

# Cellular Function Analysis

6th Edition

Cell Proliferation / Cytotoxicity

Cellular Metabolism

Mitochondria

**Oxidative Stress** 

**Cellular Senescence** 

Lipid droplet

Cell Membrane Dynamics / Morphology

Autophagy

Exosome

Labeling Kit

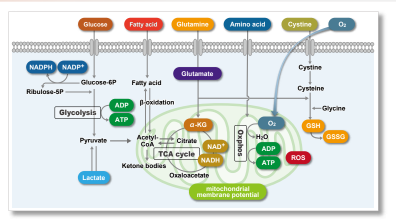


# **Cellular function analysis**

# Intracellular Metabolism ... p.9

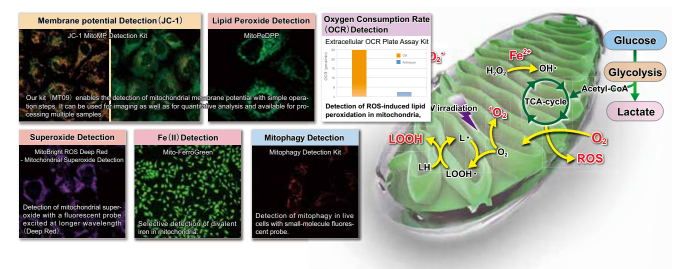
Understanding the intracellular metabolism, such as glycolysis, the TCA cycle and mitochondrial electron transport, is essential for analyzing the status of the cell.

We offer the ready-to-use kits and reagents to analyze the intracellular metabolism with various indicators evaluating the metabolites and nutrients uptake, including glucose, lactate, NAD(P)/NAD(P)H, and glutamine/cystine, as well as the mitochondrial function, including mitochondrial membrane potential, and OCR (Oxygen Consumption Rate).



#### **Mitochondria** ... p.18

Dojindo offers a broad range of reagents and kits for the research on mitochondria, including Mitophagy Detection, Oxygen Consumption Rate Assay, ROS Assay/Detection Kit, as well as the Assay/Detection of lipid peroxidation caused by the ROS. Mitochondrial membrane potential (MtMP) dependent fluorescent probes for mitochondria staining are also available.

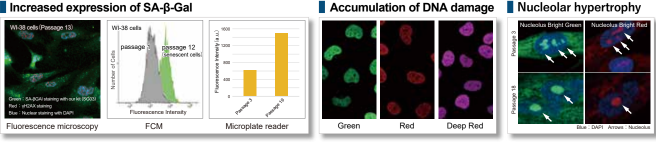


# Cellular Senescence ... p.28

Understanding the apoptosis, necrosis, autophagy and cellular senescence is critical for Cell Function Analysis. Especially, the cellular senescence has been attracting the attention in various fields since the report of the SASP (senescence-associated secretory phenotype), and the stem cell senescence.

We offer four types of kits and reagents with different indicators and methods.

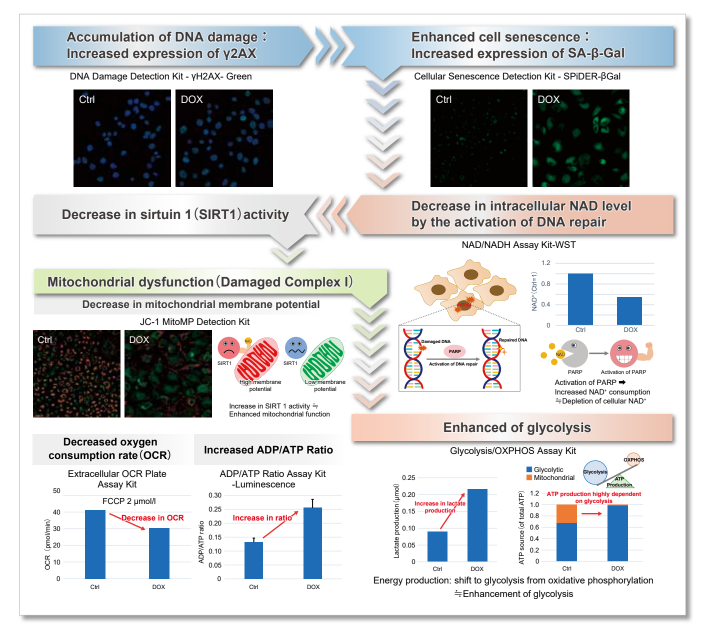
#### Increased expression of SA-β-Gal



# Experiments using multiple indicators

## Observing the intracellular metabolic changes associated with cellular senescence

In cellular senescence induced cells, an increased expression of SA-β-gal and irreversible cell-cycle arrest are observed, and those cells with accumulation of DNA damage shift the energy metabolism to glycolysis as the consequence of the decline in mitochondrial function. The changes in cell function were evaluated with doxorubicin (DOX) induced senescent A549 cells by conducting the experiments using multiple indicators.



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litochondria Research			
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# **Cell Proliferation / Cytotoxicity**

# **Cell Proliferation / Cytotoxicity Assay**

Cell Counting Kit-8

This kit uses the WST method developed by Dojindo Laboratories to measure living cells using intracellular metabolic activity as an indicator. It is widely used in cell proliferation tests and cytotoxicity tests due to its ease of measurement, long-term stability of reagents, and high reproducibility.

DOJINDO MOLECULAR TECHNOLOGIES, INC.

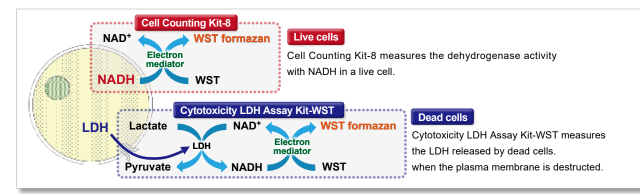
# Cytotoxicity Assay

Cytotoxicity LDH Assay Kit-WST

Plate Reader

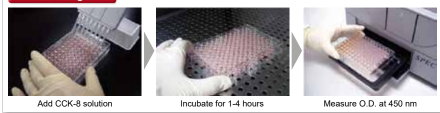
This kit measures cytotoxicity by measuring lactate dehydrogenase (LDH) activity released from cells into the medium. The LDH assay method is increasingly being used in conjunction with Cell Counting Kit-8 (WST method) and MTT method when measuring cytotoxicity.

# Detection Principle

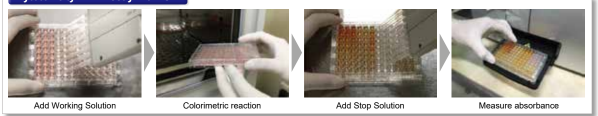


# Procedure

## Cell Counting Kit-8



### Cytotoxicity LDH Assay Kit-WST



Description	Unit	Code
	1,000 tests	
Cell Counting Kit-8	3,000 tests	 CK04
	10,000 tests	
Cytotoxicity LDH Assay Kit-WST	100 tests	
	500 tests	CK12
	2,000 tests	

Exosome

Ā



# **Cell Proliferation / Cytotoxicity Assay Kit**

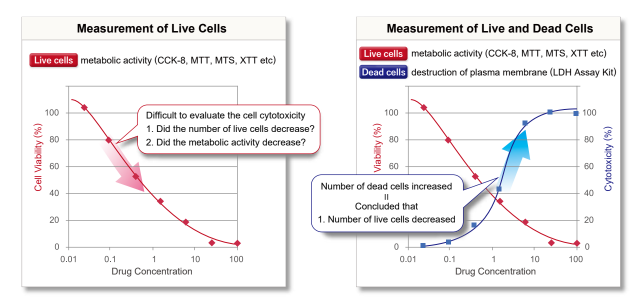
Viability/Cytotoxicity Multiplex Assay Kit



A set of Cell Counting Kit-8 and Cytotoxicity LDH Assay Kit-WST (500 tests each)

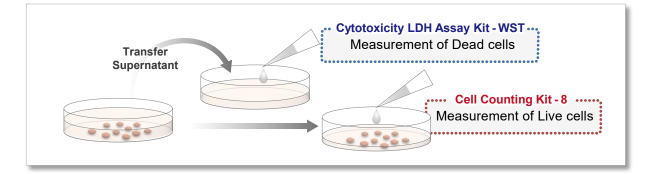
## Purpose of Using Two Detection Methods

Since Cell Counting Kit-8 (CK04) measures the metabolic activity of live cells, the data does not specifically verify cell death. To ensure the data is reflecting the cell death instead of decreased metabolic activity, Cytotoxicity LDH Assay Kit-WST (CK12) is used to increase the data reliability by measuring LDH released from dead cells.



## The same cell sample can be applied

This kit includes CCK-8 for measuring live cells and LDH Assay Kit for measuring dead cells, both of which can be evaluated by plate assay at the same absorbance wavelength. And since same samples can be used for Cell Counting Kit-8 and Cytotoxicity LDH Assay Kit-WST, it is convenient and time-efficient.



Description	Unit	Code
Viability/Cytotoxicity Multiplex Assay Kit	500 tests	CK17

Exosome

# **Cell Count Normalization**

Cell Count Normalization Kit

This kit contains a nuclear staining dye (Hoechst 33342) and an optimized buffer for plate assay, which together allow you to easily determine the cell number per well just by adding the reagent.

Plate

Reader

## Necessity of cell count normalization

When cells are analyzed in a microplate, the results obtained may differ depending on the number of cells per well. In such cases, normalization of the measured values will be necessary.



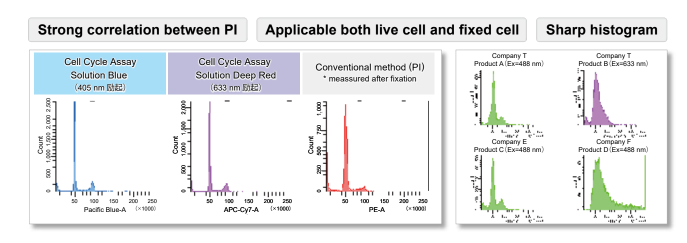
# **Cell Cycle Measurement**



Cell Cycle Assay Solution Blue / Deep Red

## Clearly Identifies Cell Cycle Stages

Live CHO cells stained by the Cell Cycle Assay Solution Blue and Deep Red were measured by flow cytometry. Similar experiments were performed using the existing reagent for cell cycle analysis and PI staining a widely used staining technique. The results obtained by the Cell Cycle Assay Solution were equivalent to PI staining results (shown below). Compared to four different products, our product obtained a sharp histogram peak in live cells.



Description	Unit	Code
Cell Cycle Assay Solution Blue	50 tests	C549
Cell Cycle Assay Solution Deep Red	50 tests	C548

Stress

Autophagy

Cell

# **Dead Cell Staining**

Dead Cell Makeup Blue / Deep Red - Higher Retention than Pl

FCM

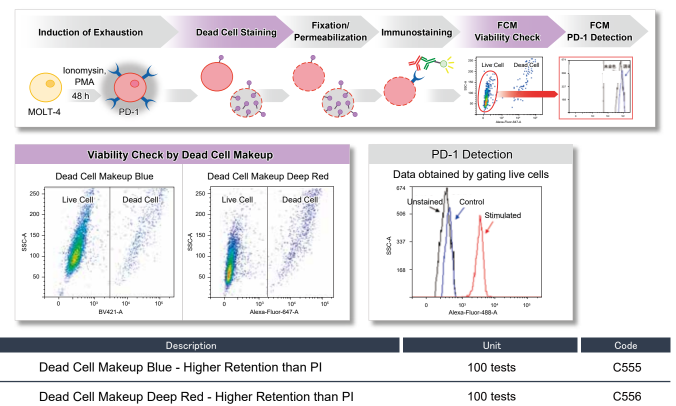
Propidium iodide (PI) is used to discriminate dead cells, but fixation or permeabilization of the membrane causes PI to leak from the cells, making it difficult to obtain accurate data. Dead Cell Makeup has the property of covalently binding to proteins on the cell surface and inside the cells, so the dye does not leak out even after fixation and permeabilization of the cells. In addition, there is a significant difference in fluorescence intensity between live and dead cells fixation, allowing to easily distinguish dead cells and exclude them from the Flow Cytometry analysis.

# Comparison with PI

PI	Dead Cell Makeup series Blue / Deep Red	
<ul> <li>Non-permeable Dye</li> <li>Intercalates into DNA/RNA in the nucleus</li> </ul>	<ul> <li>Non-permeable Dye</li> <li>Covalently binds to cell-surface and intracellular protein</li> </ul>	
<ul> <li>Commonly used for dead cell staining</li> <li>Low cost</li> </ul>	<ul> <li>Enter into dead cell and show high fluorescence intensity.</li> <li>Does not leak from cells</li> </ul>	
Excitation: 561pm / Emission: 586/15 pm (PE)	Blue Excitation: 405nm / Emission: 450/50nm (Pacific Blue, Brilliant Violet 421, Alexa Fluor 405)	
	Deep Red Excitation: 640nm / Emission: 670/30nm (APC, Alexa Fluor 647, Cy5)	
PI leaked from dead cells enters into live cells through its compromised membrane.	Dead Cell Makeup Dyes covalently binds to the protein, hence does not leak even after permeabilization.	
	Non-permeable Dye     Intercalates into DNA/RNA in the nucleus     Commonly used for dead cell staining     Low cost  Excitation: 561nm / Emission: 586/15 nm (PE)  PI leaked from dead cells enters into live cells through	

# PD-1 detection after induction of MOLT-4 cell exhaustion

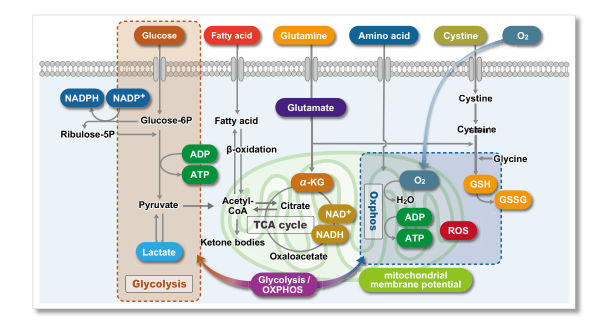
MOLT-4 cells were stimulated for 48 hours in RPMI medium containing Ionomycin (500 ng/ml) and PMA (Phorbol 12-myristate 13-acetate, 50 ng/ml). Cells were stained with Dead Cell Makeup, and PD-1 expression was detected by immunostaining (primary antibody: anti-PD-1 mouse antibody, secondary antibody: anti-mouse antibody-Alexa488). The results showed that dead cells and live cells could be clearly discriminated, and that PD-1 expression was predominantly elevated in the stimulated cell group when only live cells were gated.



# Intracellular Metabolism

Analyzing the various intracellular metabolic pathways [e.g., the glycolysis, the tricarboxylic acid (TCA) cycle, electron transport chain, etc.] is important when trying to understand cellular status. Metabolites and energy sources [e.g., glucose, lactate, and NAD(P)+/NAD(P)H] are the indicators used for analyzing intracellular metabolisms.

We offer a variety of assay kits to evaluate these indicators with necessary components for sample preparation and measurement.



Description	Unit	Code
Starter Kit		
Glycolysis/OXPHOS Assay Kit	50 tests	G270
Glycolysis/JC-1 MitoMP Assay Kit	50 tests	G272
Quantification for Intracellular Metabolism		
ATD Assay Kit Luminossanos	50 tests	A550
ATP Assay Kit-Luminescence	200 tests	A550
ADP/ATP Ratio Assay Kit-Luminescence	100 tests	A552
	50 tests	G264
Glucose Assay Kit-WST	200 tests	G264
Glutamine Assay Kit-WST	100 tests	G268
Glutamate Assay Kit-WST	100 tests	G269
α-Ketoglutarate Assay Kit-Fluorometric	100 tests	K261
	50 tests	L256
Lactate Assay Kit-WST	200 tests	L256
NAD/NADH Assay Kit-WST	100 tests	N509
NADP/NADPH Assay Kit-WST	100 tests	N510
Uptake Assay Kit		
Glucose Uptake Assay Kit-Blue	1 set	UP01
Glucose Uptake Assay Kit-Green	1 set	UP02
Glucose Uptake Assay Kit-Red	1 set	UP03
A A	20 tests	UP04
Amino Acid Uptake Assay	100 tests	UP04
Cystine Uptake Assay Kit	20 tests	UP05
	100 tests	UP05
Fatty Acid Uptake Assay Kit	100 tests	UP07

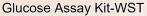
Mitochondria

Cell Membrane

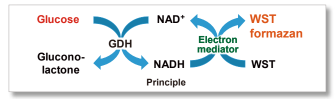
Exosome

#### Glucose

# Glucose Metabolism Assay



Glucose Assay Kit-WST is a colorimetric assay kit for quantification of glucose, one of the major indicators of energy metabolism (lower limit: 0.02 mmol/l). Glucose quantification is used as an indicator of glucose metabolism in diabetes research, as well as to monitor the changes in cellular metabolism in cancer research.



# Lactate

# Lactate Measurement

Lactate Assay Kit-WST

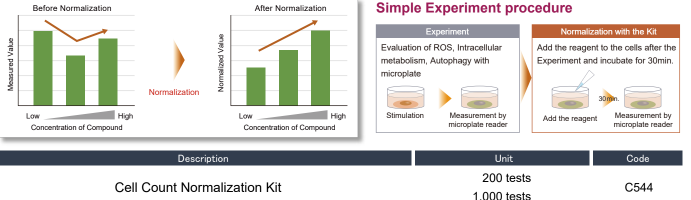


Intracellular Metabolism

L256

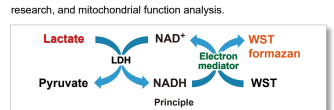
# - 178

# measured values will be necessary. With this kit, by simply adding the reagent to the cell culture medium, the nuclei within the cells are stained, and the number of cells can be easily evaluated from the obtained fluorescence intensity. Before Normalization After Normalization Simple Experiment procedure



When cells are analyzed in a microplate, the results obtained may differ depending on the number of cells per well. In such cases, normalization of the

10



Lactate Assay Kit-WST is a colorimetric assay kit for quantification

of lactate, a metabolite of glycolysis (lower limit: 0.02 mmol/l).

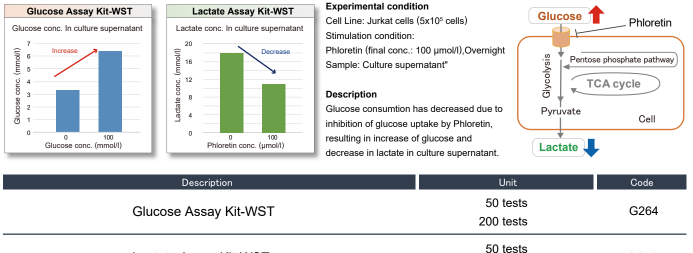
Lactate quantification is widely used in cancer research as an

indicator to monitor changes in cellular metabolic pathways. Recently, there are an increasing number of reports using lactate

as an indicator in areas such as stem cell differentiation, diabetes

# Evaluation of culture supernatant using two indicators

Phloretin, the glucose transporter inhibitor, was added to Jurkat cells and the intracellular metabolism change was evaluated using Glucose Assay Kit-WST and Lactate Assay Kit-WST.



Lactate Assay Kit-WST

(p.7)

**Cell Count Normalization** 



200 tests

# NAD+ NAD<sup>+</sup>/NADH Assay

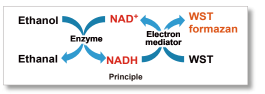


#### NAD/NADH Assay Kit-WST



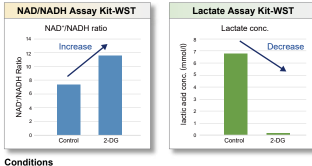
NAD/NADH Assay Kit-WST is a colorimetric assay kit for quantification of NAD+ and NADH and measuring their ratio

NAD is an important cofactor involved in redox reactions in the main metabolic pathways in cells such as glycolysis, electron transfer system and TCA cycle. Maintaining appropriate levels of its oxidized form NAD+ and its reduced form NADH is essential for cell function. Additionally, recent studies report the relations between the decreased level of NAD+ and Senescence.

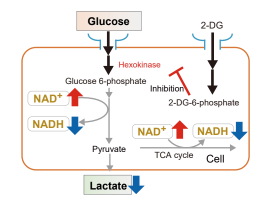


## NAD+/NADH in combination with Lactate Assay Kit

2-Deoxy-D-glucose was added to HeLa cells. After 24 hours of incubation, lactate levels in the supernatant were quantified using the Lactate Assay Kit-WST (Code L256), and the NAD+/NADH ratio was determined with the cell pellet after removing the supernatant using the NAD/NADH Assay Kit-WST.



Cell Line: HeLa cells (1x106 cells) Stimulation condition: 2-DG (final conc.: 6mmol/l), 24 hrs Sample: Culture supernatant (Lactate) Cell (NAD+/NADH ratio)



As a result, intracellular glycolysis was inhibited by 2-Deoxy-D-glucose, which led to decreased lactate levels and an increase in the NAD\*/NADH ratio

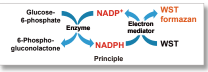
# **NADP<sup>+</sup>/NADPH** Assay NADP/NADPH Assay Kit-WST



NADP\*/NADPH Assay Kit-WST is a colorimetric assay kit that enables quantitation of the amount of total NADP+/NADPH. NADPH and NADP+, and measurement of their ratio.

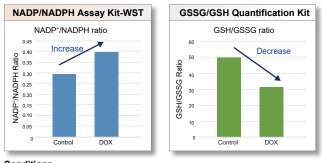
NADP is an important cofactor involved in reactions in the pentose phosphate pathway, one of the metabolic pathways in cells. NADP exists as an oxidized form NADP\* and a reduced form NADPH in cells and involved in biosynthesis of fatty acids and cholesterol as well as generation of reduced glutathione (GSH). In addition, recent study suggests that NADP+/NADPH is associated with the extension of life

span by carbohydrate restriction.



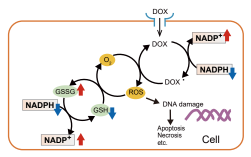
## Measurement of NADP\*/NADPH in combination with **Glutathione (GSH) Quantification Kit**

Doxorubicin (DOX) was added to Jurkat cells. After 24 hours of incubation, NADP+/NADPH ratio and reduced/oxidized glutathione (GSH/GSSG) ratio were determined using the NADP/NADPH Assay Kit-WST and the GSSG/GSH Quantification Kit (Code G257).



#### Conditions

Cell Line: Jurkat cells (3x106 cells) Stimulation condition: DOX (final conc.: 500 nmol/l), 24 hrs Sample: Cell



The results shown above are likely to be explained by the following mechanism. When DOX (doxorubicin) was added to cells, DOX radicals, along with NADP+, were generated by enzymatic reaction. DOX radicals form reactive oxygen species (ROS), which induces DNA damage and apoptosis. In the meantime, to eliminate ROS formed in cells, GSH is consumed and GSSG is increased. Moreover, NADPH is used to reduce GSSG to GSH, resulting in an increase in NADP\*.

Description	Unit	Code
NAD/NADH Assay Kit-WST	100 tests	N509
NADP/NADPH Assay Kit-WST	100 tests	N510

Autophagy

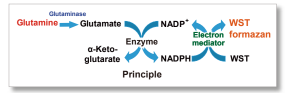
#### Glutamine

## **Glutamine Assay**

#### Glutamine Assay Kit-WST

Glutamine Assay Kit-WST is a colorimetric assay kit that enables quantification of glutamine present in cell culture supernatant or intracellular glutamine (Lower limit: 5µmol/I).

Glutamine is a source of  $\alpha\text{-ketoglutarate}$  (one of the TCA cycle intermediates). It is reported that Glutaminolysis, the process by which glutamine as a substrate is converted to α-ketoglutarate, is upregulated in cancer cells and largely contributed to scavenge the reactive oxygen species (ROS) and reduction of oxidized glutathione.



## Measurement of Glutamine/Glutamate level

A549 cells were seeded in 6-well plates and the concentration of glutamine and glutamate in culture supernatant were measured over time using the Glutamine Assay Kit-WST and Glutamate Assay Kit-WST.

The results showed that the concentration of glutamine in culture supernatant decreased with time, while the concentration of glutamate increased with time.

Glutamate

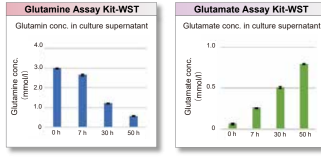
Glutamate Assay

glutamate (Lower limit: 5µmol/l).

Glutamate

Alzheimer's disease.

Glutamate Assay Kit-WST



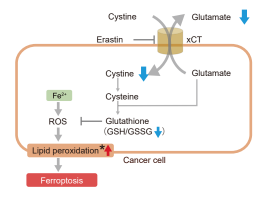
Conditions : Cell Line: A549 cells (5x10<sup>5</sup> cells) Incubation: 50 hrs

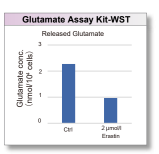
## Induction of Ferroptosis by Erastin

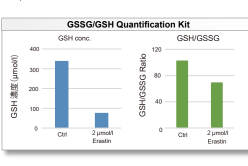
Erastin is a known inducer of ferroptosis by inhibiting the Cystine/Glutamate transporter (xCT).

Released Glutamate and intracellular Glutathione in erastin-treated A549 cells were measured using Glutamate Assay Kit-WST and GSSG/GSH Quantification Kit (Code G257).

As a result, decrease in release of Glutamate and intracellular Glutathione by inhibition of cystine uptake was obseved in erastin treated cells .







#### Conditions

Cell Line: Jurkat cells (3x10<sup>6</sup> cells) Stimulation condition: DOX (final conc.: 500 nmol/l), 24 hrs

Sample: Culture supernatant (Glutamate) Cell (GSH/GSSG)

Description	Unit	Code
Glutamine Assay Kit-WST	100 tests	G268
Glutamate Assay Kit-WST	100 tests	G269

Glutamin Cytosol Glutamate Glutamate Glutamin Mitochondria Gluta nate Acetvl CoA TCA cycle α-ketoglutarate Electron transport chair idative phosphorylation Glutaminolysis

formazan Electron Enzyme mediator α-Keto-NADPH WST glutarate Principle

Glutamate Assay Kit-WST is a colorimetric assay lit that enables you to

quantify glutamate present in cell culture supernatant or intracellular

Glutamate contributes to the biosynthesis of protein and glutathione. It also functions as important neurotransmitter, and an excess glutamate is

considered to be a cause of neurodegenerative disorders, including

A recent study showed that an iron-dependent cell death called "ferroptosis" is induced by inhibition of cystine/glutamate antiporter (xCT) responsible for cystine uptake and glutamate release. As such, cancer research which targets xCT has also been conducted in recent years.

NADP<sup>+</sup>

WST

Intracellular Metabolism

α-KG

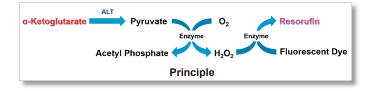
## α-Ketoglutaric Acid Measurement α-Ketoglutarate Assay Kit-Fluorometric



 $\alpha$ -Ketoglutarate Assay Kit-Fluorometric is a fluorometric assay kit for quantification of intracellular  $\alpha$ -ketoglutarate ( $\alpha$ -KG).

 $\alpha$ -KG is an important intermediate of the TCA cycle. It is used as an indicator of enhanced uptake of glucose metabolites into TCA cycle and of enhanced glutaminolysis, a pathway that supplies  $\alpha$ -KG using glutamine as a substrate.

 $\alpha$ -KG has been attracting attention as a metabolic indicator since it plays an essential role in the production of glutamate and  $\gamma$ -aminobutyric acid (GABA), and contributes scavenging the reactive oxygen species.



## ATP Measurement ATP Assay Kit-Luminescence

ATP Assay Kit-Luminescence is a luciferase luminescence assay kit for quantification of intracellular ATP. ATP is an important energy source of living cells that is synthesized in both glycolysis and mitochondrial oxidative phosphorylation.

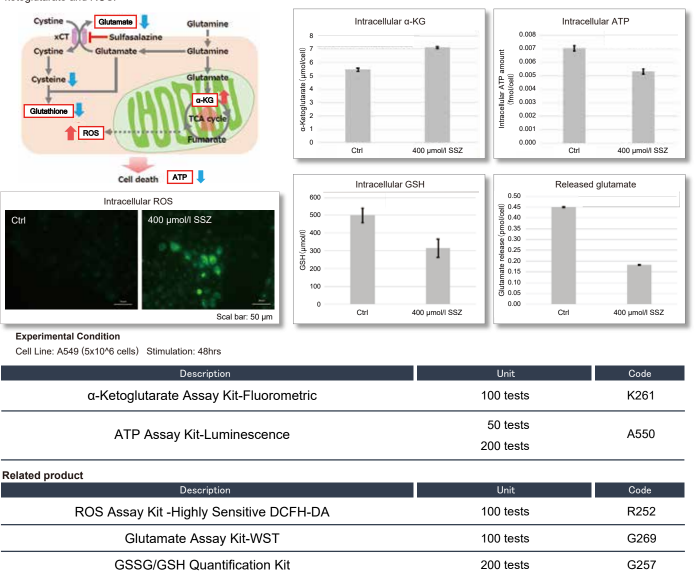
Mitochondrial dysfunction reduces ATP levels in cells and the decreased ATP levels are known to be associated with cancer, aging, and neurodegenerative diseases. Therefore, ATP level is used as an indicator for mitochondrial activity. ATP level is also focused in Cancer research since the recent studies have revealed that although cancer cells were known to rely on glycolysis for ATP synthesis, a shift from glycolysis to oxidative phosphorylation occurs when glycolysis is suppressed.



# Mesurement of α-KG and ATP

Sulfasalazine (SSZ), a known inhibitor of cystine/glutamate transporter (xCT), was added to A549 cells and the changes in intracellular  $\alpha$  -ketoglutarate ( $\alpha$ -KG), ATP, glutathione (GSH), ROS and glutamate release were observed.

The results showed that the addition of SSZ decreased intracellular ATP, glutathione (GSH) and glutamate release, and increased intracellular  $\alpha$ -ketoglutarate and ROS.



Mitochondria

Autophagy

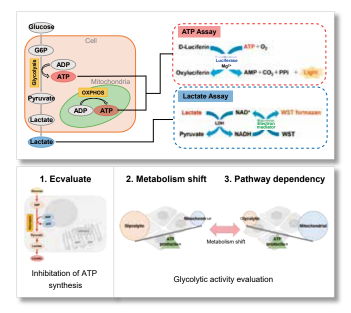
## **Glycolysis/OXPHOS**

#### Glycolysis/Oxidative phosphorylation Assay Glycolysis/OXPHOS Assay Kit

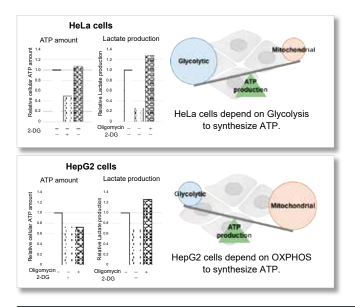


Many cancer cells produce ATP through energy metabolism that depends on the glycolytic pathway. On the other hand, it has been recently reported that cancer cells whose glycolytic pathway is suppressed survive by shifting their energy metabolism to oxidative phosphorylation(OXPHOS). These phenomena has attracted much attention because it will not only help elucidate the mechanisms of anticancer drugs but also lead to the development of therapeutic strategies in various diseases including aging and neurodegenerative disorders.

This kit evaluates glycolytic capacity, metabolism shift, and whether cells rely on the glycolytic system or oxidative phosphorylation for energy production using a plate reader. This ready-to-use kit includes all the reagents required.



#### Comparison of metabolic pathway dependence in different cell lines



#### ATP ADP

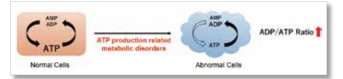
**ADP/ATP Ratio Assay** 



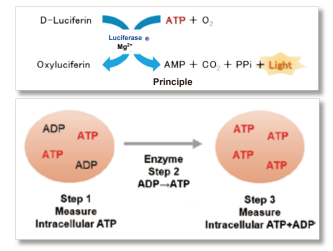
Description	Unit	Code
Glycolysis/OXPHOS Assay Kit	50 tests	G270
ADP/ATP Ratio Assay Kit-Luminescence	100 tests	A552

ADP/ATP Ratio Assay Kit-Luminescence

Normally, when the intracellular ATP production decreases, the degradation product ADP will be resynthesized to ATP to maintain the intracellular ATP concentrations. However, when the metabolism of ATP production is disrupted, ATP cannot be resynthesized from ADP, but intracellular ATP is converted to ADP and the ADP/ATP ratio increases. The change in ADP/ATP ratio is not only related to apoptosis and autophagy, but also intracellular energy metabolism, and is used as one of the indicators of metabolic activity.



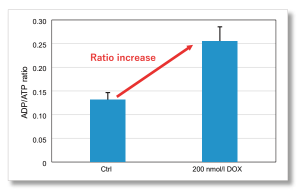
This kit can measure the ratio of ADP to ATP in cells. First, measure intracellular ATP by the luciferase luminescence method.



## ATP/ADP change in senescence-induced cells

We measured the ADP/ATP ratio of A549 cells treated with Doxorubicin (DOX) and untreated A549 cells using this kit.

As a result, it was confirmed that the ADP/ATP ratio of DOX-treated A549 cells increased predominantly, and it was possible to observe the variation of ADP/ATP ratio by senescence induction using this kit.

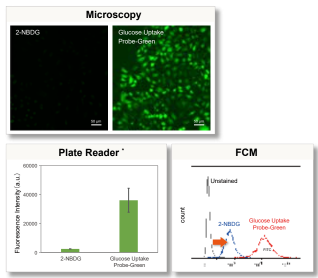


#### Glucose

### Glucose Uptake Assay Glucose Uptake Assay Kit-Blue, Green, Red



Glucose Uptake Probe allowing highly sensitive detection of cellular glucose uptake by fluorescence imaging or flow cytometry. The WI Solution in this kit can enhance cellular retention to provide more reliable experimental data. Also, compared to the existing method (2-NBDG), the measurement time can be significantly reduced.



\* Only Green and Red can be detected with a plate reader.

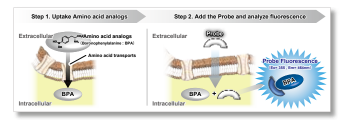
# Amino acid

### Amino Acid Uptake Assay Amino Acid Uptake Assay Kit



Amino acids are essential for intracellular protein and nucleic acid synthesis, Especially for cancer cells which are proliferating continually. Since the supply of acetyl CoA from the glycolytic pathway is decreased in cancer cells, they have a furthermore huge demand for amino acids which are an important nutrient source for the TCA cycle. A research has shown that cancer cells increase the expression of the amino acid transporter LAT1 (L-type amino acid transporter <sup>1)</sup> and take up a large number of amino acids, which is expected to be a new target for anti-Cancer drugs' discovery.

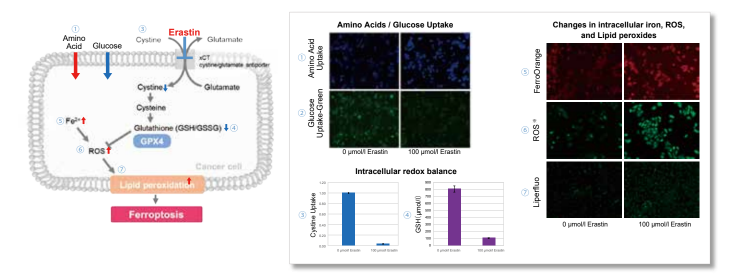
This product is suitable for fluorescence imaging, plate reader, and flow cytometry, making it possible to visualize and quantify the amino acid uptake capacity of cells and is useful for evaluation of amino acids uptake capacity and screening of amino acids transporter inhibitors.



## Extraceller Induction of Ferroptosis by Erastin

The following experimental examples show changes in each aforementioned index as a consequence of erastin stimulation. Measurements were made using Dojindo reagents.

Using erastin-treated A549 cells, we measured intracellular  $Fe^{2+}$ , ROS, lipid peroxide, glutathione, glutamate release into the extracellular space, and cystine uptake. As a result, inhibition of xCT by elastin was observed and also the release of glutamate and uptake of cystine were decreased. Furthermore, elastin treatment decreased intracellular glutathione while it increased intracellular  $Fe^{2+}$ , ROS, and lipid peroxides.



Description	Unit	Code
Glucose Uptake Assay Kit-Blue	1 set*1	UP01
Glucose Uptake Assay Kit-Green	1 set *1	UP02
Glucose Uptake Assay Kit-Red	1 set *1	UP03
Amino Acid Untoko Accov Kit	20 tests *2	UP04
Amino Acid Uptake Assay Kit	100 tests *2	0P04

\*1 <Approximate number of uses per 1 set> 35 mm dish x 10, 96-well plate x 1 \*2 < Approximate number of uses per 100 tests> 35 mm dish x 10, 96-well plate x 1

Mitochondria

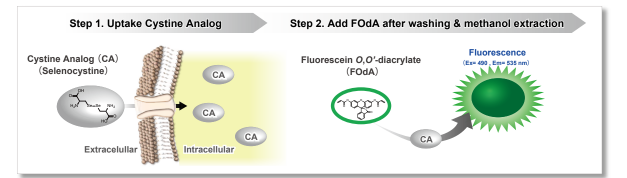
Labeling

Ā



## What you can do with the kit

The Cystine Analog (CA) in this kit can be taken up into cells via xCT, and the incorporated CA can be specifically detected using the Fluorescent Probe and Reducing Agent. Thus, the xCT activity can be measured easily.[Patent pending]



The relevant technicals are published in the following journal: Shimomura T, Hirakawa N, Ohuchi Y, Ishiyama M, Shiga M, Ueno Y, Simple Fluorescence Assay for Cystine Uptake via the xCT in Cells Using Selenocystine and a Fluorescent Probe. ACS Sensors, 2021, 6(6), 2125-2128

Description	Unit	Code
Cystine Uptake Assay Kit	20 tests *	UP05
	100 tests *	0.00

\*<Approximate number of uses per 100 tests> 96-well plate x 1



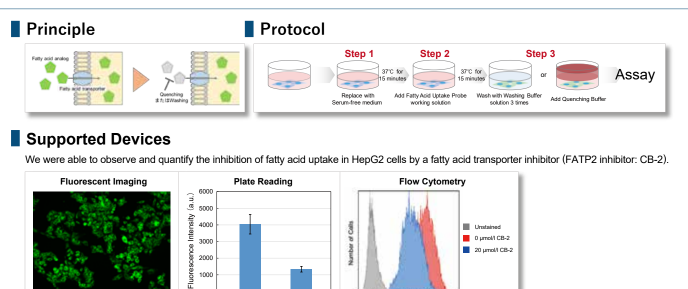
# Fatty Acid Uptake Assay

0 µmol/l CB-2

Fatty Acid Uptake Assay Kit

Micro Plate FCM Reader scop

This kit contains a fatty acid analog (Fatty Acid Uptake Probe) which can be uptaken by cells via fatty acid transporters, and fatty acid uptake ability can be detected by a fluorescence method (Principle). The Quenching Buffer enables detection without cell-washing steps (Protocol).



FITC CB-2 intreated HepG2 cells Description Unit Code **UP07** Fatty Acid Uptake Assay Kit 100 tests \*

20 µmol/l CB-2

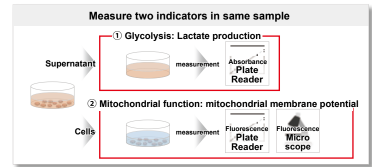
Autophagy

# **Glycolysis / Mitochondria Membrane Potential Assay**

Glycolysis/JC-1 MitoMP Assay Kit

Glycolysis/JC-1 MitoMP Assay Kit enables the measurement of lactate production (via a lactate assay) to detect changes in the glycolysis, as well as the mitochondrial membrane potential (via a JC-1 assay) to assess mitochondrial function, from the same samples, using cell culture supernatant and the cells respectively. The kit includes all necessary reagents for the assays, and a combined protocol is available.

## Simultaneous measurement of the same sample



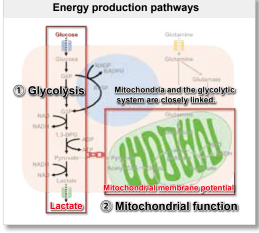


Micro

Plate Reader

## Why monitor glycolysis and mitochondrial function?

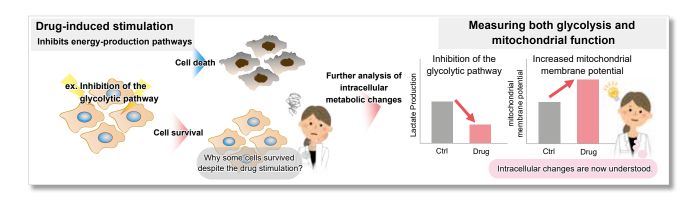
The link between mitochondrial function and cellular metabolism is well-known, with implications for a range of diseases including cancer, aging, and neurodegenerative diseases. It has been found that aging cells often rely on the glycolytic system for survival, rather than drawing on mitochondrial energy sources. Conversely, cancer cells, which typically depend heavily on glycolysis, activate mitochondrial function to ensure their survival even when the glycolytic system is inhibited. Given these observations, there is a growing necessity to investigate both mitochondrial function and the glycolytic pathway to enhance our understanding of intracellular metabolic alterations.



#### Intracellular metabolic changes induced by drug stimulation

#### Simultaneous monitoring of glycolysis and mitochondrial function

In certain instances, cells manage to survive despite sustaining damage to their glycolysis or mitochondrial function, the principal pathways for energy production. It is understood that this occurs as cells strive to persist and prevent cell death by augmenting glycolysis even when mitochondrial function is compromised, or by activating mitochondrial function when glycolysis is impaired. Simultaneously monitoring both the glycolysis and mitochondrial function, as detailed below, can gain insight into what is happening inside the cell.



Description	Unit	Code
Glycolysis/JC-1 MitoMP Assay Kit	50 tests *	G272

17

\* <Approximate number of uses> 96-well plate x 0.5

Mitochondria

\_abeling

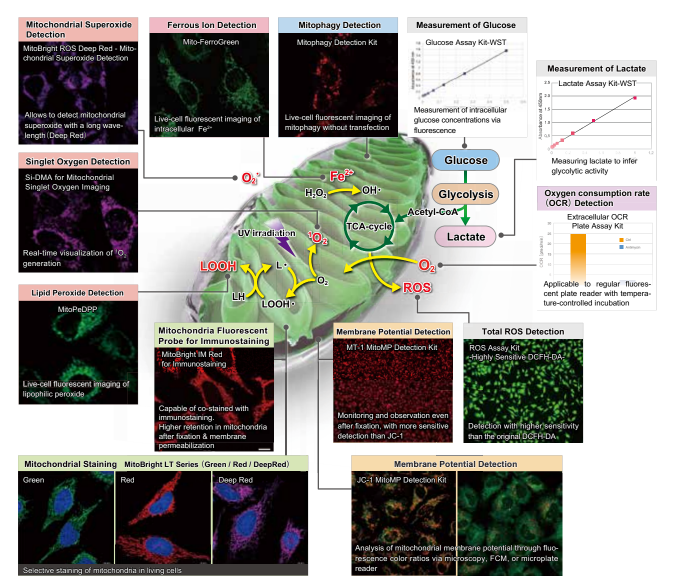
Ā

# **Mitochondria Research**

Mitochondria, which have important functions in cells, are a place for ATP production by oxidative phosphorylation, and their activity and dysfunction are closely related to cancer, senescence, and neurodegenerative diseases such as Alzheimer's disease and Parkinson's disease. Therefore, understanding the state of mitochondria is particularly important.

# Product lineup for Mitochondrial research

Mitochondrial research products include the detection of Mitophagy, which is a quality control system for damaged mitochondria, the detection of ROS: reactive oxygen species (Si-DMA, MitoBright ROS), and the detection of peroxidation caused by oxidation by ROS. Detection of lipid peroxides (MitoPeDPP), as well as various fluorescent probes for mitochondrial staining and membrane potential dependent fluorecent probes (MitoBright LT series Green/Red/Deep Red), MitoBright IM, JC-1, MT-1 ) are available.



# **Mitochondrial Staining**

MitoBright LT Green / Red / Deep Red

MitoBright LT dyes are designed to exhibit mitochondria retention for long-term visualization. In addition, the MitoBright LT dyes show stronger fluorescence signals compared with other commercially available dyes that contain the chloromethyl moiety. The MitoBright LT dyes offer three different color options (Green, Red and Deep Red), and are provided as a ready-to-use DMSO solution. A working solution can easily be prepared in a single dilution step with growth medium or HBSS.

### Stained in serum-contained media

HeLa cells were stained with MitoBright LTs or an existing reagent and observed after 4 days. MitoBright LT remained unchanged and observable even after 7 days, while the existing reagent's intensity decreased.

		MitoBright LT			Company(T)	
	Green	Red	Deep Red	Green	Red	Deep Red
Intial staining	Rey.	( <b>Ö</b> -\$				
Day 4						

Description	Unit	Code
MitoBright LT Green	400 µl *	MT10
MitoBright LT Red	400 µl *	MT11
MitoBright LT Deep Red	400 µl *	MT12

\*<Estimated number of use per 400µl> 35 mm dish x 200

Micro

scope

Micro

scop

FCM

# **Mitochondria Fluorescent Probe**

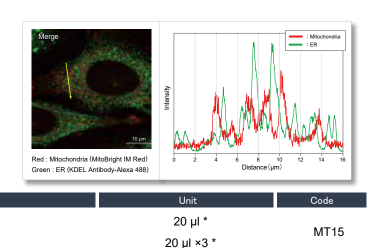
MitoBright IM Red for Immunostaining

MitoBright IM has a structure that allows it to be easily retained in mitochondria even during the immunostaining process after staining live cells. MitoBright IM is a new reagent that overcomes the problem of combined use with immunostaining.

19

#### Clear observation of various organelles

Mitochondria of Hela cells were stained with MitoBright IM, then fixed and permeabilzed. After that, cells were co-stained with immunostaining using KDEL antibody, a marker protein of the endoplasmic reticulum(ER). The fluorescence intensity was measured in the area indicated by the blue arrow in the fluorescence image (right). As a result, the morphology of the mitochondria and the neighboring endoplasmic reticulum was clearly observed.



MitoBright IM Red for Immunostaining

Cytotoxicity

Metabolism

Autophagy

Labeling

Ā

# JC-1 Mitochondrial Membrane **Potential Detection**

JC-1 MitoMP Detection Kit

# **MT-1** Mitochondrial Membrane **Potential Detection**

MT-1 MitoMP Detection Kit

Mitochondria is an important organelle that uses oxygen to synthesize ATP, producing the necessary energy for live cells to thrive. Decreased mitochondrial activity and mitochondrial dysfunction are associated with cancer, aging, and neurodegenerative diseases such as Alzheimer's and Parkinson's disease. Therefore, mitochondrial membrane potential (MMP) has been widely studied as a promising target for mitochondria-related diseases.

Micro

scope

Micro

scope

Plate

Reade

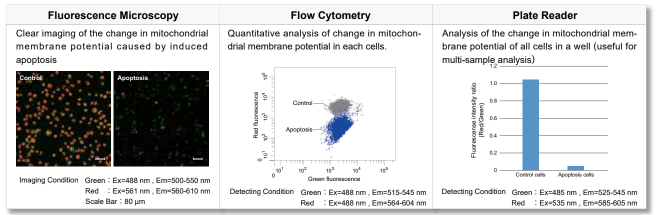
FCM

FCM

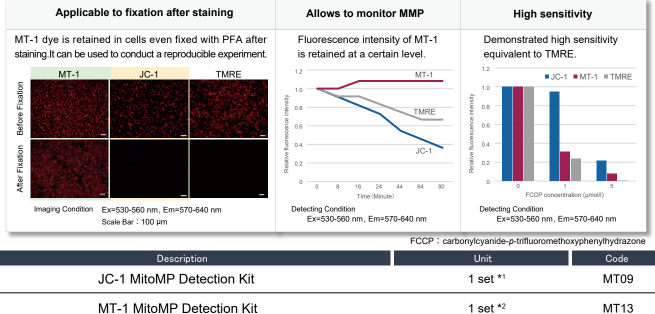
Product	Features	Sensitivity	Fixation	Monitoring	Applicable Instruments
JC-1 MitoMP Detection Kit	Recomended for starting-up	~			Micro scope FCM Plate Reader
MT-1 MitoMP Detection Kit	Recommended for more detailed analysis	イ (High)	$\checkmark$	$\checkmark$	Micro scope FCM

#### Applicable to various instruments JC-1 MitoMP Detection Kit

Jurkat cells treated with Staurosporine, an apoptosis inducing reagent, were stained with JC-1 MitoMP Detection Kit.



#### More detailed analysis MT-1 MitoMP Detection Kit



### MT-1 MitoMP Detection Kit

\*1 < Approximate number of uses (Reagent concentration: 2 µmol/I)> 96 well plate x 5, 35 mm dish x 25, \*2 < Approximate number of uses> 35 mm dish x 30

\_abeling

**MT13** 

# Metabolism

Mitochondria

# Autophagy

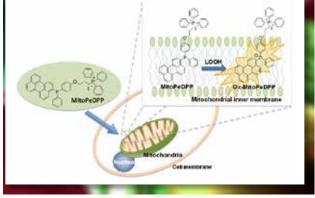
Exosome

# **Mitochondrial Lipid Peroxide Detection**

MitoPeDPP

MitoPeDPP is a fluorescent dye which penetrates cell membrane and accumulated in mitochondria. It is oxidized specifically by the lipid peroxide in mitochondrial membrane and emits strong fluorescence. MitoPeDPP enables the imaging of the lipid peroxide that were oxidized under oxidative stress conditions (ROS, etc.) in mitochondrial membrane. It is important to evaluate the reactive oxygen species (ROS) in mitochondria where oxygen for energy production is consumed. MitoPeDPP is also used to monitor lipid peroxidation in ferroptosis research. \*This probe has been developed by Dr. Shioji et al., Fukuoka University, Department of Chemistry

# Principle



Detection of mitochondrial lipid peroxide using rotenone Additional information is available on our website.

Micro

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0 min	180 min	
	18	
10 A	• 1900 - 64	
Ç.		
MitoPeDPP	1 (F	4
		HONE.

Description	Unit	Code
MitoPeDPP	5 µg ×3 *	M466

 $^{\star}$  <Approximate number of uses per 5  $\mu g >$  96 well plate x 1-5, 35 mm dish x 5-25

# Oxygen Consumption Rate(OCR) Plate Assay Kit

Extracellular OCR Plate Assay Kit

\* OCR : Oxygen Consumption Rate

# Comparison with Flux Analyzer

\* This product was commercialized under the guidance of

Dr. Toshitada Yoshihara, Gunma University

Flux Analyzer (XFe24) and this kit were measured on the same day under the same conditions (cell type, cell number, and FCCP concentration). As a result, correlated data of oxygen consumption rate changes was obtained for XFe24 and this kit.



Cell Type : HepG2 Cell Number : 5×10<sup>4</sup> cells/well FCCP Conc. : 2 µmol/l

The Extracellular OCR Plate Assay Kit includes an Oxygen Probe, which has the property of increasing phosphorescence intensity as the oxygen concentration in the medium decreases, and Mineral Oil blocks the influx of oxygen from the air.

After measuring the phosphorescence intensity according to the extracellular oxygen concentration with a fluorescent microplate reader, the OCR of the cells is calculated (automatic calculation sheet) based on the Stern-Volmer equation.



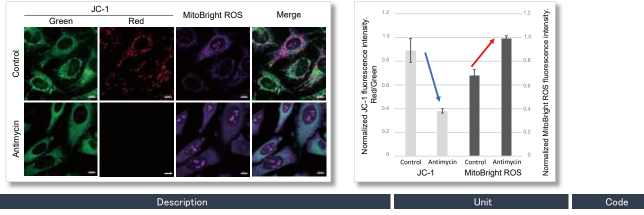
# **Mitochondrial Superoxide Detection**

MitoBright ROS Deep Red - Mitochondrial Superoxide Detection \*Same product as Product code 'MT14'

MitoBright ROS Deep Red selectively reacts with superoxide within mitochondria and emits fluorescence with a longer wavelength than existing products.

# Simultaneous evaluation of mitochondrial superoxide and membrane potential

HeLa cells were washed with HBSS, co-stained with MitoBright ROS Deep Red and mitochondrial membrane potential staining dye (JC-1: code MT09 or MT-1: code MT13), and the generated mitochondrial ROS and membrane potential were observed simultaneously. As a result, the decrease in mitochondrial membrane potential and the generation of mitochondrial ROS were simultaneously observed.



MitoBright ROS Deep Red - Mitochondrial Superoxide Detection \*1 100 nmol \*2

\*1. Same product as Product code 'MT14'. \*2. < Approximate number of uses (Reagent concentration: 2 µmol/l)> 96 well plate x 5, 35 mm dish x 25

# DOINTO MOLECULAR TECHNOLOGUES, INC. Mitochondrial Singlet Oxygen Imaging



Micro

scope

Plate

Reader

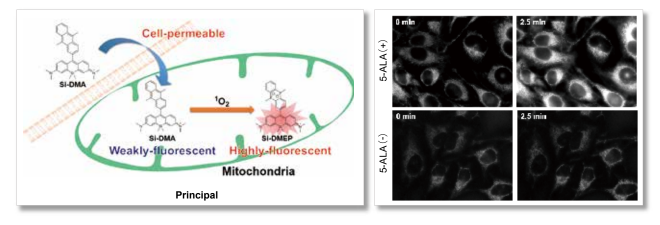
FCM

Si-DMA for Mitochondrial Singlet Oxygen Imaging

Majima et al. synthesized a new far-red fluorescence probe, "Si-DMA", composed of silicon-containing rhodamine and anthracene moieties. Under the presence of  ${}^{1}O_{2}$ , fluorescence of Si-DMA increases 17 times due to endoperoxide formation at the anthracene moiety. Among seven different ROS, Si-DMA can selectively detect the  ${}^{1}O_{2}$ . In addition, Si-DMA was able to visualize the real-time generation of  ${}^{1}O_{2}$  from protoporphyrin IX in mitochondria with 5-aminolevulinic acid (5-ALA).

\* This probe was developed by Dr. Tetsurou Majima et al., Osaka University

# Product Description / Singlet oxygen detection after the addition of 5-ALA



Description	Unit	Code
Si-DMA for Mitochondrial Singlet Oxygen Imaging	2 µg *	MT05

\* < Approximate number of uses > µ-Slide 8 well x 22 to 90

MT16

Labeling Kit

Autophagy

# **Mitophagy Detection**

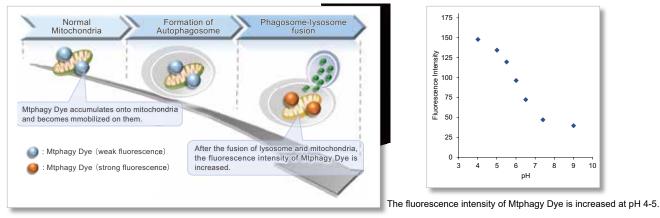
Mitophagy Detection Kit

DOJINDO MOLECULAR TECHNOLOGIES, INC.



## Increased fluorescence intensity during mitophagy

This kit is composed of Mtphagy Dye and Lyso Dye. Mtphagy Dye accumulates in intact mitochondria, and immobilized on it with chemical bond and exhibits a weak fluorescence from the influence of surrounding condition. When Mitophagy is induced, the damaged mitochondria fuses to lysosome and then Mtphagy Dye emits a high fluorescence. To confirm the fusion of Mtphagy Dye–labeled mitochondria and lysosome, Lyso Dye included in this kit can be used for co-staining.



Description	Unit	Code
Mitophagy Detection Kit	1 set *	MD01
Mtphagy Dye	5 µg ×3	MT02

 $^{\star}$  <Approximate number of uses> 96 well plate x 5, 35 mm dish x 25 (Reagent concentration: 2  $\mu mol/l)$ 

# **Mitochondrial Iron Detection**

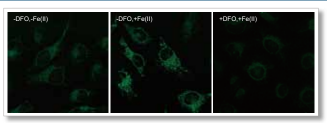
Mito-FerroGreen



Mito-FerroGreen is a novel fluorescent probe for the detection of ferrous ion (Fe2+) in mitochondria where Fe-S clusters and heme proteins are synthesized, and enables live cell fluorescent imaging of mitochondrial Fe2+. Mito-FerroGreen has no no chelating ability. Mito-FerroGreen and Fe2+ react irreversibly, which is different from the detection principle of calcium-iron probes such as Fluo-3. \* This product was commercialized under the advisory of Dr. Hideko Nagasawa and Dr. Tasuku Hirayama (Gifu Pharmaceutical University).

## Detection of mitochondrial Fe2+

By adding Mito-FerroGreen to HeLa cells cultured in MEM growth medium containing serum, intrinsic iron in mitochondria was detected by fluorescence (left figure). On the other hand, strong fluorescence of Mito-FerroGreen was observed in HeLa cells treated with iron (iron (II) ammonium sulfate) (middle figure), while Mito-FerroGreen fluorescence was not observed in cells treated with iron chelating



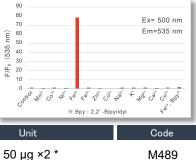
Ex=488 nm, Em=500-565 nm

reagent (DFO: deferoxamine) and iron. Thus, difference in the amount of mitochondrial iron were confirmed as difference in fluorescence intensity. The protocol of this experiment is discribed in the manual (available on our website).

## High selectivity of Fe<sup>2+</sup>

High selectivity of Fe<sup>2+</sup> and high signal are obtained.

Add 2 µl of 1 mmol/l Mito-FerroGreen, 2 µl of 10 mmol/l various metals, and 20 µl of 1 mg/ml esterase to 1 ml of 50 mmol/l HEPES Buffer (pH 7.4) and fluorescence intensity was measured after 1 hour reaction at room temperature.



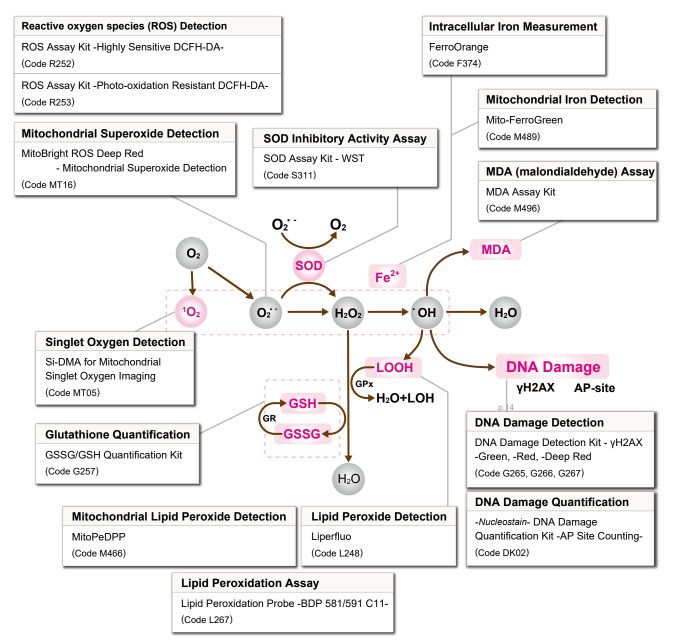
Description Mito-FerroGreen

\* <Approximate number of uses per 50  $\mu g\!>\,\mu\text{-silde}$  8 well x 6

# **Oxidative Stress**

Oxygen is used for energy production and along the process, reactive oxygen species (ROS) are generated. ROS play an essential role in signaling pathways and the immune system, while excess ROS is associated with diseases and cellular senescence. Recent studies suggested that ferroptosis is a new type of cell death characterized by iron dependency and increased ROS. Thus, ROS detection has been attracting considerable interest in ferroptosis research.

# Oxidative stress related products



# **Oxidative Stress**

# **Photo-oxidation Resistant Detection of total ROS**

ROS Assay Kit -Photo-oxidation Resistant DCFH-DA-

The dye that is employed in this kit allows ROS detection with higher sensitivity than DCFH-DA; It does not leak from cells because the fluorescent dye can immobilize protein via a chemical bond, and it is resistant to photo-oxidation compared with DCFH-DA.

Micro

scope

Micro

scope

∎Nevi∎

Fluorescent Plate Reader

Fluorescent

Plate Reader

FCM

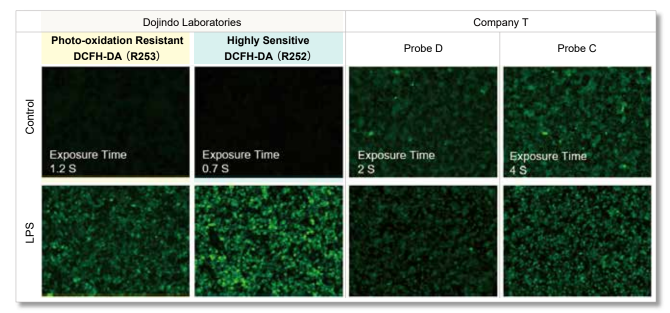
FCM

# **Highly Sensitive total ROS detection**

ROS Assay Kit -Highly Sensitive DCFH-DA-

The dye employed in the kit allows ROS detection with higher sensitivity than DCFH-DA. Moreover, the Loading Buffer included in this kit maintains cellular health during assays.

# Comparison of Fluorescent Sensitivity



# Comparison with Existing Reagents

	Dojindo Laboratories       ROS Assay Kit     ROS Assay Kit       -Photo-oxidation Resistant     -Highly Sensitive       DCFH-DA-     DCFH-DA-		Comp	any T
Product Name			Probe D	Probe C
Photo-oxidation Resistant	O Highest resistant ability	× Photo-oxidation	X Photo-oxidation	$\triangle$ Photo oxidation
Cell fixation	O Highest retention ability	X Leakage from cell	X Leakage from cell	OK for fixation
Sensitivity (Intracellular)	O Better sensitivity	O Highest sensitivity	$\triangle$ Lower sensitivity	$\triangle$ Lower sensitivity
Code	R253	R252	_	_

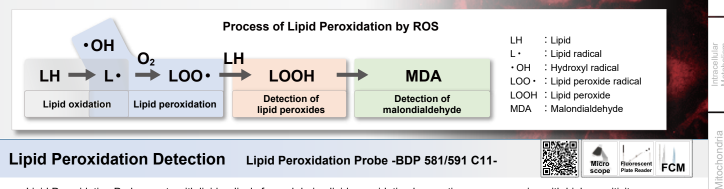
Description	Unit	Code
ROS Assay Kit -Highly Sensitive DCFH-DA-	100 tests	R252
ROS Assay Kit -Photo-oxidation Resistant DCFH-DA-	100 tests	R253

25

Mitochondria

Labeling Kit

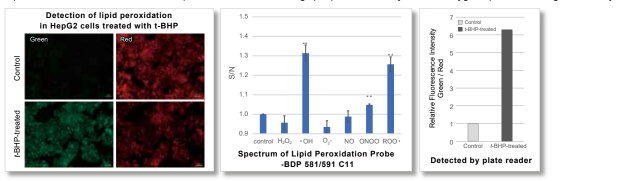
Reactive Oxygen Species (ROS) refers to a variety of reactive molecules derived from molecular oxygen. Various ROS are generated in vivo and each of them differs in its properties, reactivities and interactions, thus, it is essential to select an appropriate reagent to detect and analyze the ROS molecules accurately.



### Lipid Peroxidation Detection

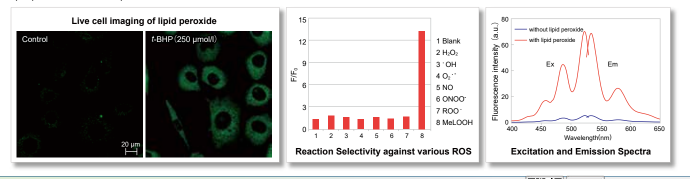
#### Lipid Peroxidation Probe -BDP 581/591 C11-

Lipid Peroxidation Probe reacts with lipid radicals formed during lipid peroxidation by reactive oxygen species with high sensitivity.



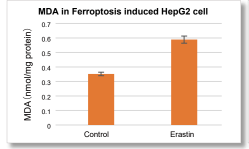
#### Lipid peroxide Detection Liperfluo

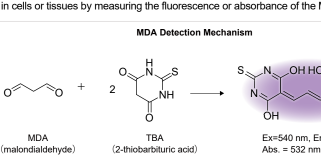
Liperflow can be applied to lipid peroxide imaging by a fluorescence microscopy and a flow cytometric analysis of live cells and is used to monitor lipid peroxidation in ferroptosis research.



## Malondialdehyde (MDA) Detection MDA Assay Kit

MDA Assay Kit uses the TBARS method to detect MDA in cells or tissues by measuring the fluorescence or absorbance of the MDA-TBA adduct.







Micro

Micro scope

Fluorescent Plate Reader

FCM

Labeling Kit

Proliferation Cytotoxicity

**Oxidative stress** 

Lipid droplet

Membrane

Cell

Autophagy

Description	Unit	Code
Lipid Peroxidation Probe -BDP 581/591 C11-	200 tests *1	L267
Liperfluo	1 set (50 µg×5) *2	L248
MDA Assay Kit	100 tests *3	M496

\*1 < Approximate number of uses > 96 well plate x 2, \*2 < Approximate number of uses > 5-50 / 50 µg (cannot be stored), \*3 < Approximate number of uses > 96-well plate x 1

# **Fluorescent probe** for intracellular iron detection

FerroOrange

FerroOrange is a novel probe that enables live-cell fluorescent imaging of intracellular Fe<sup>2+</sup>. By simply adding it to cultured cell, it penetrates the cell membrane and selectively interacts with intracellular Fe2+. FerroOrange is used in ferroptosis research. \* This product was commercialized under the advisory of Dr. Hideko Nagasawa and Dr. Tasuku Hirayama (Gifu Pharmaceutical University).

# Live-cell imaging of intracellular iron

The fluorescence intensity of FerroOrange was increased in HeLa cells treated with Ammonium iron (II) sulfate compared to untreated cells; conversely, its fluorescence intensity was decreased in cells treated with Bpy, the chelator of iron.



<Detection conditions> Ex: 561 nm, Em: 570-620 nm

Scale bars : 20 µm

Description	Unit	Code
<b>F O</b>	1 tube *	
FerroOrange	3 tubes *	F374

\*<Approximate number of uses per 1 tube (24  $\mu g$ ) >  $\mu$ -Slide 8 well x 20

Plate

Reader

Micro

scope

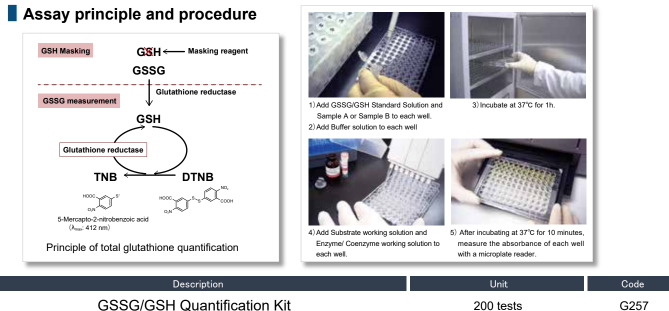
Plate

Reader

# DOJINDO MOLECULAR TECHNOLOGIES, INC. **Glutathione Quantification**

**GSSG/GSH** Quantification Kit

GSSG/GSH Quantification Kit contains Masking Reagent of GSH. GSH can be deactivated in the sample by simply adding the Masking Reagent. Therefore, GSSG is quantified by measuring the absorption ( $\lambda$  max = 412 nm) of DTNB (5,5 Edithiobis (2-nitrobenzoic acid) using the enzymatic recycling system. GSH amount can also be determined by subtracting GSSG from the total amount of glutathione.



## **GSSG/GSH** Quantification Kit

200 tests

Cytotoxicity

Mitochondria

Autophagy

# Highly Sensitive Detection of Cellular Senescence

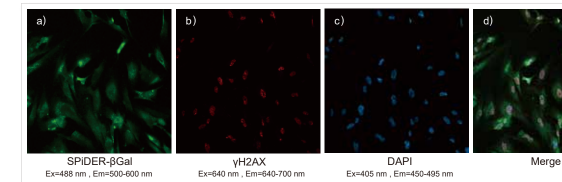


Cellular Senescence Detection Kit - SPiDER-βGal

Cellular Senescence Detection Kit – SPiDER- $\beta$ Gal is a fluorometric assay kit to detect SA- $\beta$ -gal with high sensitivity and ease of use. SPiDER- $\beta$ Gal is a reagent to detect  $\beta$ -galactosidase which possesses a high cell-permeability and a high intracellular retention enabling the use in both live and fixed cells. This kit can be applied to quantitative analysis by flow cytometry, and co-staining with SA- $\beta$ -Gal and other markers.

## **Multistaining** -Co-staining of SA-β-gal and DNA Damage marker in WI-38 cells-

WI-38 cells (passage 10) were stained with this kit, DNA damage marker  $\gamma$ H2AX as one of the senescence markers, and DAPI for indicating the location of nucleus. Details of this experiment are available on our website.



Description

Cellular Senescence Detection Kit - SPiDER-βGal

\* <Approximate number of uses > 35 mm dish x 10

III.Se k∣

Code

SG03

Plate

Reade

Unit

10 assays \*

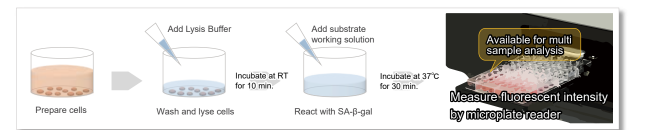
# Senescent Cell Detection (for Plate Assay)

Cellular Senescence Plate Assay Kit - SPiDER-βGal

This product is a simple detection kit that can be used by plate assay to measure senescence-associated  $\beta$ -galactosidase (SA- $\beta$ -gal) activity. By simply adding SPiDER- $\beta$ Gal, a reagent for detection of  $\beta$ -galactosidase, to a 96 well plate, you can quantify SA- $\beta$ -gal activity and evaluate multiple samples.

## Simple and easy protocol - Just add the reagents into well -

Cells prepared in advance are lysed with the Lysis Buffer included this kit. Fluorescence intensity according to the SA- $\beta$ -gal activity are obtained by simply adding the fluorescent substrate SPiDER- $\beta$ Gal to the cell lysate.



\* Precautions when using this kit

Cell counts may need to be normalized. When cells are analyzed in a microplate, the results obtained may differ depending on cell numbers per well. In such cases, normalization of the measured values obtained from cell counting and total protein will be necessary. In the Cell Count Normalization Kit (code: C554), cell numbers can be easily measured by the fluorescence intensity induced by a reagent added to the cell culture medium for staining nuclei. (Protocol for combined analysis is available on our website.)

Description	Unit	Code	
Callular Sanaganga Blata Agagy Kit SBiDEB &Cal	20 tests	SG05	
Cellular Senescence Plate Assay Kit - SPiDER-βGal	100 tests	3605	

Intracellular Metabolism

Mitochondria

Labeling Kit

# **yH2AX** Detection Kit

DNA Damage Detection Kit - yH2AX Green / Red / Deep Red

DNA Damage Detection Kit is an all-in-one kit for the detection of  $\gamma$  H2AX, an indicator of DNA Damage, using primary and secondary antibodies.

# Visualizing DNA Damage



## Co-staining with Cellular Senescence Marker

Gamma-H2AX and SA- $\beta$ -gal (senescence-associated  $\beta$ -galactosidase) of WI-38 (Passage 19) cells were detected with this kit and Cellular Senescence Detection Kit - SPiDER- $\beta$  Gal (code: SG03).

As a result, increased fluorescence derived from  $\gamma$  H2AX and enhanced SA- $\beta$ -gal activity were observed.

Comparison with WI-38 (Passage 1) cells is available on our website.

Description	Unit	Code
DNA Damage Detection Kit - γH2AX - Green	1 set *	G265
DNA Damage Detection Kit - γH2AX - Red	1 set *	G266
DNA Damage Detection Kit - γH2AX - Deep Red	1 set *	G267

 $^{\star}$  5ml of staining solution can be prepared  $\,$  / 1 set

最新情報へ G265 同仁 検索

顕微鏡

# **Nucleolus Fluorescent Staining**

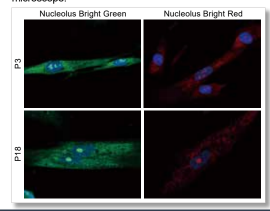
Nucleolus Bright Green/Red



Nucleolus Bright dyes are small molecules that bind to RNA in the nucleolus and emit fluorescnece. The nucleolus can be observed without any washing steps after staining with Nucleolus Bright dyes.

## **Detection in Senescent Cells**

Different passage of WI-38 cells were fixed with 4% PFA and washed with PBS, then membrane permeabilized with 1% Triton X-100. Nucleolus Bright Green (N511) or Nucleolus Bright Red (N512) and nuclear staining dye, DAPI were added and were imaged using a confocal microscope.



As a result, most passages 3 cells (P3) had multiple nucleoli in one nucleus, whereas passages 18 cells (P18) had a single enlarged nucleolus.

#### Staining Condition

Cells were immersed in 4% PFA for 5 minutes, Triton X-100 for 20 minutes. Then incubated in each fluorescent probe for 5 minutes.

#### **Detection Condition**

Nucleolus Bright Green	:	Ex=488 nm , Em=500-600 nm
Nucleolus Bright Red	:	Ex=561 nm , Em=565-650 nm
DAPI	:	Ex=405 nm , Em=450-495 nm

Description	Unit	Code
Nucleolus Bright Green	60 nmol *	N511
Nucleolus Bright Red	60 nmol *	N512

\*<Approximate number of uses per 60 nmol> 35 mm dish x 30 (Reagent concentration: 1  $\mu$  mol/l)

Oxidative Stress

Labeling

Ā

# **Lipid Droplet**

# Fluorescent reagent for Lipid droplet staining

Lipi series Blue / Green / Red / Deep Red

Lipi probes are small molecules that emit strong fluorescence in a hydrophobic environment such as Lipid droplets, which can be observed without any washing steps after staining with Lipi probes. It can be used with both live and fixed cells.

Live cell imaging	Lipi-Blue	Lipi-Green FL	Lipi-Red	Lipi-Deep Red
Oleic-acid-treated HeLa cells were stained with each	r.	м <sup>с</sup>		FL.
Lipi sereis probe.		·		
Staining condition				
HeLa cells were cultured overnight in a medium containing	20 µm	20 µm	<u>20 µm</u>	11) <u>er</u>
bleic acid. The cells were washed with PBS and stained	DIC + FL	DIC + FL	DIC + FL	
with each Lipi product working solution (Lipi-Blue / Green /	a and a		The second second	
Deep Red: 0.1 μmol/l, Lipi-Red: 1 μmol/l) for 15 minutes.	Castal	- All S	la 5	Report 1
Jpper row: fluorescence image, lower row: merged image with				
ohase contrast image. Detection conditions are shown on our website.	_20 µm	20 µm	<u>20 µm</u>	10 te
				Scale bar:20 µm
Description			Unit	Code
Lipi-Blue		1	0 nmol *	LD01
Lipi-Green		1	0 nmol *	LD02
			-	
Lipi-Green		10	0 nmol *	LD02

# Lipid Droplet AssayKit for Quantification

Lipid Droplet Assay Kit Blue / Deep Red

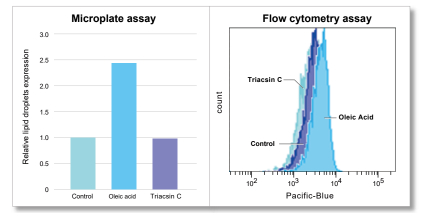


Micro

scope

Lipid Droplet Assay Kit is a fluorometric assay kit for quantification of lipid droplets. This kit considerably shortens the entire process compared to the colorimetric method and can be used for both live and fixed cells.

# Microplate / Flow cytometry assay Lipid Droplet Assay Kit - Blue



Changes in lipid droplets by the addition of oleic acid or Triacsin C (acyl-CoA synthetase inhibitor, an inhibitor of lipid droplet formation) to HeLa cells (plate assay) and A549 cells (flow cytometry) were examined using the Lipid Droplet Assay Kit-Blue. As a result, the oleic acid-treated cells showed an increase in the number of LDs, compared to control and Triacsin C-treated cells.

#### **Detection Conditions**

Microplate assay : Ex=376-386 nm, Em=435-455 nm Flow cytometry assay : Ex=405 nm, Em=425-475 nm

Description	Unit	Code
Lipid Droplet Assay Kit - Blue	1 set *	LD05
Lipid Droplet Assay Kit - Deep Red	1 set *	LD06

30

\* <Approximate number of uses> 96 well plate x 1, 40 assays for flow cytometry

Autophagy

# **Cell Membrane Dynamics**

The cell membrane plays an important role in selective transport of intracellular and extracellular substances. Since cell membrane disfunction or disruption of Endocytosis is related to cell status and linked to diseases, it is useful to capture the cell membrane dynamics in various research areas, such as neurodegenerative diseases, cancer, and senescense.

## Cell Membrane related products

#### **Endosomal Traficking**

AcidSensor Labeling Kit - Endocytic Internalization Assay (Code A558)

- pH sensor Labeling
- Co-staining with Endocytosis Detection Dye

#### Visualizing Endocytosis

#### ECGreen-Endocytosis Detection (Code E296)

- Visualizing phagocytosis
- Trackinging the virul infection pathway

# Isolating Exosomes

Exosome Isolation Kit (Code EX10/11)

Easy to use, no technique required

#### **Visualizing Exosomes**

ExoSparkler Exosome Membrane/Protein Labeling Kit (Code EX01/02/03/04/05/06)

初期エンドク

- Tracking of internalized exosome
- Flow Cytometry Analysis

Degradation pathway Recycling pathway Early endsome Recycling endosome Code Ex10 Naturation Late endsome Eusion with Lysosomes Lysosome

#### **Visualizing Cell Membrane**

PlasMem Bright Series (Code P504/505)

- Neurotoxicity
- Membrane labeling for counting cell number

#### **Visualizing Lysosomes**

LysoPrime (Code L261/264), pHLys Red (Code L265)

- Lysosomal function (pH) and localization
- Quantification of lysosomal mass

#### Lysosomal pH Detection

#### Lysosomal Acidic pH Detection Kit (Code L268)

- · High sensitivity for lysosomal pH change
- · Including positive control, convenient for first time users

31

Mitochondria

Labeling

Ā

# **Cell Membrane Staining**

PlasMem Bright Green / Red

PlasMem Bright dyes are designed to overcome the problems with conventional small molecule dyes, enabling to stain plasma membranes for over a day. Furthermore, PlasMem Bright dyes are more water-soluble and can be diluted with culture medium.

# A solution for plasma membrane staining

Small-molecule fluorescent dyes are widely used as a method for plasma membrane staining. However, short-lived stain retention in the plasma membrane and low water solubility are often problematic. Our PlasMem Bright dyes offer solutions to these problems.

#### **Features**

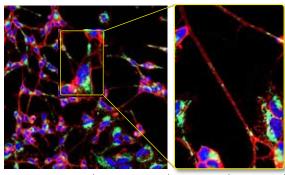
- · Applicable to live cells and fixation after staining
- · Higher retention with low toxicity
- · Simple procedure: just add reagents to culture medium

Micro

scope

## Clear visualization of plasma membrane

Prolonged Retention in the Plasma Membrane



Red : Plasma Membrane (PlasMem Bright Red), Blue : Nuclear (Hoechst 33342) Green : Mitochondria (MitoBright LT Green)

Detection Conditions PlasMem Bright Red: Ex = 561 nm , Em = 560-700 nm

## High retention on plasma membrane

**Dojindo PlasMem Bright Series** 

Just After staining

After 24 hrs

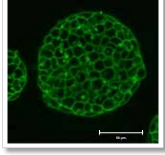
## Plasma membrane staining inside an ES cell colony

Mouse ES cells were cultured in a gelatin-coated glass bottom dish for 4 days, and the colonies were stained with PlasMem Bright

Green (200x dilution) for 15 minutes and observed under a confocal microscope (Zeiss: LSM710) after medium

exchange. As a result, the membranes of the cells inside the colonies could be visualized with PlasMem Bright Green.

**Alternative Products** 



Detection Conditions PlasMem Bright Green: Ex = 488 nm , Em = 500-560 nm \* This data was kindly provided by Dr. Otsugu Ishizu, Keio University School of Medicine.

# HeLa cells stained with each plasma membrane staining reagent were incubated for 24 hrs and their fluorescent image

Autophagy

**Cell Membrane** 

	Labelin
Scale bar∶10 µm	
Code	
D504	

Description	Unit	Code
PlasMem Bright Green	100 µl *	P504
PlasMem Bright Red	100 µl *	P505

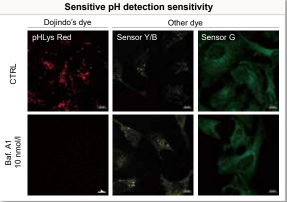
\* < Approximate number of uses per 100  $\mu$ l> 35 mm dish x 10 /  $\mu$ -Slide 8 well x 10

was compared. PlasMem Bright series had higher retention on plasma membrane than other products. \* Retention on plasma membrane may vary depending on the cell type. (Difference was observed between Hela cells and SH-SY5Y cells.) The lysosomal staining dye series includes pHLys Red, which shows lysosomal pH-dependent fluorescence change, and pH-resistant LysoPrime Green and Deep Red. Combining these dyes and simultaneously analyzing lysosomal mass and pH, enables more detailed analysis of lysosomal function.



Lysosomal pH Detection 1 pHLys Red – Lysosomal Acidic pH Detection





Merged with LAMP1 Merged with LAMP1

pHLys Red

Precise localization

Other Dye

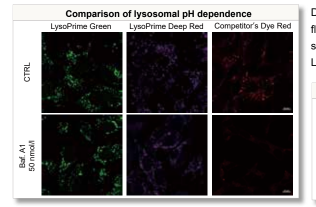
Lysosome Staining Dye ② LysoPrime Green / Deep Red – High Specificity and pH Resistance

pH in live cells. It is also applicable

long-term imaging due to its high

to experiments that require

retention ability.



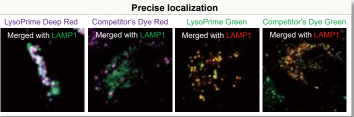
Dojindo's LysoPrime Green overcomes known problems with existing fluorescent lysosome probes, such as lack of specificity for lysosomes and staining dependent on the lysosomal pH. In addition, the high-retentivity of LysoPrime Green enables long-term imaging experiments.

Live imaging using small molecule fluorescent probes has been widely

used for lysosomal live cell analysis, but the low specificity and retention

ability due to pH change have been cited as issues. The pHLys Red is a small molecule fluorescent probe with high lysosomal specificity and

sensitivity to pH changes, enabling a more accurate analysis of lysosomal



#### Lysosomal Acidic pH Detection Combination of ① and ② Lysosomal Acidic pH Detection Kit

: LysoPrime Green

: pHLys Red

#### All-in-one Kit including Lysosome Acidification Inhibitor

- Lysosome staining Dye
- Lysosomal pH Detection Dye
- Inhibitor of lysosomal pH acidification :Bafilomycin A1

For more information	tion ờ	0.55 (0 342 (342
L266 dojindo	Search	

Description	Unit	Code
① Dye for lysosomal pH detection	1 tube *1	
pHLys Red – Lysosomal Acidic pH Detection	3 tubes *1	
② Dye for lysosome mass and localization detection		
	10 µl ×1 *1	L261
LysoPrime Green – High Specificity and pH Resistance	10 µl ×3 *1	E201
LysoPrime Deen Red – High Specificity and nH Resistance	1 tube *1	1.004
LysoPrime Deep Red – High Specificity and pH Resistance	3 tubes *1	L264
Accurate pH detection of lysosomes		
Lysosomal Acidic pH Detection Kit	1 set *2	L266

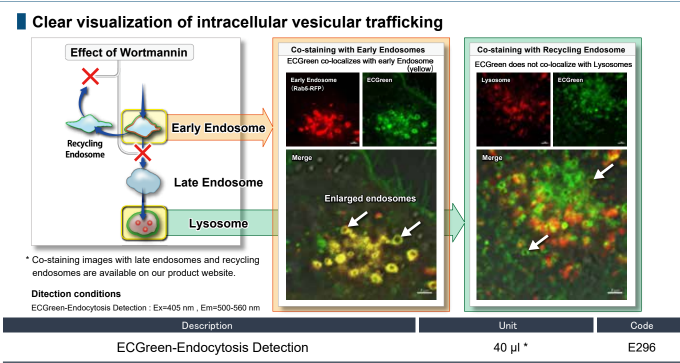
Cytotoxicit

Exosome

# **Endocytosis Detection**

**ECGreen-Endocytosis Detection** 

ECGreen-Endocytosis Detection is a pH dependent fluorescent dye that localizes to vesicle membrane. The visualization of endocytosis using the ECGreen is a more direct method than fluorescent analogs and allows visualization of endocytosis from the stage of early endosomes.



\* <Approximate number of uses per 40  $\mu$ l $>\,$  35mm dish x 20,  $\mu$ -Slide 8 well x 20

Micro

scope

Micro

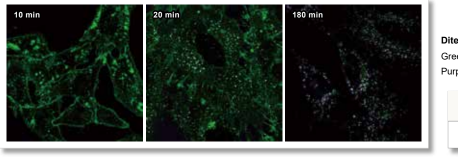
# pH Sensor Labeling Kit

AcidSensor Labeling Kit - Endocytic Internalization Assay

This kit is an all-in-one kit that allows visualization of the endocytic uptake of a target substance. The NH2-Reactive AcidSensor (fluorescent probe) included in the kit has an intramolecular active ester group that forms a stable covalent bond when mixed with an amino group-containing target substance (protein).

## Co-staining for Observation with Endosomes - Cellular uptake of labeled IgG over time

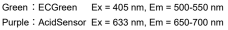
AcidSensor-labeled mouse IgG stained with this kit and Dojindo's endocytosis detection dye, ECGreen -Endocytosis Detection ((code: E296)) were added to HeLa cells, and observed at 10, 20, and 180 minutes after staining. The results showed that the AcidSensor (Deep Red) and endosomal membranes (Green) were localized at the same place, indicating that mouse IgG was uptaken by the cells via the endocytic pathway.



#### Ditection conditions

3 samples \*

E296 dojindo



A558

Description

AcidSensor Labeling Kit Endocytic Internalization Assay

\* < Approximate number of uses per 1 sample > Labels 50-200 µg of protein/antibodies. \* Protein/Antibody is not included.

Lipid droplet

Mitochondria

**Oxidative Stress** 

Autophagy

Labeling Kit

Mitochondria

Oxidative Stress

Lipid droplet

Cell Membrane

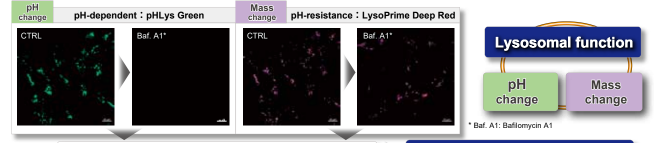
# Lysosomal Acidic pH Detection Kit

Lysosomal Acidic pH Detection Kit-Green/Deep Red

Since lysosomal dysfunction is deeply involved in the onset and progression of neurodegenerative diseases and other disorders, detailed analysis of lysosomes has attracted much attention in elucidating pathological conditions and developing therapeutic agents. Furthermore, recent studies have shown that a decrease in acidity inside lysosomes in mouse models of the neurodegenerative disease Alzheimer's disease results in the inability to degrade waste products inside autophagosomes and the accumulation of toxic amyloid- $\beta^*$ , thus increasing the need to confirm the pH of lysosomes. \* *Nature Neuroscience*, **2022**, *25*, 688–701.

## Lysosomal pH and mass - accurate analysis of lysosomal function

With existing reagents, it was difficult to examine whether lysosomal mass or their function (pH) fluctuated because the discussion was based on changes in the fluorescence brightness of a single dye. This kit contains pHLys Green, which is highly specific to lysosomes and shows pH-dependent fluorescence change, and pH-resistant LysoPrime Deep Red (code: L264). Using these two dyes, lysosomal pH and mass can be measured from the same sample enabling the detailed analysis of lysosomal function.



Lysosomal mass remains the same and pH changes

Impaired Lysosomal function

Micro

FCM

## Comparison with existing reagents

Compared to existing staining dyes, Dojindo's lysosomal detection dyes selectively accumulate on lysosomes and continue to stain for more than 24 hours. These dyes can be used in combination, and lysosomal mass and pH can be more accurately confirmed.

	Lysosomal Acidic pH Detection Kit -Green/Deep Red (L268)		Lysosomal Acidic pH Detection Kit (L266)		Company T	Company T
Dye/ Wave length	<b>pHLys Green</b> Ex=488 nm Em=490-550 nm	LysoPrime Deep Red * Ex=633 nm Em=640-700 nm	<b>pHLys Red</b> * Ex=561 nm Em=560-650 nm	LysoPrime Green * Ex=488 nm Em=500-600 nm	Lysosomes pH sensor	Lysosomes Staining dye
Purpose	pН	Mass	pН	Mass	pН	Mass
lysosome pH sensitivity	$\checkmark$	Resistant to pH change	$\checkmark$	Resistant to pH change	Less sensitivity	Less sensitivity
lysosome Specificity	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$
lysosome retention	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$
Supported Devices	Supporte	d Devices	Micro scope	Plate Reader	Micro scope	Micro scope

Indicator	Detection Color	Description	Unit	Code
pH / Mass	Green / Purple	Lysosomal Acidic pH Detection Kit -Green/Deep Red	1 set*	L268

		* <estimated 1="" number="" of="" per="" set="" use=""> 3</estimated>	5 mm dish x 10, μ-Slide 8 well	x 10, 96-well Plate x 2	
Related product					
Indicator	Detection Color	Description	Unit	Code	
pH / Mass	Red / Green	Lysosomal Acidic pH Detection Kit	1 set	L266	
pН	Pod	nHI va Rad Lyaccomal Acidia nH Dataction	1 tube	L265	
рп	Red	pHLys Red- Lysosomal Acidic pH Detection	3 tubes	L205	
	Durala	LysoPrime Deep Red	1 tube	L264	
Mass	Purple	- High Specificity and pH Resistance	3 tubes	L204	
	Green	LysoPrime Green	10 µl	L261	
	Green	- High Specificity and pH Resistance	10 µI×3	L201	

# Autophagy

# **Autophagosome Detection**

**DAPGreen - Autophagy Detection** 

**DAPRed - Autophagy Detection** 

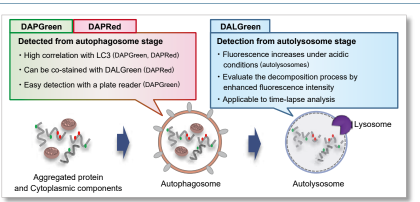
# **Autolysosome Detection**

**DALGreen - Autophagy Detection** 

Autophagy is an intracellular degradation system, where dysfunctional proteins and organelles are degraded. DAPRed, DAPGreen, and DALGreen are fluorescent reagents that can easily detect autophagy by simply adding it to cultured cells.

# Principle

DAPRed and DAPGreen, are incorporated inside of the membrane when an autophagosome membrane is formed. Its fluorescence is enhanced under lipophilic condition.On the other hand, DALGreen emits enhanced fluorescence under acidic condition after the autophagosome is fused with the lysosome. In this way, DAPRed, DAPGreen, and DALGreen enables to monitor the process of autophagosome by simply adding



FCM

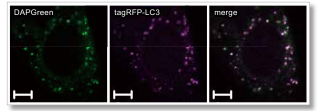
# **Related Product Information**

	Applicable instruments Fluorescent Flow Microplate Microscopy cytometer reader		Fluorescent properties	Volume / the number of usable assays	Existing methods	
DAPGreen	0	0	0	Ex = 425-475 nm Em = 500-560 nm	5 nmol x 1 / 35 mm dish: 25 (when used in 1.0 µmol/l)	LC3-GFP
DAPRed	0	×	×	Ex = 500-560 nm Em = 690-750 nm	5 nmol x 1 / 35 mm dish: 25 (when used in 1.0 µmol/l)	MDC Cyto-ID etc.
DALGreen	0	0	×	Ex = 350-450 nm Em = 500-560 nm	20 nmol x 1 / 35 mm dish: 10 (when used in 1.0 µmol/l)	

## High Correlation with LC3

DAPGreen

After adding DAPGreen to the RFP-LC3 expressed Hela cells, cells were treated with rapamycin to induce autophagy. Fluorescent imaging was conducted by confocal microscopy after 4 hrs. from autophagy induction.



**Imaging Condition** Ex=488 nm , Em=500-563 nm Scale bar : 10 μm

# Time-lapse imaging DALGreen

After staining with DALGreen, HeLa cells were cultured in amino acid-free medium, and changes in cells and fluorescence images were observed for up to 6 hours.

As a result, an increase in DALGreen fluorescence was confirmed in cells in which autophagy was induced.



Time-lapse video is now available on our website Imaging Condition Ex=405 nm, Em=500-550 nm Confocal Imaging Cytometer (Yokogawa Electric Corporation : CQ1)

Description	Unit	Code
DAPGreen - Autophagy Detection	5 nmol	D676
DAPRed - Autophagy Detection	5 nmol	D677
DALGreen - Autophagy Detection	20 nmol	D675

Lipid droplet

Mitochondria

**Oxidative Stress** 

# Substrate for Immunofluorescence

CLAMP F405-Signal Boosting

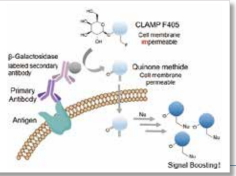


A highly sensitive CLAMP method (quinone methide-based catalyzed signal amplification) can be applied to live/fixed cells or tissue sections. In this method, using primary antibody,  $\beta$ -galactosidase-labeled secondary antibody and newly developed fluorescent substrate CLAMP F405, the cells expressing a specific low-expressed cell surface protein can be selectively detected with high sensitivity.

Note: This method cannot confirm the localization of surface antigens.

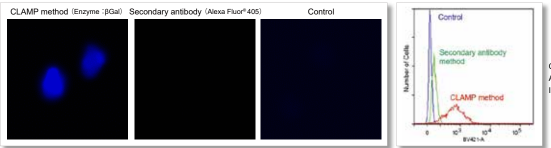
\*This product was developed with technical guidance and information provided by Prof. Yoshiki Katayama at Kyushu University.

Reference : Noguchi, K. et al., "β-Galactosidase-Catalyzed Fluorescent Reporter Labeling of Living Cells for Sensitive Detection of Cell Surface Antigens", *Bioconjugate Chem.*, **2020**, 31(7), 1740–1744.



# Highly sensitive detection of cellular surface antigen expression

A fluorescence detection method using a fluorescence-labeled antibody is widely known as a specific detection method for cell surface proteins (fluorescence immunostaining method). However, it is difficult to apply this method for low expressed surface proteins due to low sensitivity. CLAMP F405-Signal Boosting allows you to detect these proteins.



Cell Line : HeLa Cells Antigen : CD44 Instrument : (Left) Fluorescent microscope Ex=340–380 nm, Em=435–485 nm (Right) Flow Cytometer Ex=405 nm, Em=425–575 nm

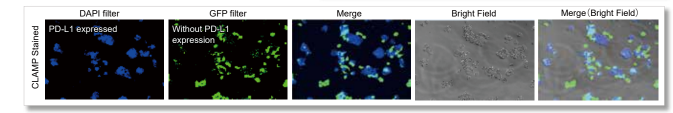
# Application Data

#### highly sensitive, high selectivity, and high retention

PD-L1 expressed HepG2 cells and CFSE-stained control cells were prepared and mixed. The results showed the CLAMP stained cells did not localize in the CFSE stained cells, indicating that the CLAMP method accurately differentiates PD-L1 expressed HepG2 cells, which was difficult using the secondary antibody method.

\* CFSE: 5- or 6- (N-Succinimidyloxycarbonyl) fluorescein 3',6'-diacetate

HopG2 Cells The CLAMP method HepG2 Cells 



#### FFPE Tissue Sections - Human Small Intestine

Using the CLAMP method and TSA (Tyramide signal amplification) method detected the  $\alpha$ SMA ( $\alpha$ -smooth muscle actin) and keratin on FFPE tissue sections of human small intestine samples. The results showed that the CLAMP method can be used for co-staining with other staining method. Please refer to the reference for a detailed protocol on the CLAMP method for tissue sections.

Reference: Hirata, M., et al. "Galactosidase-catalyzed fluorescence amplification method (GAFAM): sensitive fluorescent immunohistochemistry using novel fluorogenic  $\beta$ -galactosidase substrates and its application in multiplex immunostaining." *Histochem Cell Biol* 159, 233–246 (2023)



\* The data was kindly provided by Dr. Masahiro Hirata, Department of Diagnostic Pathology, Kyoto University Hospital.

Description	Unit	Code
CLAMP F405-Signal Boosting	10 µl *	C554

Cytotoxicity

Mitochondria

Oxidative

-ipid droplet

Cell Membrane

 $^{\ast}$  Primary antibody and  $\beta\text{-}Galactosidase$  labeled secondary antibody is not included in this kit.

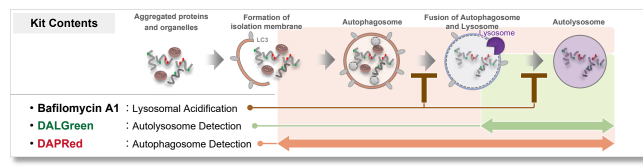
# Autophagic Flux Assay

Autophagic Flux Assay Kit

Autophagic Flux Assay Kit contains autophagosome and autolysosome detection dye (DAPRed), autolysosome detection dye (DALGreen), and lysosomal acidification inhibitor (Bafilomycin A1). This kit allows the accurate evaluation of autophagic flux by monitoring autophagosome formation through autolysosome formation <sup>1)</sup>. 1) H. Sakurai, *et al., iScience*, **2023**, *26*, 107218.

## What this kit can do

This all-in-one kit includes DAPRed (code: D677) for the detection of autophagosomes and autolysosomes, DALGreen (code: D675) for the detection of autolysosomes, and Bafilomycin A1 for inhibiting lysosome acidification. By simply adding the reagents, you can monitor the process from autophagosome formation to autolysosome formation  $2^{0,3}$ .



#### Analyzing the autophagic flux by fluorescence changes

Autophagic flux can be analyzed more in detail by using this kit in conjunction with autophagy induction or inhibition. It enables to evaluate the stages in the autophagic flux by observing the fluorescence change of DAPRed and DALGreen simultaneously.

Fluorescence chang	e relative to control	Indication from observed fluorescence changes	Typical inhibitors <sup>4), 5)</sup>	
DAPRed	DALGreen	indication non observed nuclescence changes	Typical inflibitors 0,0	
1	1	Autophagy induction or autolysosome accumulation	E64d/Pepstatin A	
+	+	Inhibition in autophagosome formation step	3-MA	
🕇 or 📫	₽	Inhibition of autolysosome formation	Bafilomycin A1	

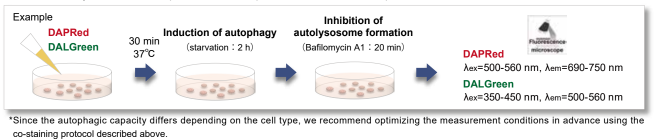
# Simple Operation

 2) X. Chen, et al., Am J Transl Res., 2020, 12(9):4902-4922.
 4) N

 3) C. Oh, et al., J Neurosci., 2022, 42(14), 3011-3024.
 5) D

4) N. Mizushima, et al., Cell, 2010, 140(3), 313-326.
5) D. J. Klionsky, et al., Autophagy, 2021, 17(1), 1-382.

Add DAPRed and DALGreen working solution to cells at the same time, then induce/inhibit the autophagy. Then treat with Bafilomycin A1 and it is ready to observe. No operations such as plasmid transfection are required.



\*This kit includes Bafilomycin A1 for evaluation of the condition.

	Description	Unit	Code
Autopl	hagic Flux Assay Kit	1 set *	A562
Related product	* 〈Approximate number	of uses per 1 set $ angle$ 35 mm dish x 5, µ-Slide 8 v	vell x 5, 96-well plate x 1
	Description	Unit	Code
Autophagosome detection dye	DAPGreen - Autophagy Detection	5 nmol *1	D676
	DAPRed - Autophagy Detection	5 nmol <sup>∗1</sup>	D677
Autolysosome detection dye	DALGreen - Autophagy Detection	20 nmol *2	D675
Lysosomal pH detection kit	Lysosomal Acidic pH Detection Kit -Green/Deep Red	1 set *3	L268
	Lysosomal Acidic pH Detection Kit	1 set *3	L266

(Approximate number of uses) \*1:35 mm dish x 25, \*2:35 mm dish x 10, \*3: per 1 set - 35 mm dish x 10, μ-Slide 8 well x 10, 96-well plate x 2

Proliferation Cytotoxicity

Intracellular Metabolism

Labeling Kit

# Exosome

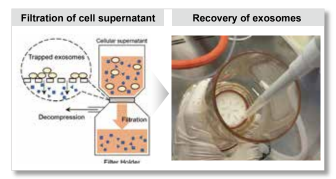
# **Exosome Isolation**

Exolsolator Exosome Isolation Kit / Exolsolator Isolation Filter



Exo*lsolator* Exosome Isolation Kit can collect exosomes from cell supernatants with a recovery rate equivalent to the ultracentrifugation (UC) method. Unlike the UC, Exo*lsolator* Exosome Isolation Kit requires only the filtration procedure, thus exosomes are obtained quickly without any complicated operations.

# Simple and easy operation



Exo*lsolator* Exosome Isolation Kit includes Filter Holder and Isolation Filter that can collect exosomes from cell supernatant by adding PBS to the filter surface after filtration. Its recovery rate is high and easy to use, no complicated technique is required throughout the process.

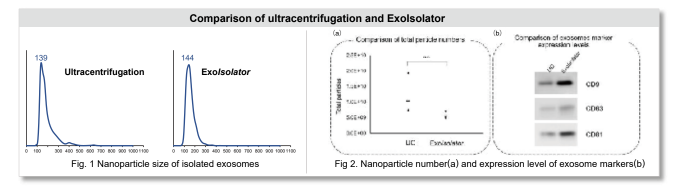
To purchase additional filters

\* Estimated sample processing volume: For culture supernatant, 25 ml is recommended for one filter.



## Recovery rate equivalent to ultracentrifugation

Ultracentrifugation is the most commonly used method to isolate exosomes. We isolated the exosomes from the supernatant of HEK293S using both of ultracentrifugation method and the Exo*lsolator* method. The particle size distribution (Fig. 1), the number of particles (Fig. 2(a)) and the expression level of exosome markers (Fig. 2(b)) of the isolated exosomes were tested and compared. The results showed that the Exo*lsolator* recovered exosomes with equivalent particle size distribution and the number of particles to the ultracentrifugation method, and the amount of exosome marker expression per protein was higher, indicating that Exo*lsolator* recovered exosomes with higher purity than the ultracentrifugation method.



# Kit contents

#### For first-time users



Description	Unit	Code
Exolsolator Exosome Isolation Kit	3 tests	EX10
Exo <i>lsolator</i> Isolation Filter	10 pieces	EX11

Mitochondria

Membrane

Ā

# **Exosome Membrane / Protein Fluorescent Staining**

ExoSparkler Exosome Membrane / Protein Labeling Kit Green / Red / Deep Red

ExoSparkler series can be used for staining the membrane or proteins of the purified exosomes and enables the imaging of labeled exosomes taken up by cells.

## ExoSparkler series does not cause extracellular aggregation

Co-staining with Mem Dye-Deep Red (Purple) Co-staining with Mem Dye-Deep Red (Purple) and Alternative Product "P" (Green)

and Alternative Product "P" (Red) Extracellular fluorescent spots Extracellular fluorescent spots

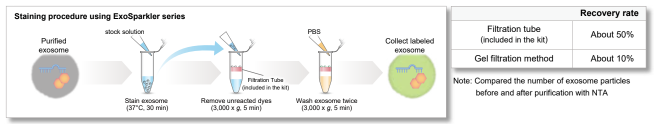
Exosomes stained with ExoSparkler's Mem Dye-Deep Red or an alternative product (green or red) were added to each well containing HeLa cells. The labeled exosomes taken into HeLa cells were observed by fluorescent microscopy. As a result, extracellular fluorescent spots suspected of dye aggregations were observed in each well containing the exosomes stained with the alternative product (green or red).

#### **Detection conditions**

Mem Dye-Deep Red (Purple): Ex 640nm/Em 640-760nm Alternative Product "P" (Green): Ex 561nm/Em 560-620nm Alternative Product "P" (Red): Ex 640nm/Em 650-700nm

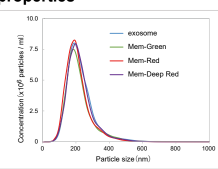
## All-in-one kit for preparing the labeled exosomes

ExoSparkler series contains filtration tubes for the removal of unreacted dyes after fluorescence labeling, as well as an optimized protocol for labeling exosomes. Our ExoSparkler series makes it possible to prepare fluorescence labeling of exosomes using the simple procedure.



## ExoSparkler Mem Dyes have little effect on exosome properties

NTA (nanoparticle tracking analysis) and zeta potential were measured to determine the changes in exosomes before and after staininge with Mem Dyes. As a result, the Mem-Dye series (green, red, deep red) had little effect on exosome properties. Comparison data is available on our website.



NTA comparison of dye-stained (Mem-Dye) or unstained exosomes

Description	Unit	Code
Exosome membrane		
Exo <i>Sparkler</i> Exosome Membrane Labeling Kit-Green	5 samples *	EX01
Exo <i>Sparkler</i> Exosome Membrane Labeling Kit-Red	5 samples *	EX02
Exo <i>Sparkler</i> Exosome Membrane Labeling Kit-Deep Red	5 samples *	EX03
Exosome protein		
Exo <i>Sparkler</i> Exosome Protein Labeling Kit-Green	5 samples *	EX04
ExoSparkler Exosome Protein Labeling Kit-Red	5 samples *	EX05
ExoSparkler Exosome Protein Labeling Kit-Deep Red	5 samples *	EX06

Micro scon

Mitochondria

Autophagy

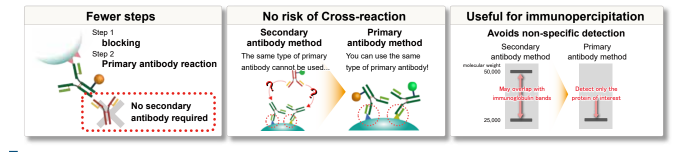
\* Protein amount : 1-10 μg / sample, Particle count : 10 to 100 x 10<sup>8</sup> / sample (As purified exosome using ultracentrifugation)

# Dojindo Labeling Kit series



Dojindo Labeling Kit series is a kit that allows you to directly label your primary antibodies with fluorescent dyes, biotin, or enzymes.

#### Features



#### Product Lineup

References using each product are available on our website.

Biotin	Sample amount / type	Target	Description	Unit	Code	Detectior
	10 µg antibody	-NH2	Ab-10 Rapid Biotin Labeling Kit *	3 samples	LK37	
	50-200 µg	-NH2	Biotin Labeling Kit -NH <sub>2</sub> *	3 samples	LK03	Micro scope
	antibody • protein	-SH	Biotin Labeling Kit -SH	3 samples	LK10	
Biotin	1 mg antibody • protein	-NH2	Biotin Labeling Kit - $\rm NH_{2}$ (for 1mg) *	1 sample	LK55	FCM Plate Reader
	1-5 mg antibody ∙ protein	-NH2	Biotinylation Kit (Sulfo-OSu)*	1 set	BK01	Western Blot
fluorescent dye	Sample amount / type	Target	Description	Unit	Code	Detectior
	10 µg antibody	-NH2	Ab-10 Rapid Fluorescein Labeling Kit *	3 samples	LK32	
Fluorescein	50-200 µg antibody ∙ protein	-NH <sub>2</sub>	Fluorescein Labeling Kit -NH $_2$ *	3 samples	LK01	-
	10 µg antibody	-NH <sub>2</sub>	Ab-10 Rapid HiLyte Fluor™ 555 Labeling Kit -NH₂ *	3 samples	LK35	
	50-200 µg antibody ∙ protein	-NH2	HiLyte Fluor™ 555 Labeling Kit -NH $_2$ *	3 samples	LK14	Micro
HiLyte Fluor	10 µg antibody	-NH <sub>2</sub>	Ab-10 Rapid HiLyte Fluor™ 647 Labeling Kit-NH₂ *	3 samples	LK36	
-	50-200 µg antibody • protein	-NH <sub>2</sub>	HiLyte Fluor™ 647 Labeling Kit-NH₂ *	3 samples	LK15	FCM
ICG	50-200 μg antibody ∙ protein	-NH2	ICG Labeling Kit-NH $_2$ *	3 samples	LK31	_
fluorescent protein	Sample amount / type	Target	Description	Unit	Code	Detection
	10 µg antibody	-NH2	Ab-10 Rapid R-Phycoerythrin Labeling Kit *	3 samples	LK34	
R-Phycoerythrin	50-200 µg	-NH <sub>2</sub>	R-Phycoerythrin Labeling Kit-NH <sub>2</sub> *	3 samples	LK23	Micro
	antibody • protein	-SH	R-Phycoerythrin Labeling Kit-SH	3 samples	LK26	scope
A.U	50-200 µg	-NH <sub>2</sub>	Allophycocyanin Labeling Kit-NH <sub>2</sub> *	3 samples	LK21	FCM
Allopycocyanine	antibody • protein	-SH	Allophycocyanin Labeling Kit-SH	3 samples	LK24	
Enzyme	Sample amount / type	Target	Description	Unit	Code	Detectio
	10 µg antibody	-NH <sub>2</sub>	Ab-10 Rapid Peroxidase Labeling Kit *	3 samples	LK33	
	50-200 µg	-NH2	Peroxidase Labeling Kit-NH <sub>2</sub> *	3 samples	LK11	
Peroxidase	antibody • protein	-SH	Peroxidase Labeling Kit-SH	3 samples	LK09	Micro scope
	1 mg antibody • protein	-NH2	Peroxidase Labeling Kit - $\rm NH_2~(for~1mg)~\star$	1 sample	LK51	Plate Reader
Alkaline	50-200 µg	-NH <sub>2</sub>	Alkaline Phosphatase Labeling Kit-NH <sub>2</sub> *	3 samples	LK12	Western
Phosphatase	antibody • protein	-SH	Alkaline Phosphatase Labeling Kit-SH	3 samples	LK13	Blot

Proliferation Cytotoxicity

Mitochondria

\* Labeling reaction may interfere the antigen recognition site of the antibody. If you have any questions, please contact our Customer Support or your sales representative.

# **Applicable Instruments**

Search for products by your device



# Plate Reader

r			
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## Western Blot

Antibody / Protein Labeling

```
Micro
scope
     Microscope
```

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