineRED-treated cells were significantly higher than that of TAMRA negative control experiment. DFMO-treatment reduced fluorescent intensity.



High-throughput cell-based detection by fluorescent plate reader

786-O cells were seeded in 96 well and culture for 1 day. Confluent cells were treated with indicated drugs for 16 hours in phenol-red free DMEM (PRf-DMEM) without FBS and further treated with final 10 μ M of PolyamineRED or TAMRA as a negative control for 4 hours. Cells were washed by PBS 3 times and added with fresh PBS. Fluorescent intensity (Ex 545 \pm 5 / Em 585 \pm 10) by the fluorescent plate reader with a transparent mode. Fluorescent signals were normalized by background signal of wells of non-dye treated cells. Fluorescent signals from PolyamineRED-treated cells were significantly higher than the TAMRA negative control experiment.



Reference

1. K. Vong, K. Tsubokura, Y. Nakao, T. Tanei, S. Noguchi, S. Kitazume, N. Taniguchi, K. Tanaka, *Chem. Commun.*, **52**, 8403 (2017). Cancer cell targeting driven by selective polyamine reactivity with glycine propargyl esters.

Related product

AcroleinRED <Cell-based Acrolein Detection Reagent>

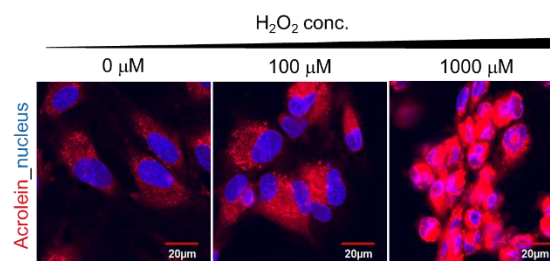
Acrolein is one of the most toxic oxidative stress marker and AcroleinRED is the world first cell-based acrolein detection reagent. As polyamines are one of the major source of acrolein, AcroleinRED and PolyamineRED are good set for oxidative stress research.

Catalog No. FDV-0022

Size 0.5 mg

Features

- Recommended Ex/Em: 560 nm/585 nm
- Easy and quick protocol
- Enable to semi-quantify intracellular acrolein



LipiRADICAL Green <Lipid Radical Detection Reagent>

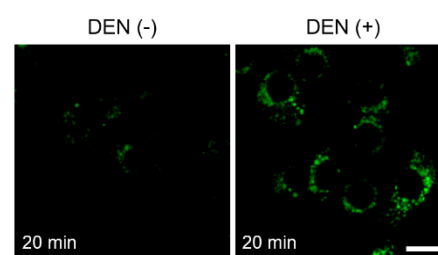
LipiRADICAL Green is a specific fluorescent dye for lipid-derived radicals which are the most upstream factor of lipid peroxidation (LPO). LipiRADICAL Green can be applied into both *in vitro* assay and cell-based assay to monitor lipid radical productions.

Catalog No. FDV-0042

Size 0.1 mg

Features

- Recommended Ex/Em: ~480 nm / 520 nm
- Enable to detect very unstable lipid-derived radicals
- Compatible with *in vitro* assay and in cell-based assay
- An innovative reagent for comprehensive identification of lipid-derived radicals by lipidomics



CellFluor™ GST <Cell-based GST Activity Assay Reagent >

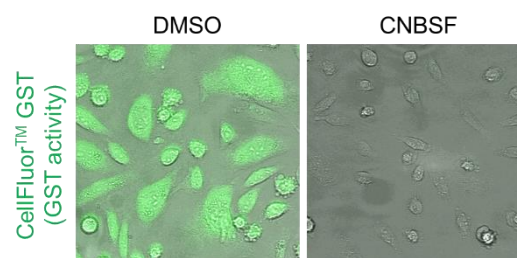
CellFluor™ GST is a novel fluorescent probe for monitoring wide GST members' activity both *in celluo* or *in vitro*. CellFluor™ GST releases green fluorophore rhodamine 110 upon GST activities. This probe has cell-permeability and can detect intracellular GST activity.

Catalog No. FDV-0031

Size 0.1 μmol

Features

- Easy and quick protocol
- Broad specificity for various GST family members
- Ex/Em: 496 nm/520 nm
(Compatible with commercial FITC filters)



Disclaimer/免責事項

This product has been commercialized by Funakoshi Co., Ltd. based on the results of academic research, and the advertisement text, figures and manuals (hereinafter “Product information”) have been prepared based on published research reports on June, 2018. The academic interpretation at the time of creation of the Product Information may change in accordance with future developments in the relevant research field and expansion of various scientific findings, and the latest version and certainty of the Product Information are not guaranteed. The specifications of this product and the Product Information are subject to change without notice. Please contact us for the latest information.

本製品は学術研究成果を基にフナコシ株式会社が製品化したもので、2018年6月時点における公開研究報告を基に広告文章およびマニュアル(以下、製品資料)を作成しています。今後の当該研究分野の発展および各種学術知見の拡大にともない、製品資料作成時の学術的解釈が変更になる可能性があり、最新性・確実性を保証するものではありません。また、本製品の仕様および製品資料を予告なく変更する場合がございます。最新の情報に関しましては、弊社までご確認いただけますようお願い申し上げます。



E-mail Newsletter
Sign Up

Japanese



English



