

# **Biochemical Reagents**



# **Contents**

Nucleic Acid Isolation/ Electrophoresis	
Total RNA Isolation Reagent: Sepasol-RNA I Super G Tail Lysis Buffer Nuclease Decontamination: RNase Quiet Coprecipitant of DNA/RNA: Gene-Packman Coprecipitant Agarose for Nucleic Acid Electrophoresis Bullet PAGE Plus Precast Gel DNA Ladder Markers Dual Green Nucleic Acid Stain. Ethidium Bromide Solution (0.44 mg/ml) Phenol solution for DNA/RNA extraction	69111213
Cell Culture	
Cell Culture Medium Custom Cell Culture Media Reduced Serum DMEM/Ham's F-12 Medium AscleStem series AscleStem® hPSC Maintenance Supplement AscleStem® Dissociation Solution for hPSCs AscleStem® Hepatocyte Differentiation & Maintenance Basal Medium AscleStem® Neuronal Medium & Supplement(50x) AscleStem® 0.1%-Gelatin Solution 0.01%-Polyethyleneimine Coating Solution Mitomycin C Solution (1 mg/ml) for preparation of feeder cells Cell Dissociation Reagent: Accutase™ EZSPHERE® for 3D cell culture with microwells Serum-free Cell Freezing Media: Cell Reservoir One ES/iPS cell Freezing Media: Cell Reservoir One, Vitrify Balanced Salines Sterile water Media additives. Reagents for cell dissociation Antibiotics Cell Count Reagent SF, based on WST-8 MTT Cell Count Kit, based on reduction of MTT. LDH Cytotoxicity Assay Kit Trypan Blue Solution Reagents for apoptosis research Reagents for alkaline phosphatase staining Medium for Bacteria, Plusgrow II. IPTG and X-gal Solutions Gelling Agent for Plant Study: Gellan Gum.	
Cell Extraction / Protein Assay	
Zymolyase <sup>™</sup> (from Arthrobacter Luteus)  Cell Lysis Solution: Cell Lysis Buffer  Protease Inhibitor Cocktail  Phosphatase Inhibitor Cocktail  Determination of Protein Concentration: Protein Assay  Protein Assay CBB Clean Up Kit  Reductant adaptable reagent for protein assay BCA Kit	44 46 47 48

# Contents

Protein Purification	
COSMOGEL® Ig-Accept Protein G COSMOGEL® GST-Accept High Performance Magnetic Nanoparticles: FG beads® TurboTEV Protease & Turbo3C Protease	52 
Protein Electrophoresis	
Bullet PAGE Plus Precast Gel  Extra PAGE One Precast Gel  Electrophoresis Tank for Precast Gel  WIDE RANGE Gel Preparation Buffer (4x) for PAGE  Stacking Gel Buffer Solution (4x) with Blue Color  Rapid Running Buffer Solution  Polyacrylamide Gel Casting Reagents  Running Buffers  Sample Buffer Solution for SDS-PAGE (6x)  Molecular Weight Markers  PAGE Clean Up Kit  CBB Stain Solution  Silver Staining Kit.	59 67 62 63 64 64 65 65 66 66
Western Blotting	
Bullet Semi-dry Transfer One Blocking One and Blocking One P Bullet Blocking One for Western Blotting Bullet ImmunoReaction Buffer Signal Enhancer HIKARI for Western blotting and ELIS HRP-conjugated secondary antibody Chemiluminescent Western Blotting Substrates Colorimetric Western Blotting Substrates WB Stripping Solution Epitope Tag Antibody	75 76 77 A 86 87 87 87 87 87 88
Immunohistochemistry	
4% - Paraformaldehyde Phosphate Buffer Solution HistoVT One (10x, pH 7.0)	9 <sup>.</sup> 99
Streptavidin Biotin Complex Peroxidase Kit	94

# Total RNA Isolation Reagent: Sepasol®-RNA I Super G

- » Easy to identify interphase
- » Less than 1hr for RNA isolation
- » Enables to extract DNA and proteins along with RNA from a single sample

# The second secon

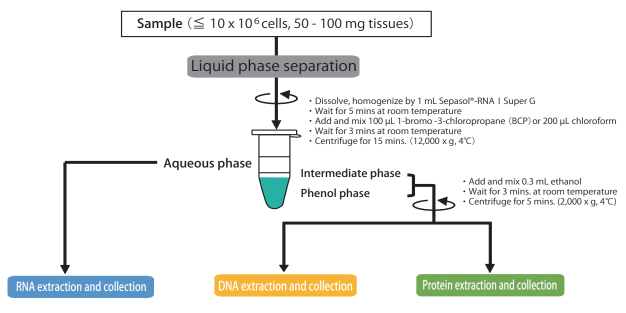
### **Phenol Phase Color**

Sepasol-RNA I Super G



Easy to identify interphase

### **Procedure**



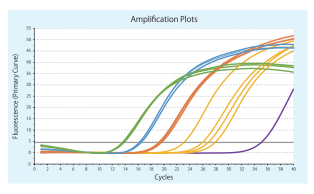
For details, refer to the instruction manual.

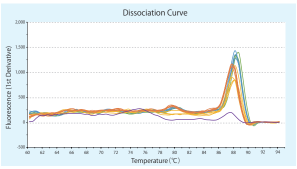
### **Performance evaluation**

Total RNA was extracted from HL-60 cells at 8 x  $10^6$  cells, 1 x  $10^6$  cells, and 0.125 x  $10^6$  cells using this product. Additionally, for the 0.125 x  $10^6$  cells of HL-60 cells, samples were prepared with Gene-Packman Coprecipitant (#12680-30) as well as general alcohol precipitation. The extracted total RNA was treated with DNase according to the instruction manual attached and proceed to RT-qPCR. Please note that throughout the entire process, all cell number samples were treated under identical conditions, including the volume of the dissolution solution.

### [ Result (RT-qPCR) ]

Using this product, it was possible to extract mRNA in a cell number-dependent manner. Moreover, although RNA yield decreased and greater variability was observed (yellow lines) when the cell number was low, such as 0.125 x 10<sup>6</sup> cells, Gene-Packman co-precipitant (#12680-30) increased the yield and reduced variability (orange lines).





Sample	Ct value
— 8 x 10 <sup>6</sup> cells	13.55 - 13.8
— 1 x 10 <sup>6</sup> cells	16.12 - 16.35
- 0.125 x 10 <sup>6</sup> cells	23.04 - 27.87
— 0.125 x 10 <sup>6</sup> cells (co-precipitation)	19.17 - 19.59
Non-Template Control	34.99

\*Threshold: 4.47

### (Conditions)

Primer:

[Gene target] β-Actin

[Forward primer] 5'-AAGAGAGGCATCCTCACCCT-3' [Reverse primer] 5'-TACATGGCTGGGGTGTTGAA-3'

[PCR product] 218 bp

PCR enzyme (reverse transcriptase enzyme): One Step TB Green® PrimeScript RT-PCR Kit II (Perfect Real Time) (Takara Bio #RR086A)

(Thermal Cycler Dice, TB Green and PrimeScript are registered by Takara Bio.)

### PCR condition:

1.Hold (reverse transcription) 42°C (5 mins.)→ 95°C (10 sec.), 1 cycle

2. 2 Step PCR 95°C (5 sec.) $\rightarrow$  60°C (30 sec.), 40 cycles

3.Dissociation 95°C (15 sec.)  $\rightarrow$  60°C (30 sec.)  $\rightarrow$  95°C (5 sec.), 1 cycle

Real-time PCR equipment

Thermal Cycler Dice® Real Time System III

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
		09379-26	10 ml
Canada (San Al Comana Coffee historical association facility than 1)	Б	09379-84	100 ml
Sepasol®-RNA I Super G (for biological samples [cell, tissue, etc.])	R	09379-97	200 ml
		09379-55	500 ml
Sepasol®-RNA II Super (for liquid samples [blood, etc.])	R	30487-46	100 ml

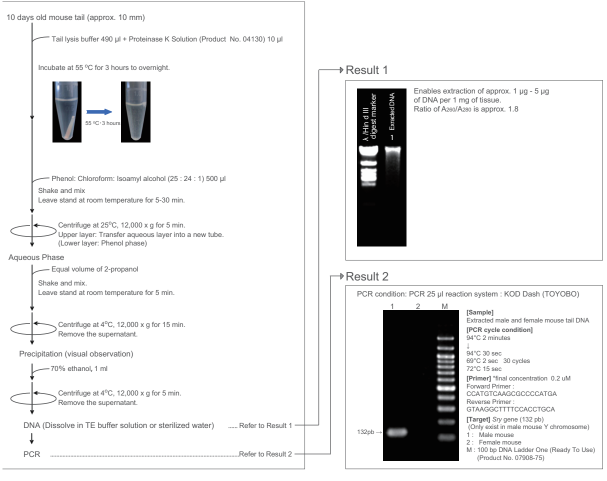
# **Tail Lysis Buffer**

Tail Lysis Buffer is ready-to-use solution that enables simple genotyping procedure.

- » Ready-to-use solution
- » DNase, RNase free

### **Application: Genotyping of mouse tail**

### **Procedure**



*Sry* gene (132 bp), which only exists in male mouse Y-chromosomes was increased in male derived DNA, but was not increased in female derived DNA. The result shows that PCR operates correctly.

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Tail Lysis Buffer	RT	06169-95	500 ml

### **Related prduct**

Product Name	Storage	Product No.	PKG Size
Dratainana II frans Tritirachium album	R	29442-14	100 mg
Proteinase K from Tritirachium album	K	29442-85	500 mg
Dratainage W/Dagambinant\ Calutian	R	15679-06	2 ml
Proteinase K(Recombinant) Solution	K	15679-64	10 ml

### **Nuclease Decontamination: RNase Quiet**

» Free from contamination : removes nucleases and nucleic acids

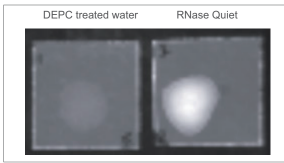
» Easy to wipe : no detergent contained

» Easy to use : spray type

» Non-carcinogenic : no DEPC contained



### **Application 1: Decontamination of cover glass**



### Conditio

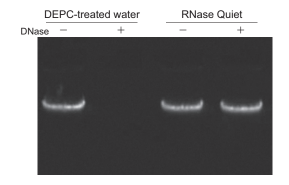
- 1. Apply 100  $\mu$ l of RNase A solution (1 mg/ml) to cover glasses and dry them.
- Spray with DEPC-treated water or RNase Quiet and wait for 1 minute. Wipe thoroughly with a clean paper towel and then rinse with RNase-free sterile water
- 3. Apply 50  $\mu$ l of RNA solution (40  $\mu$ g/ml) on the cover glasses and incubate them at 37°C for 30 minutes.
- 4. Apply 1  $\mu$ I ethidium bromide solution (20  $\mu$ g/mI) to the cover glasses.
- 5. Observe with UV detector.

### Application 2: RNase and DNase decontamination of 1.5 ml micro-tubes

[RNase decontamination test]

DEPC-treated water
RNase - + - +

[DNase decontamination test]



### Condition

① Add the following solutions and dry them up.

RNase decontamination test

- (+): Add 10  $\mu$ l of RNase A solution (10 mg/ml) to a 1.5 ml micro-tube.
- (-): Add 10 µl of only the solution used to dissolve RNase A to a 1.5 ml micro-tube as a control.

DNase decontamination test

- (+): Add 10  $\mu l$  of DNase I (27.3 U/10  $\mu l)$  to a 1.5 ml micro-tube, and dry it up.
- (–): Add 10  $\mu l$  of only the solution used to dissolve DNase I to a 1.5 ml micro-tube as a control.
- ② Add 1 ml of DEPC-treated water or RNase Quiet and wait for 1 minute.
- ③ Remove the solution from the tubes and rinse those with 1 ml of DEPC-treated water.
- (4) Remove the DEPC-treated water from the tubes and add 25 µl of RNA solution (40 µg/ml) for RNase decontamination test; 25 µl of DNA solution (40 µg/ml) and 1 µl of 50 mmol/L magnesium chloride for DNase decontamination test, and incubate them at 37°C for 30 minutes.
- $\ensuremath{\mathfrak{D}}$  Reacted samples are electrophoresed on agarose gel and detected using EtBr.

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
RNase Quiet (with spray nozzle)	RT	09147-14	475 ml
RNase Quiet Refill	RT	09477-94	475 ml

# Coprecipitant of DNA/RNA: Gene-Packman Coprecipitant

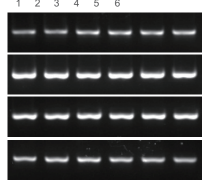
- » Almost complete nucleic acid recovery
- » No requirement of low-temperature incubation
- » Endotoxin, DNase and RNase tested
- » High nucleic acid pellet visibility

### No Effect of Gene-Packman Coprecipitant to PCR

In order to figure out that Gene-Packman Coprecipitant does not effect PCR performance, Gene-Packman Coprecipitant was consciously added to PCR reaction solutions which amplify 1,000 bp DNA fragments. This data shows that Gene-Packman does not affect PCR performance.

Amount of Gene-Packman Coprecipitant to PCR reactin solution:

Lane 1: 0 μl, Lane 2: 0.2 μl, Lane 3: 0.5 μl, Lane 4: 1.0 μl, Lane 5: 3.0 μl and Lane 6: 5.0 μl.



Blend Taq by TOYOBO

KOD Dash by TOYOBO

KOD FX by TOYOBO

Takara EX Taq by TAKARA BIO

PCR condition

Reaction Volume : 25 µl reaction

Tempate DNA : 15 ng

Denaturation : 94°C for 0.5 min.

Annealing : 55°C for 0.5 min.

Extension : 72°C for 1 min.

Cycle : 25 times

The agarose gel image above shows no interference with Gene-Packman Coprecipitant to PCR performance.

(Blend Taq®, KOD Dash® and KOD FX® are registered by Toyobo. TaKaRa Ex Taq® are registered by Takara Bio.)

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Gene-Packman Coprecipitant	R	12680-30	1 KIT

# **Agarose for Nucleic Acid Electrophoresis**

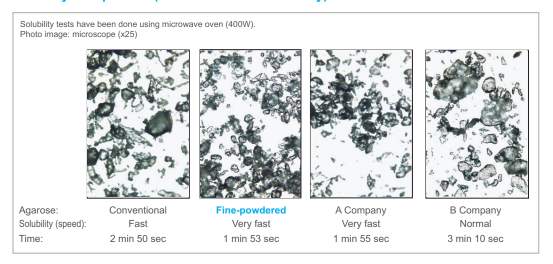
### Fine-powdered Agarose

» High solubility : Smaller average particle size for easy disolution

» Simple : Easy-to-weigh

» High sharpness : Sharp bands observable

## Solubility Comparison (Particle size and solubility)



### **Specification**

Type :  $\geq$  1kbp Sulfate (%) :  $\leq$  0.2

Gel Strength :  $\geq$  2,500 g/cm $^2$  (at 1.5%)

 $\begin{array}{ll} \mbox{Gel Point (°C)} & : 36 \pm 1.5 \\ \mbox{Electroendosmosis (-mr)} & : 0.09 \text{-} 0.13 \\ \end{array}$ 

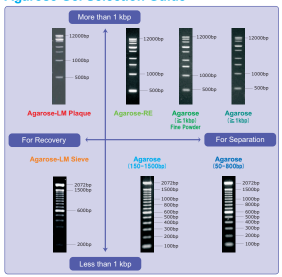
### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
		02468-24	10 g
Agarose for ≥ 1kbp fragment (Fine Powder)	RT	02468-66	100 g
		02468-95	500 g

[Storage] RT = Room Temperature, R = Refrigerator, F = Freezer

## Other Agaroses

### **Agarose Gel Selection Guide**





# Immunohistochemistry

### **Ordering Information**

	Product Name	Storage	Product No.	PKG Size
\garose-LM Plaque for ≧	≧ 1kbp fragment			
Туре	: Low Melting Agarose ≧ 1kbp fragment			
Sulfate (%)	: ≦ 0.5		01650-02	25 g
Gel Strength	: ≥ 250 g/cm² (at 1.5%)	RT	01650-86	100 g
Gel Point (°C)	: ≦ 30		01030-80	100 g
Melting Temp. (°C) Electroendosmosis (-mr)	: ≤ 65.5 : ≤ 0.12			
DNase, RNase tested				
Agarose-LM Sieve for ≦	1kbp fragment			
Type	: Low Melting Agarose ≦ 1kbp fragment			
Sulfate (%)	: ≦ 0.5		0.1051.00	0.5
Gel Strength	: ≥ 1,000 g/cm² (at 4%)	RT	01651-92	25 g
Gel Point (°C)	: ≦ 35		01651-76	100 g
Melting Temp. (°C)	: ≦ 65			
Electroendosmosis (-mr) DNase, RNase tested	: ≦ 0.12			
,	agment, for Restriction and Ligation			
Type Sulfate (%)	: ≧ 1kbp : ≦ 0.2		01149-92	25 g
Gel Strength	0.2 : $\ge 2,500 \text{ g/cm}^2 \text{ (at 1.5\%)}$	RT	01149-76	100 g
Gel Point (°C)	: 36 ± 1.5		01149-05	500 g
Electroendosmosis (-mr)	: 0.09 - 0.13			
Tested for Silver staining, Dna	ase, RNase and Enzyme reaction			
Agarose for $\geqq$ 1kbp fragr	ment (Fine Powder) refer to pxx			
Туре	: ≧ 1kbp		02468-24	10 g
Sulfate (%)	: ≦ 0.5	RT	02468-66	100 g
Gel Strength	: ≧ 2,500 g/cm² (at 1.5%)		02468-95	500 g
Gel Point (°C)	: 36 ± 1.5			3
Electroendosmosis (-mr)	: 0.09 - 0.13			
Agarose for $\ge$ 1kbp fragr	ment		04400 00	0.5
Туре	: ≧ 1kbp		01163-92	25 g
Sulfate (%)	$1 \le 0.2$	RT	01163-76	100 g
Gel Strength Gel Point (°C)	: ≧ 2,500 g/cm² (at 1.5%) : 36 ± 1.5		01163-05	500 g
Electroendosmosis (-mr)	: 0.09 - 0.13			
Agarose for 150-1,500bp	fragment			
Туре	: 150-1,500bp			
Sulfate (%)	: ≦ 0.1		01153-22	25 g
Gel Strength	$\ge 2,000 \text{ g/cm}^2 \text{ (at 1.5\%)}$	RT	01153-64	100 g
Gel Point (°C)	: ≦ 36.5		01100-04	100 g
Electroendosmosis (-mr)	: ≦ 0.12			
DNase, RNase tested				
Agarose for 50-800bp fra	gment			
Type	: 50-800bp		01447 40	0.5
Sulfate (%)	0.1 $0.1$ $0.2$ $0.1$	RT	01147-12	25 g
Gel Strength Gel Point (°C)	: ≧ 750 g/cm² (at 1.5%) : 30		01147-96	100 g
Electroendosmosis (-mr)	: 30 : ≤ 0.12			
DNase, RNase tested	·			
Agarose-LE, Classic type				
Type	: LE, Classic		01157-82	25 g
Sulfate (%)	: ≦ 0.2	RT	01157-66	100 g
Gel Strength	: ≥ 2,500 g/cm² (at 1.5%)	141	01157-95	500 g
Gel Point (°C)	: 36 ± 1.5		01137-83	500 g
Electroendosmosis (-mr)	: 0.09 - 0.13			
Agarose-ME, Classic type	9			
Туре	: ME, Classic		01158-72	25 g
Sulfate (%)	: ≦ 0.25	RT	01158-56	100 g
Gel Strength	: ≧ 2,000 g/cm² (at 1.5%)		01158-85	500 g
Gel Point (°C) Electroendosmosis (-mr)	: 36 ± 1.5 : 0.16 - 0.19			3
Agarose-LM (melting tem				
Type	: Low Melting Agarose			
Sulfate (%)	: ≦ 0.2	_	01161-12	25 g
Gel Strength	$0.2$ : $\leq 5.50 \text{ g/cm}^2 \text{ (at 1.5\%)}$	RT	01161-54	100 g
Gel Point (°C)	: 26 ± 2		01101-04	100 g
Molting Town (°C)	: ≦ 65			
Melting Temp. (°C) Electroendosmosis (-mr)	: ≦ 0.12			

# **Bullet PAGE Plus Precast Gel**

Bullet PAGE Plus is a precast gel for high-speed electrophoresis.

For protein electrophoresis, refer to p54.

### » Only 12 minutes electrophoresis time for nucleic acid with 400 V

### » Surfactant-free, so can be used for nucleic acid analysis.

\*Please use 25 mM Tris and 192 mM glycine buffer for nucleic acid electrophoresis.

	Gel	Gradient gel				
Gel cor	ncentration	5-10 %	5-15 %	5-20 %	7.5-15 %	10-20 %
Product	13 wells	21789-34	21791-84	21793-64	21795-44	21797-24
number	17 wells	21790-94	21792-74	21794-54	21796-34	21798-14
- C	nucleic acids AGE)	1,500 → 500 → 100 → 20 → 20 →	1,500→ 500→ 100→	1,500→ 500→ 100→	1,500- 500- 100- 20-	1,500- 500- 100- 20-

G	Gel	Homogeneous gel			
Gel cond	centration	7.5%	10%	12.5%	15%
Product	13 wells	21799-04	21801-44	21807-84	21853-74
number	17 wells	21800-54	21806-94	21811-14	21854-64
- C	nucleic acids (GE)	1,500 - 100	1,500-> 500-> 100->	1,500- 500- 100- 100-	(bp)

### **Ordering Information**

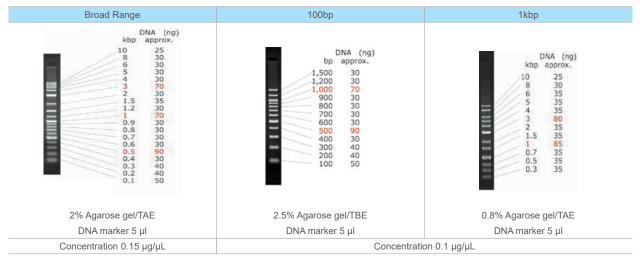
ordering information			
Product Name	Storage	Product No.	PKG Size
Bullet PAGE Plus Precast Gel, 5-10%, 13wells	R	21789-34	10 Sheets
Bullet PAGE Plus Precast Gel, 5-10%, 17wells	R	21790-94	10 Sheets
Bullet PAGE Plus Precast Gel, 5-15%, 13wells	R	21791-84	10 Sheets
Bullet PAGE Plus Precast Gel, 5-15%, 17wells	R	21792-74	10 Sheets
Bullet PAGE Plus Precast Gel, 5-20%, 13wells	R	21793-64	10 Sheets
Bullet PAGE Plus Precast Gel, 5-20%, 17wells	R	21794-54	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5-15%, 13wells	R	21795-44	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5-15%, 17wells	R	21796-34	10 Sheets
Bullet PAGE Plus Precast Gel, 10-20%, 13wells	R	21797-24	10 Sheets
Bullet PAGE Plus Precast Gel, 10-20%, 17wells	R	21798-14	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5%, 13wells	R	21799-04	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5%, 17wells	R	21800-54	10 Sheets
Bullet PAGE Plus Precast Gel, 10%, 13wells	R	21801-44	10 Sheets
Bullet PAGE Plus Precast Gel, 10%, 17wells	R	21806-94	10 Sheets
Bullet PAGE Plus Precast Gel, 12.5%, 13wells	R	21807-84	10 Sheets
Bullet PAGE Plus Precast Gel, 12.5%, 17wells	R	21811-14	10 Sheets
Bullet PAGE Plus Precast Gel, 15%, 13wells	R	21853-74	10 Sheets
Bullet PAGE Plus Precast Gel, 15%, 17wells	R	21854-64	10 Sheets

### **DNA Ladder Markers**

- » Covers wide range from 0.1 kbp to 10 kbp
- » Emphasis of 0.5, 1 and 3 kbp bands
- » Ready-to-use markers containing 2 loading dyes



### **Product Contents**



### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
DNA Ladder One (Broad Range)(Ready-to-use)	R	08362-85	500 µl
100bp DNA Ladder One (Ready-to-use)	R	07908-75	500 µl
1kbp DNA Ladder One (Ready-to-use)	R	08232-85	500 µl

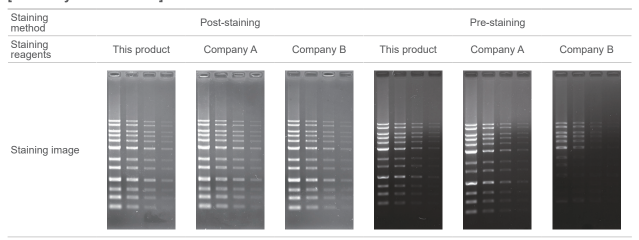
### **Dual Green Nucleic Acid Stain**

- » Can be detected by UV excitation
- » Can be applied in both pre-stain and post-stain procedures
- » Can be used with agarose and polyacrylamide gels
- » All double-stranded DNA, single-stranded DNA and RNA can be detected.

### Comparison

In agarose gel, DNA detection was performed by using post-staining and pre-staining methods. This product has equivalent or superior performance compared to products from other companies.

### [Detect by UV excitation]



### [Detect by Bule LED]

Staining method		Post-staining			Pre-staining	
Staining reagents	This product	Company A	Company B	This product	Company A	Company B
Staining image						

(Conditions)

Sample : 1kbp DNA Ladder One(Ready To Use) (#08232-85) Total nucleic acid quantity: 600 ng, 200 ng, 67 ng, 20 ng from left

Gel : 1% Agarose gel

 ${\tt Electrophoresis\ buffer\ : Prepare\ Tris-Acetate-EDTA\ Buffer\ (\#35430)\ at\ 1x\ concentration\ (dye-free)}$ 

Electrophoresis : Run at 100 V until BPB reaches about 3/4 of the gel.

 $Staining : [Pre-staining] \ After \ dissolving \ the \ agarose, \ each \ dye \ was \ added \ to \ make \ a \ 20,000x \ dilution, \ and \ then \ the \ gel \ was \ solidified.$ 

[Post-staining] TAE (1x) to make a 10,000x dilution, then place the gel after electrophoresis and shake for 30 minutes (without destaining).

 $\label{thm:decomposition} \mbox{Detection: ChemiDoc Touch MP} (\mbox{Bio-Rad laboratories}) \mbox{ The exposure time is detested automatically}.$ 

[UV excitation] SYBR® Green mode (UV transmission, 590 / 110 nm detect on filter)

[Blue LED excitation] Alexa Fluor® 488 mode (Blue incident light, 532 / 28 nm detect on filter)

 ${\rm SYBR}^{\rm e}, {\rm Alexa\ Fluor}^{\rm e}\ {\rm is\ registered\ trademark\ of\ Molecular,\ Probes\ and\ Incorporated}.$ 

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Dual Green Nucleic Acid Stain	RT	20599-41	1 ml

# Ethidium Bromide Solution (0.44 mg/ml)

Ethidium Bromide Solution (0.44 mg/ml) is easy and safe to use because of its eye-drop bottle. It is used in adjustment of nucleic acid staining after electrophoresis or gel containing ethidium bromide.

### How to use

Adjust the concentration of ethidium bromide solution as follows

Concentration of Ethidium Bromide	Adjusting Solution	Ethidium Bromide Solution (0.44 mg/ml)
0.1 μg/ml	200 ml	1 drop
0.2 μg/ml	100 ml	1 drop
0.5 μg/ml	40 ml	1 drop



### Note:

- 1. 1 drop of Ethidium Bromide Solution (0.44 mg/ml) is 45  $\mu$ l.
- 2. In situations where Ethidium Bromide Solution (0.44 mg/ml) is used in concentrations other than shown in the above table, remove the nozzle, collect the appropriate amount with a micropipette and dilute accordingly.
- 3. For adjustments of even greater ethidium bromide solution volumes, use Ethidium Bromide Solution (10 mg/ml) (Product No. 14631-94).

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Ethidium Bromide Solution (0.44 mg/ml) eye-drop-bottle	R	02393-94	10 ml
Ethidium Bromide Solution (10 mg/ml)	R	14631-94	10 ml

# Phenol solution for DNA/RNA extraction

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size	
For DNA				
Phenol, Saturated with TE Buffer	Buffer-saturated phenol at pH 6.6. Easily adjusted to pH 7.9 adding the attached buffer before use.	R	25969-54 25969-96	100 ml 400 ml
Phenol, Saturated with TE Buffer	It contains 8-quinolinol as a stabilizer. The phenol layer (lower layer) is yellow in color and can be easily distinguished from the clear water layer (upper layer). pH 8 already adjusted.	R	26829-54 26829-96	100 ml 400 ml
Phenol:Chloroform:Isoamyl Alcohol 25:24:1 Mixed, pH6.7	Proteins and lipids can be removed. Easily adjusted to pH 8 adding the attached buffer before use.	R	25967-74 25967-16	100 ml 400 ml
Phenol:Chloroform:Isoamyl Alcohol 25:24:1 Mixed, pH7.9	It contains 8-quinolinol as a stabilizer. The phenol layer (lower layer) is yellow in color and can be easily distinguished from the clear water layer (upper layer). pH 8 already adjusted.	R	25970-14 25970-56	100 ml 400 ml
For RNA				
Phenol, Saturated with Citrate Buffer	Buffer-saturated phenol at pH4.3.	R	25968-64	100 ml
Phenol:Chloroform:Isoamyl Alcohol 25:24:1 Mixed, pH5.2	Proteins and lipids can be removed. Good for recovery of RNA with long poly(A).	R	26058-54 26058-96	100 ml 400 ml

### **Related products**

Product Name	Storage	Product No.	PKG Size
Reagent for lysation			
Proteinase K from Tritirachium album	R	29442-14	100 mg
Floremase K nom mulachum album	IX.	29442-85	500 mg
Proteinase K(Recombinant) Solution	R	15679-06	2 m
Totelliase N(Necombiliant) Solution		15679-64	10 m
8mol/l-Guanidine Hydrochloride Solution	RT	17356-24	100 m
6mol/I-Guanidine Thiocyanate Solution	RT	16689-04	100 m
100g/l-Hexadecyltrimetylammonium Bromide Solution (10% (w/v) CTAB Solution)	RT	17472-94	100 m
Zymolyase <sup>™</sup> 20T	R	07663-91	1 (
Zymolyase <sup>™</sup> 100T	R	07665-55	500 m
Buffer for electrophoresis			
Tris-Acetate-EDTA Buffer (10x) [TAE Buffer]		35430-61	1
TIIS-ACETATE-ED TA BUITET (TOX) [TAE BUITET]	RT	35430-74	5
Tris-Acetate-EDTA Buffer (50x) [TAE Buffer]	RT	32666-81	1
Tris-Borate-EDTA Buffer (5x) [TBE Buffer]	RT	35432-41	1
Tris-Borate-EDTA Buffer (10x) [TBE Buffer]	RT	35440-31	1
This-boliate-LDTA build (Tox) [TBL build]	IXI	35440-44	5
Running Buffer Solution(10x) for PAGE	RT	30340-91	1
MOPS Buffer Stock Solution (10x) (pH 7.0)	RT	22089-45	500 n
Reagent for polyacrilamid gel			
40(w/v)%-Acrylamide/Bis Mixed Solution(19:1), Nuclease tested	R	06140-45	500 m
30(w/v)%-Acrylamide/Bis Mixed Solution(19:1), Nuclease tested	R	07175-75	500 n
10(w/v)%-Ammonium Peroxodisulfate Solution	F	02634-34	10 n
N,N,N',N'-Tetramethylethylenediamine	RT	33401-72	25
Denaturing reagent			
Formaldehyde Solution	RT	16223-55	500
Urea	RT	35940-65	500
Formamide	F	02020-64	100 n

# **Cell Culture Medium**

- » Animal origin-free
- » Sterility tested for bacteria, fungus and mycoplasma
- » Endotoxin tested
- » pH tested

### **Product list**

Media name	Product No.	Glucose	L-Gln	Phenol Red	HEPES	Sodium Pyruvate	Other	GIBCO	Sigma
	08459		0	0	-	-		11965-092	D5796
	08458		0	0	-	0		11995-065	D6429
	11584		-	0	-	0		10313-021	D6546
	08457		0	0	0	-		12430-054	
	11585	High	-	0	0	-			D6171
	08488	ingii	-	0	-	-		11960-044	D5671
DMEM	08489		-	-	-	-		31053-028	D1145
	16971		0	0	-	0	with 1,500 mg/L sodium hydrogen carbonate		
	16972		-	0	-	-	without calcium	21068-028	
	08456	Low	0	0	-	0		11885-084	D6046
	08490	LOW	-	-	-	0		11054-020	
	09891	No	0	0	-	-		11966-025	
	11581	0	0	0	-	0		11320-033	D8062
	08460	0	0	0	0	0		11330-032	D8437
DMEM/	11583	0	-	0	0	0			D6421
Ham's F-12	05177	0	0	-	0	0		11039-021	
	11582	0	0	-	-	0			
	09893	No	0	0	-	0			
Ham's F-12	17458	0	0	0	-	0		11765-054	N6658
G-MEM	12965	0	0	0	-	-		11710-035	
	21442	0	0	0	-	-		11095-080	M4655
MEM	21443	0	0	0	-	-	with non-essential amino acids		
	09848	No	0	0	-	-	with non-essential amino acids		
α-ΜΕΜ	21445	0	0	0	-	0		12561-056	
α-IVI⊏IVI	21444	0	0	0	-	0	with nucleosides	12571-063	
IMDM	11506	0	0	0	0	0		12440-053	16529
	30264	0	0	0	-	-		11875-093	R8758
	30263	0	0	0	0	-		22400-089	
RPMI 1640	05176	0	-	0	-	-		21870-076	R0883
A IVII IOHU	06261	0	0	-	-	-		11835-030	R1780
	16970	0	0	0	0	0	with 4,500 mg/L glucose	A1049101	
	09892	No	0	0	-	-		11879-020	

Compositions of each product are available on online catalog, "e-Nacalai Search Version" at www.nacalai.com

For media additives, refer to p32.

### **Ordering Information**

Product Name	Grade	Storage	Product No.	PKG size
DMEM(4.5g/I Glucose)	CD (F TC)	Б	08459-35	500 ml
with L-Gln, without Sodium Pyruvate, liquid	SP (For TC)	R	08459-64	10 x 500 ml
DMEM(4.5g/I Glucose)	CD (For TC)	В	08458-45	500 ml
with L-Gln and Sodium Pyruvate, liquid	SP (For TC)	R	08458-16	10 x 500 ml
DMEM(4.5g/l Glucose) with Sodium Pyruvate, without L-Gln, liquid	SP (For TC)	R	11584-85	500 ml
DMEM(4.5g/l Glucose) with L-Gln and HEPES, without Sodium Pyruvate, liquid	SP (For TC)	R	08457-55	500 ml
DMEM(4.5g/I Glucose)	SP (For TC)	R	11585-75	500 ml
with HEPES, without L-Gln and Sodium Pyruvate, liquid				
DMEM(4.5g/l Glucose) without L-Gln and Sodium Pyruvate, liquid	SP (For TC)	R	08488-55	500 ml
DMEM(4.5g/l Glucose) without L-Gln, Sodium Pyruvate and Phenol Red, liquid	SP (For TC)	R	08489-45	500 ml
DMEM(4.5g/l Glucose) with L-Gln, Sodium Pyruvate and 1.5g/l Sodium Hydrogen Carbonate, liquid	SP (For TC)	R	16971-55	500 ml
DMEM(4.5g/l Glucose) Calcium free without L-Gln and Sodium Pyruvate, liquid	SP (For TC)	R	16972-45	500 ml
	/ )	_	08456-65	500 ml
DMEM(1.0g/l Glucose) with L-Gln and Sodium Pyruvate, liquid	SP (For TC)	R	08456-36	10 x 500 ml
DMEM(1.0g/l Glucose) with Sodium Pyruvate, without L-Gln and Phenol Red, liquid	SP (For TC)	R	08490-05	500 ml
DMEM(No Glucose) with L-Gln, without Sodium Pyruvate, liquid	SP (For TC)	R	09891-25	500 ml
DMEM/Ham's F-12	, ,			
with L-Gln and Sodium Pyruvate, without HEPES, liquid	SP (For TC)	R	11581-15	500 ml
DMEM/Ham's F-12 with L-Gln, Sodium Pyruvate and HEPES, liquid	SP (For TC)	R	08460-95	500 ml
DMEM/Ham's F-12 with Sodium Pyruvate and HEPES, without L-Gln, liquid	SP (For TC)	R	11583-95	500 ml
DMEM/Ham's F-12	SP (For TC)	R	05177-15	500 ml
with L-Gln, Sodium Pyruvate and HEPES, without Phenol Red, liquid	3F (F0FFC)		03177-13	300 1111
DMEM/Ham's F-12 with L-Gln and Sodium Pyruvate, without HEPES and Phenol Red, liquid	SP (For TC)	R	11582-05	500 ml
DMEM/Ham's F-12(No Glucose) with L-Gln and Sodium Pyruvate, liquid	SP (For TC)	R	09893-05	500 ml
Ham's F-12 with L-Gln, liquid	SP (For TC)	R	17458-65	500 ml
G-MEM with L-Gln, liquid	SP (For TC)	R	12965-65	500 ml
MEM with Earle's Salts and L-Gln, liquid	SP (For TC)	R	21442-25	500 ml
MEM with Earle's Salts, L-Gln and Non-Essential Amino Acids, liquid	SP (For TC)	R	21443-15	500 ml
MEM(No Glucose) with Earle's Salts, L-Gln and Non-Essential Amino Acids, liquid	SP (For TC)	R	09848-05	500 ml
α-MEM with L-Gln, without Ribonucleosides and Deoxyribonucleosides, liquid	SP (For TC)	R	21445-95	500 ml
α-MEM with L-Gln, Ribonucleosides and Deoxyribonucleosides, liquid	SP (For TC)	R	21444-05	500 ml
IMDM with L-Gln and HEPES, liquid	SP (For TC)	R	11506-05	500 ml
<u> </u>	, ,		30264-85	500 ml
RPMI 1640 with L-GIn, liquid	SP (For TC)	R	30264-56	10 x 500 ml
RPMI 1640 with L-Gln and HEPES, liquid	SP (For TC)	R	30263-95	500 ml
RPMI 1640 without L-Gln, liquid	SP (For TC)	R	05176-25	500 ml
RPMI 1640 with L-GIn, without Phenol Red, liquid	SP (For TC)	R	06261-65	500 ml
	. ,			
RPMI 1640(4.5g/l Glucose) with L-Gln, Sodium Pyruvate and HEPES	SP (For TC)	R	16970-65	500 ml

### **Custom Cell Culture Media**

### For researchers who want

- · Specific compositions that appeared in the literature
- To modify the composition of commercially available cell culture media
- To get rid of phenol red due to its estrogenic effect

### **Specification**

Product Form	Liquid
Minimum PKG Size	500 ml
Guaranteed items	pH, Osmotic pressure, Sterilized, Endotoxin tested, Mycoplasma tested
Lead terms	8-10 weeks

### **How to Order**

Please visit our website at http://www.nacalai.co.jp/global/reagent/custom/Custom\_Services.html and fill out the request form.

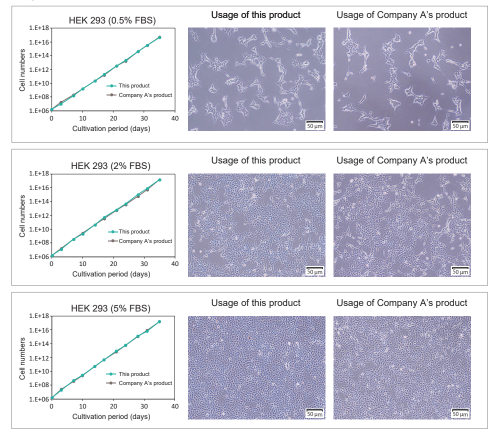
### Reduced Serum DMEM/Ham's F-12 Medium

- » Cultivable under low-serum conditions
- » Suitable as a basal medium for serum-free cultivation
- » Reduce costs associated with serum usage
- » Minimize variations between batches, reducing serum-related effects

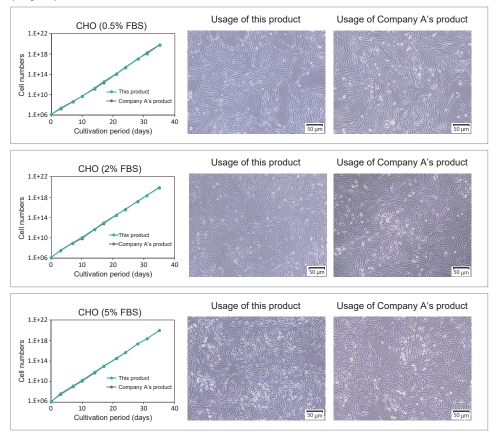
### Comparison of cell proliferation rate and cell morphology

HEK 293 and CHO cells were cultured using both this product and Company A's serum-reduced medium. Similar to the medium from Company A, our product demonstrated that cultivation under low-serum conditions is feasible without affecting either cell proliferation rate or morphology

# Proliferation curves and cell morphology of HEK 293 cells cultured under various serum concentrations (Day 35)



# Proliferation curves and cell morphology of CHO cells cultured under various serum concentrations (Day 35)



### **Application**

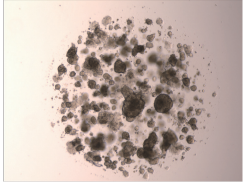
### **Cultivation of Colorectal Cancer Patient-Derived Organoids (Under Serum-Free Conditions)**

We prepared droplets (n=4) embedding dispersed colon cancer patient-derived organoid cells using Corning<sup>®</sup> Matrigel Basement Membrane Matrix Growth Factor Reduced and TrypLE Express. Then, 1 ml of medium \*was added on top, and the cultures were incubated.

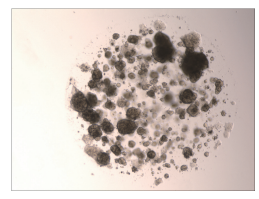
Visual assessment of proliferation did not reveal significant differences between the two conditions.

\*2 mM L-Alanyl-L-Glutamine was added to both of this product and Company A's product.

### Usage of this product



### Usage of Company A's product



Data courtesy of Associate Professor Junpei Kondo, Department of Molecular Biochemistry, Division of Health Sciences, Graduate School of Medicine, Osaka University

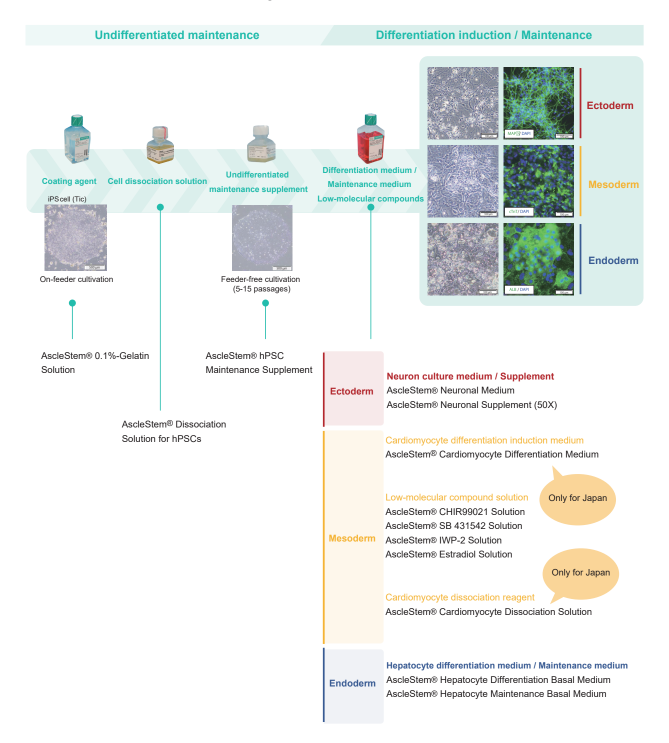
### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Reduced Serum DMEM/Ham's F-12 Medium	R	21906-55	500 ml

### **AscleStem series**

### Line-up and application of AscleStem

The AscleStem® series offers a wide range of products, from undifferentiated maintenance to differentiation induction-related reagents. Using AscleStem® series products, we performed undifferentiated maintenance and differentiation induction to the trilineage in Tic line of human iPS cells.

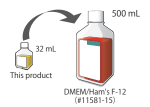


# AscleStem® hPSC Maintenance Supplement

AscleStem® hPSC Maintenance Supplement (#21979-44), is a feeder-free culture supplement for maintaining undifferentiated human pluripotent stem cells (hPSCs). It can be easily prepared by simply adding it to DMEM/Ham's F-12 medium. This product is based on the composition of a clinical grade undifferentiated maintenance medium developed through collaborative research with Osaka University. Please refer to the provided references for more information.

- » Simply add it into DMEM/Ham's F-12 (#11581-15)
- » Available to reduce the cost
- » Applicable for the weekend skip protocol
- » Medium components are disclosed \* Concentration is confidential

### **Protocol**



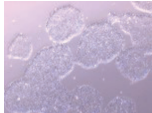
After thawed this product (stored in a frozen state) in a refrigerator, add the entire contents to 500 ml of DMEM/Ham's F-12. Please use it after thorough stirring.

### **Application**

### Cell morphology and alkaline phosphatase activity

The cell morphology of the 253G1 hiPSCs was confirmed after 2 weeks of culture with this product (prototype). Alkaline phosphatase staining was also performed to confirm that undifferentiated ability was not affected.









### Reference

Hua Y, Yoshimochi K, Li J, Takekita K, Shimotsuma M, Li L, et al. Development and evaluation of a novel xeno-free culture medium for human-induced pluripotent stem cells. Stem Cell Res Ther. 2022;13:223. doi:10.1186/s13287-022-02879-z https://pubmed.ncbi.nlm.nih.gov/35658933/

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
AscleStem® hPSC Maintenance Supplement	F	21979-44	32 ml

# AscleStem® Dissociation Solution for hPSCs

AscleStem® Dissociation Solution for hPSCs (#21777-84) is a cell dissociation solution for human pluripotent stem cells (hPSCs) composed of collagenase, trypsin, and serum substitutes. It allows for gentle dissociation of colonies during passaging of hPSCs without making single cells.

- » Allows for gentle passaging with pipetting only
- » Mycoplasma tested

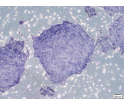


<Cell detachment during passaging>
Observed passaging of human iPS cells in on-feeder culture with this product.

### **Application**

We confirmed the undifferentiated potential of human iPS cells, which were passaged ten times using this product, by performing alkaline phosphatase staining.





201B7 on SNL76/7 feeder cells

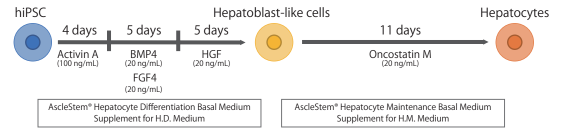
### **Ordering Information**

1 Toduct Name	luraye	Product No.	FNG SIZE
AscleStem® Dissociation Solution for hPSCs	F	21777-84	30 ml

# AscleStem<sup>®</sup> Hepatocyte Differentiation & Maintenance Basal Medium

- » Enables human pluripotent stem cells to differentiate into hepatocytes
- » Maintenance Basal Medium is suitable for drug screening applications

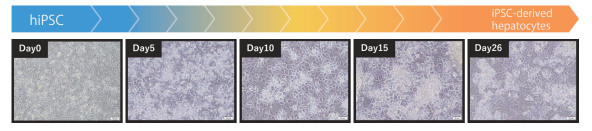
### Protocol for hiPSC differentiation to hepatocytes



### **Application**

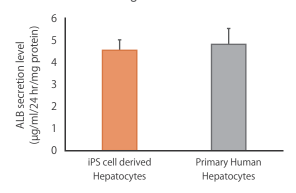
Using AscleStem® Hepatocyte Differentiation/Maintenance medium, the Tic hiPSCs was induced to differentiate into hepatocytes, and albumin production was compared with that of primary cultured human hepatocytes.

### Imaging: Phase microscopy



### Comparison of albumin production levels

This figure shows similar albumin production levels in hepatocytes differentiated from Tic hiPSCs using AscleStem<sup>®</sup> Hepatocyte Differentiation/Maintenance medium and primary human hepatocytes cultivated 24 hours after thawing.



### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
AscleStem® Hepatocyte Differentiation Basal Medium	R	20607-05	500 ml
Supplement for H.D. Medium	F	20612-54	5 ml
AscleStem® Hepatocyte Maintenance Basal Medium	R	20683-05	500 ml
Supplement for H.M. Medium	F	20615-24	30 ml

# AscleStem® Neuronal Medium & Supplement(50x)

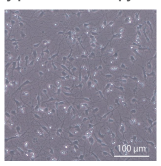
- » Serum-free media
- » Suitable for iPSC-derived neuronal cell culture

### Differentiation from neural progenitor cells to neurons

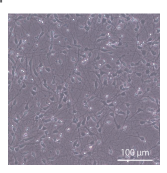
hiPSCs-derived neural progenitor cells (day 17 after differentiation induction) were seeded at 500,000 cells/cm² and induced to differentiate into neurons by adding GlutaMAX and other inducers to our medium/ our supplement or Company A's medium / their supplement. Immunostaining was performed to confirm the expression of MAP2, a marker of mature neurons.

We confirmed that hiPSC-derived neurons cultured in AscleStem® Neuronal Medium and AscleStem® Neuronal Supplement(50x) were comparable in hiPSC-derived neurons cultured in Thermo's system.

### Observation by phase microscopy 26 days after differentiation

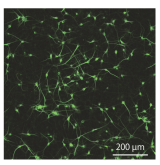


Company A's medium / their supplement

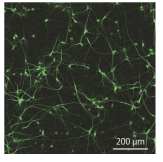


AscleStem® Neuronal Medium / AscleStem® Neuronal Supplement

### Imaging by immunohistochemistry 56 days after differentiation



Company A's medium / their supplement



AscleStem® Neuronal Medium / AscleStem® Neuronal Supplement

Data courtesy of Assistant Prof. Kaneyasu Nishimura from Division of Integrated Pharmaceutical Sciences, Kyoto Pharmaceutical University

### Differentiating motor neurons from hiPSCs

Using our supplement or Company A's supplement, human iPS cells were induced to differentiate into motor neurons. Immunostaining was performed to confirm the induction of motor neurons by choline acetyltransferase (ChAT) and the elongation of nerve axons by  $\beta$  III tubulin.

### **Protocol**



# Observed cell morphology and cell number and survival rate at detachment (28 days after start of differentiation induction)

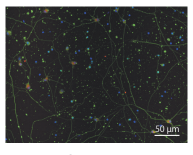


Company A
Survival rate: 41.3%
Cell number: 4.63 x 10<sup>4</sup> cells/cm<sup>2</sup>

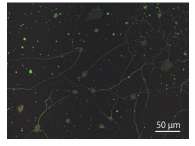


AscleStem® Neuronal Supplement Survival rate: 50.2% Cell number: 4.87 x 10<sup>4</sup> cells/cm<sup>2</sup>

### IHC (β III tubulin / ChAT / DAPI : 31 days after differentiation)



Company A



AscleStem® Neuronal Supplement

Morphology, degree of axon elongation, and ChAT expression levels were comparable using AscleStem® Neuronal Supplement compared to Company A's.

Data courtecy of Stem Cell & Device Laboratory, Inc. (SCAD)

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
AscleStem® Neuronal Medium	R	21168-35	500 ml
AscleStem® Neuronal Supplement (50x)	F	21169-54	10 ml

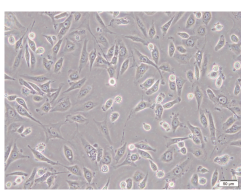
[Storage] RT = Room Temperature, R = Refrigerator, F = Freezer

# AscleStem® 0.1%-Gelatin Solution

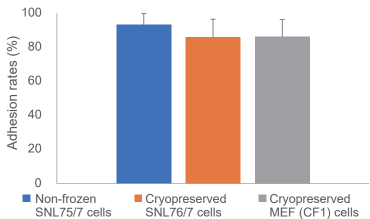
- » Ready-to-use
- » Good for cultivation of feeder cells

### Application1: Adhesion of feeder cells

After coating a cell culture plate with AscleStem® 0.1%-Gelatin Solution, non-freezed SNL76/7 cells, cryopreserved SNL76/7 cells or MEF (CF1) cells were cultivated, then adhesion rates were measured.







### Application2: Maintenance culture of hiPSCs

SNL76/7 cells and MEF (CF1) cells were seeded onto plates coated with this product, on which the 253G1 hiPSCs was maintained cuture and confirmed colony morphology and undifferentiated characteristics (alkaline phosphatase staining).

### [Morphological observation]

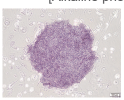


253G1 on SNL76/7 feeder cells

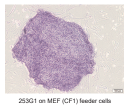


253G1 on MEF (CF1) feeder cells

### [Alkaline phosphatase staining]



253G1 on SNL76/7 feeder cells



### **Ordering Information**

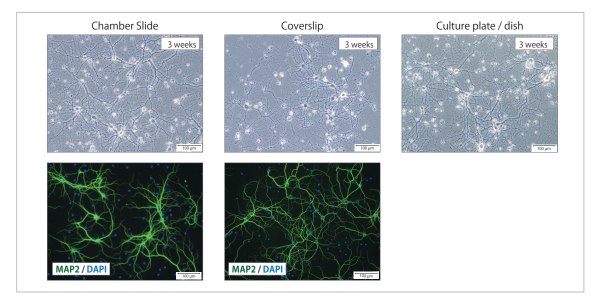
Product Name	Storage	Product No.	PKG Size
AscleStem® 0.1%-Gelatin Solution	RT	19895-75	500 ml

# 0.01%-Polyethyleneimine Coating Solution

- » Applicable for primary neuronal cell culture
- » Suitable for various types of cell culture labware
- » Store at room temperature

### **Application**

Primary cultured neurons from the mouse fetal brain hippocampus were successfully maintained and cultured on various cell culture labware coated with this product.



### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
0.01%-Polyethyleneimine Coating Solution	RT	21958-14	100 ml

# Mitomycin C Solution (1 mg/ml) for preparation of feeder cells

- » Ready-to-use: Sterility-tested for cell culture, does not solidify in freezer
- » High stability: 2 years in freezer, protected from light



### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Mitomycin C Solution (1mg/ml)	F	20898-21	1 ml

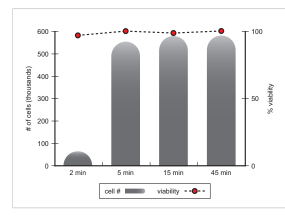
### **Related Products**

	Product Name	Storage	Product No.	PKG Size
Y-27632		_	08945-71	1 mg
1-27032		Г	08945-84	5 mg

# **Cell Dissociation Reagent: Accutase™**

- » Gentle and efficient dissociation of any adherent cell line
- » No mammalian or bacterial components are contained
- » No neutralization steps by serum or trypsin inhibitors are required
- » Works extremely well on embryonic and neuronal stem cells

### **Application**



Human MG63 Fibrosarcoma cells cultured in DMEM + 10% FBS were treated with Accutase™.

Treatment resulted in rapid cell detachment, a single cell suspension, and high viability.

Accutase<sup>TM</sup> is gentle on cells; viability was  $97 \pm 3\%$  even after 45 minutes in Accutase<sup>TM</sup>.

### **Cell Lines dissociated with Accutase**<sup>™</sup>

- hESCs
- · vascular endothelial cells
- hepatocyte progenitors
- · adherent CHO cells
- 293 cells
- 3T3
- HeLa
- M24 metastatic melanoma
- gliomas D54

- fibroblasts
- · hepatocytes
- · primary chick embryo neuronal cells
- · adherent BHK cells
- · L929 cells
- Verd
- NT2
- A375 metastatic melanoma
- HT1080 fibrosarcoma cells

- keratinocytes
- · vascular smooth muscle cells
- · bone marrow stem cells
- macrophages
- immortalized mouse testicular germ cells
- · COS
- MG63
- gliomas U251
- Sf9 insect cells

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Accutase™	F	12679-54	100 ml

### **Related Products**

Product Name	Storage	Product No.	PKG Size
Accumax	F	13766-74	100 ml
2.5g/l-Trypsin Solution	F	35555-54	100 ml
5.0g/I-Trypsin/5.3mmol/I-EDTA Solution	F	35556-44	100 ml
2.5g/I-Trypsin/1mmol/I-EDTA Solution	F	35554-64	100 ml
2.5g/I-Trypsin/1mmol/I-EDTA Solution, with Phenol Red	F	32777-44	100 ml
0.5g/I-Trypsin/0.53mmol/I-EDTA Solution	F	35553-74	100 ml
0.5g/I-Trypsin/0.53mmol/I-EDTA Solution, with Phenol Red	F	32778-34	100 ml
0.2g/l-EDTA Solution	R	14367-74	100 ml

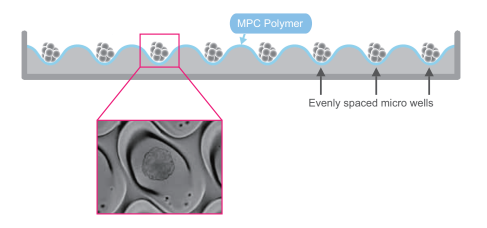
# **EZSPHERE®** for 3D cell culture with microwells

Three dimensional (3D) cell culture systems have gained in popularity as invaluable tools in broad applications of cell biology. 3D multi-cellular cell aggregates (Spheroid) can be formed by using a low attachment culture surface. However, variability in forming spheroids has been a persistent problem. EZSPHERE® is specifically designed to form a large number of uniformly sized spheroids and embryoid bodies (EBs).



- » Coated with very low binding 2-methacryloyloxyethyl phosphorylcholine (MPC) polymer
- » Form uniformly sized spheroids efficiently in the round shape wells

### **Well Cross-section**



### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
EZSPHERE® SP, 35 mm Dish	RT	4000-900SP	10 ea
EZSPHERE® SP, 60 mm Dish	RT	4010-900SP	10 ea
EZSPHERE® SP, 100 mm Dish	RT	4020-900SP	10 ea
EZSPHERE® SP, 24-well Plate	RT	4820-900SP	5 ea
EZSPHERE® SP, 96-well Plate	RT	4860-900SP	5 ea
EZSPHERE® SP, 35 mm Dish Type 902	RT	4000-902SP	10 ea
EZSPHERE® SP, 35 mm Dish Type 903	RT	4000-903SP	10 ea
EZSPHERE® SP, 35 mm Dish Type 904	RT	4000-904SP	10 ea
EZSPHERE® SP, 35 mm Dish Type 905	RT	4000-905SP	10 ea
EZSPHERE® SP Microplate 6-well EZ-TRY	RT	TCI-4810-EZ-TRY-SP-N	5 ea
EZSPHERE® SP Microplate 24-well EZ-TRY	RT	TCI-4820-EZ-TRY-SP	5 ea

[Storage] RT = Room Temperature, R = Refrigerator, F = Freezer

EZSPHERE® cell culture dishes are produced by AGC

# Serum-free Cell Freezing Media: Cell Reservoir One

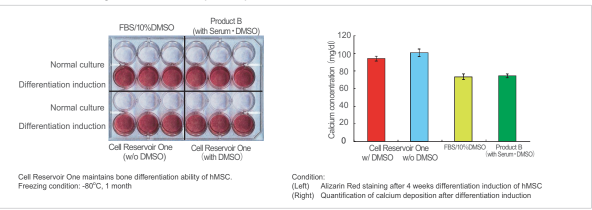
Cell Reservoir One is a serum-free cell culture freezing medium, which contains a water-soluble glycoprotein sericin isolated from the silkworm cocoon as a major constituent. Sericin shows the same high efficacy of cryopreservation as with FBS, and reduces the cell toxicity of DMSO. As DMSO is known to have adverse effects on cellular functions, especially embryonic stem cells, Cell Reservoir One is available both with and without DMSO.

- » No programmed freezer or special vessel necessary
- » Ready-to-use solution
- » Serum-free with no animal-derived components
- » High cell recovery and viability



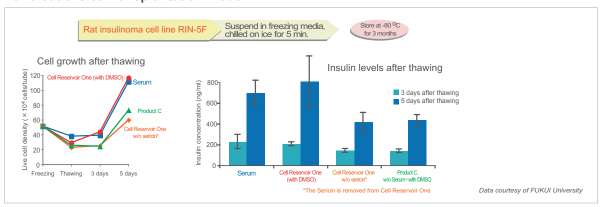
### **Application 1**

### Human Mesenchymal Stem Cell (hMSC): Bone Differentiation



### **Application 2**

### **Pancreatic Islet Transplantation Model**



### **Procedure for Cell Freezing**

- 1. Collect cells in logarithmic growth phase.
- 2. Suspend the cells in Cell Reservoir One (5 x 10<sup>5</sup> 1 x 10<sup>7</sup> cells in 1 ml of Cell Reservoir One).
- 3. Dispense the suspension to a cryo-cell tube.
- 4. Store it at -80°C without pre-freezing.

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Cell Reservoir One (with DMSO)	R	07485-44	100 ml
Cell Reservoir One (without DMSO)	R	07579-24	100 ml

# ES/iPS cell Freezing Media: Cell Reservoir One, Vitrify

Vitrification has become an important alternative to standard slow programmable freezing methods for cryopreservation of primate ES cell lines including Human iPS cells because of the higher survival rates of cells after thawing. However, the vitrification requires an ultra-rapid freezing protocol, usually less than 15 seconds between making cell suspensions and freezing in liquid nitrogen. Cell Reservoir One (Vitrify) is a novel serum-free cell culture freezing medium for vitrification method, which contains a water-soluble glycoprotein sericin isolated from the silkworm cocoon as a major constituent. It provides high survival rates of primate cells, such as Monkey ES cells and Human iPS cells even with a longer freezing protocol; up to 60 second from the cell collection to freezing in liquid nitrogen.

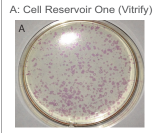
\*Cell Reservoir One (Vitrify) is produced in corporation with SEIREN. (Patented)

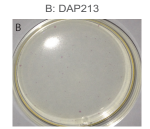
- » High viability with a longer freezing protocol (up to 60 seconds)
- » Low toxicity to cells (DMSO and acetamide free)

### **Application**

Comparison of survival rate of Human iPS cells (201B7 cell line\*) \*Takahashi, K. et al. Cell, Nov 30;131(5):861-872 (2007)

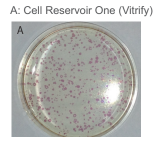
### Freezing protocol: 60 seconds

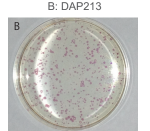




Human iPS cells were cryopreserved for more than 2 weeks in Cell Reservoir One (Vitrify) or DAP213. Viability was detected using Alkaline Phosphatase 4 days after thawing. Cell Reservoir One (Vitrify) showed high survival rate, while most of cells in DAP213 were dead.

### Freezing protocol: 15 seconds







Human iPS cells were cryopreserved for more than 2 weeks in Cell Reservoir One (Vitrify), DAP213 or Company A's product. Viability was detected using Alkaline Phosphatase 4 days after thawing. Cell Reservoir One (Vitrify) showed the highest viability.

Data courtesy of a customer

### Conclusion

Cell Reservoir One (Vitrify) showed high viability with both 15 and 60 seconds of freezing protocol. With 60 seconds protocol, the survival rate of cells in Cell Reservoir One (Vitrify) was significantly higher than other freezing media.

Freezing Medium		The Number of Colony					
		Vitrification	Slow Freezing Method				
		60 Seconds	15 Seconds	Slow Freezing Method			
Α	Cell Reservoir One (Vitrify)	672	563	-			
В	DAP213	37	479	-			
С	Company A	-	-	172			

### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Cell Reservoir One, Vitrify	R	11325-62	25 ml

### **Related product**

Product Name	Grade	Storage	Product No.	PKG size
Dimethyl Sulfoxide	SP (For TC)	RT	13408-64	5 x 5 ml
Glycerol	SP (Biotechnology grade)	RT	09886-05	500 ml
Propylene Glycol	SP (Biotechnology grade)	RT	11740-25	500 g

### **Balanced Salines**

» Quality control items: pH, Osmotic pressure, Sterility test, Endotoxin test, Mycoplasma test

D-PBS(-) w/o Ca and Mg, liquid 14249-95



D-PBS(-) w/o Ca and Mg, powder 07269-84



D-PBS(+) Preparation Reagent (Ca,Mg Solution) (100x) 02492-94



### **Product list and ordering information**

Product Name	KCI	Ca • Mg	Phenol Red	Storage	Product No.	PKG size	GIBCO	Sigma
D. DDS/ ) without Co and Ma. liquid				RT	14249-95	500 ml	14190-144	D8537
D-PBS(-) without Ca and Mg, liquid	$\bigcirc$	-	-	Γί	14249-24	10 x 500 ml	14190-250	
D-PBS(-) without Ca and Mg, liquid(10x)	0	-	-	RT	11482-15	500 ml	14200-075	D1408
D-PBS without Ca and Mg, Powder	0	-	-	RT	07269-84	100 g	21600-069	D5652
Phosphate Buffered Saline without KCI(pH 7.2)	-	-	-	RT	11480-35	500 ml	20012-027	
Phosphate Buffered Saline without KCl(10x)(pH 7.2)	-	-	-	RT	11481-25	500 ml	70013-032	
Phosphate Buffered Saline without KCI(pH 7.4)	-	-	-	RT	13397-85	500 ml	10010-023	
D-PBS(+)Preparation Reagent(Ca, Mg Solution)(100x)	-	0	-	RT	02492-94	30 ml		
HBSS(+) with Ca, Mg and Phenol Red, liquid	0	0	0	RT	17459-55	500 ml	24020-117	H9269
HBSS(+) with Ca, Mg, without Phenol Red, liquid	0	0	-	RT	09735-75	500 ml	14025-092	H8264
HBSS(-) without Ca and Mg, with Phenol Red, liquid	0	-	0	RT	17460-15	500 ml	14170-112	H9394
HBSS(-) without Ca, Mg and Phenol Red, liquid	0	-	-	RT	17461-05	500 ml	14175-095	H6648

# Sterile water

» Quality control items: Sterility test, Endotoxin test, Nuclease test, Protease test

### **Product list and ordering information**

Product Name	Grade	Product No.	Storage	PKG size	GIBCO	Sigma
Water deionized & sterilized	SP (For molecular biology)	06442-95	RT	500 ml	15230-162	W3500
Water deionized & sterilized	SP (For molecular biology)	20620-31	F	10 x 1 ml	15230-162	-

### **Media additives**

### Prepared media additive solutions

» Quality control items: Sterility test, Endotoxin test, Mycoplasma test

### **Product list and ordering information**

Product Name	Grade	Storage	Product No.	PKG size	GIBCO	Sigma
200mmol/l L-Alanyl-L-glutamine Solution(100x)	SP (For TC)	R	04260-64	100 ml	35050-061	G8541
200mM-L-Glutamine Stock Solution	SP (For TC)	F	16948-04	100 ml	25030-081	G7513
0.1mol/l-Calcium Chloride Solution	SP (For TC)	RT	16973-64	20 ml		
45(w/v)%-D-(+)-Glucose Solution	SP (For TC)	RT	16974-54	100 ml		G8769
1mol/I-HEPES Buffer Solution	SP (For TC)	R	17557-94	100 ml	15630-080	H0887
MEM Non-Essential Amino Acids	SD (For TC)		06344-14	20 ml	11140-076	
Solution(100x)	SP (For TC)	R	06344-56	100 ml	11140-050	M7145
100mM-Sodium Pyruvate Solution(100x)	SP (For TC)	R	06977-34	100 ml	11360-070	S8636

### Amino acids, vitamins and others

» Quality control items: Endotoxin test, cell culture test

### **Product list and ordering information**

Product Name	Grade	Storage	Product No.	PKG size
Amino Acids				
L-α-Alanine, Animal-Free	SP (For TC)	RT	12998-32	25 g
L-Alanyl-L-glutamine, Animal-Free	SP (For TC)	R	18189-52	25 g
I Argining Animal Frag	CD (For TC)	RT	11984-32	25 g
L-Arginine, Animal-Free	SP (For TC)	KI	11984-45	500 g
L-(+)-Arginine Monohydrochloride, Animal-Free	SP (For TC)	RT	12999-22	25 g
L-Asparagine Monohydrate, Animal-Free	SP (For TC)	RT	13000-42	25 g
L-Aspartic Acid, Animal-Free	SP (For TC)	RT	13002-22	25 g
L-Cysteine Hydrochloride Monohydrate, Animal-Free	SP (For TC)	R	11983-42	25 g
L-Cysteme nydrochlonde Mononydrate, Animai-Free	SP (FOLIC)	K	11983-55	500 g
L-Cystine Dihydrochloride, Animal-Free	SP (For TC)	RT	13003-12	25 g
L-Glutamic Acid, Animal-Free	SP (For TC)	RT	13012-92	25 g
L-Glutamine, Animal-Free	SP (For TC)	RT	13004-02	25 g
L-Glutamine, Animai-Free	5P (F0I 1C)	KI	13004-15	500 g
Glycine, Animal-Free	SP (For TC)	RT	12997-42	25 g
L-Histidine(free base), Animal-Free	SP (For TC)	RT	13014-72	25 g
L-Histidine Monohydrochloride Monohydrate, Animal-Free	SP (For TC)	RT	13017-42	25 g
L-Hydroxyproline, Animal-Free	SP (For TC)	RT	13018-32	25 g
L-(+)-Isoleucine, Animal-Free	SP (For TC)	RT	13035-02	25 g
L-Leucine, Animal-Free	SP (For TC)	RT	13036-92	25 g
L-Lysine Monohydrochloride, Animal-Free	SP (For TC)	RT	13037-82	25 g
L-Methionine, Animal-Free	SP (For TC)	RT	13038-72	25 g
L-(-)-Phenylalanine, Animal-Free	SP (For TC)	RT	13039-62	25 g
L(-)-Proline, Animal-Free	SP (For TC)	RT	13040-22	25 g
L-Serine, Animal-Free	SP (For TC)	RT	13041-12	25 g
L-Threonine, Animal-Free	SP (For TC)	RT	13042-02	25 g
L-Tryptophan, Animal-Free	SP (For TC)	RT	13043-92	25 g
I. T	CD (F TO)		11985-22	25 g
L-Tyrosine Disodium Salt Dihydrate, Animal-Free	SP (For TC)	R	11985-35	500 g
L-Valine, Animal-Free	SP (For TC)	RT	13046-62	25 g

Vitamins				
p-Aminobenzoic Acid, Animal-Free	SP (For TC)	RT	13047-52	25 g
L(+)-Ascorbic Acid, Animal-Free	SP (For TC)	RT	13048-42	25 g
D-Biotin, Animal-Free	SP (For TC)	RT	13049-61	1 g
Choline Chloride, Animal-Free	SP (For TC)	R	13050-34	5 g
myo-Inositol, Animal-Free	SP (For TC)	RT	13051-82	25 g
Pyridoxine Hydrochloride, Animal-Free	SP (For TC)	RT	13053-04	5 g
Vitamin B1 Hydrochloride, Animal-Free	SP (For TC)	RT	13052-14	5 g
Vitamin B12, Animal-Free	SP (For TC)	R	13054-94	100 mg
Others				
D-(+)-Glucose, Animal-Free	SP (For TC)	RT	13057-35	500 g
Glutathione(Reduced Form), free acid,	SP (For TC)	R	13056-61	1 a
Animal-Free	3F (10110)		13030-01	1 g
Hypoxanthine, Animal-Free	SP (For TC)	RT	13055-71	1 g
Sodium Pyruvate, Animal-Free	SP (For TC)	R	13058-12	25 g

### Low Molecular Compounds

### (Solution)

» Quality control items: Sterility test, Mycoplasma test, Endotoxin test

### **Product list and ordering information**

Product Name	Grade	Storage	Product No.	PKG size
AscleStem® CHIR99021 Solution(10mM), Animal-Free	SP (For TC)	F	21068-74	5 x 100 μL
AscleStem® SB 431542 Solution(10mM), Animal-Free	SP (For TC)	F	21235-74	5 x 100 μL
AscleStem® IWP-2 Solution(5mM), Animal-Free	SP (For TC)	F	21236-64	5 x 100 μL
AscleStem® Estradiol Solution(1mM), Animal-Free	SP (For TC)	F	21237-25	500 μL
10mmol/L-Y-27632 Solution	OD (F . TO)		18190-96	100 µL
1011111101/L-1-27032 Solution	SP (For TC)	Г	18190-54	1.5 ml

### (Powder)

» Quality control items: refer to the specification sheets for each product.

### **Product list and ordering information**

Product Name	Grade	Storage	Product No.	PKG size	
170 Fetradial Animal Free	SP	В	10100 00	25 mg	
17β-Estradiol, Animal-Free	(For biochemical research)	R	18192-92	25 mg	
NA/D O A I F	SP	D.	10175.01	5	
IWP-2, Animal-Free	(For biochemical research)	R	18175-64	5 mg	
CD 424540 Animal Fra	SP	DT	10170.51	_	
SB 431542, Animal-Free	(For biochemical research)	RT	18176-54	5 mg	
V.07000 A I. F.	SP	F		40	
Y-27632, Animal-Free	(For biochemical research)	F	18188-04	10 mg	

### Recombinant Proteins

### (Solution)

» Quality control items: Sterility test, Endotoxin test

### **Product list and ordering information**

Product Name	Grade	Storage	Product No.	PKG size
			18585-36	10 µg
Activin A Solution, Human, Recombinant	SP (For TC)	F	18585-94	50 μg
			18585-81	1 mg
hEGE (harris EGE) Calatian Harran Darambin ant (454 a.s.)			19155-07	10 µg
bFGF (basic FGF) Solution, Human, Recombinant (154 a.a.), Animal-Free	SP (For TC)	F	19155-36	5 x 10 μg
Animai-Free			19155-81	1 mg
			21065-04	10 µg
SCF Solution, Human, Recombinant	SP (For TC)	F	21065-46	50 µg
			21065-91	1 mg
			21185-34	10 µg
KGF (FGF-7) Solution, Human, Recombinant	SP (For TC)	F	21185-76	50 µg
			21185-21	1 mg
			21957-24	10 µg
VEGF165 Solution, Human, Recombinant	SP (For TC)	F	21957-66	50 µg
			21957-11	1 mg
			22247-94	10 µg
BMP-4 Solution, Human, Recombinant	SP (For TC)	F	22247-36	50 µg
			22247-81	1 mg

### (Powder)

» Quality control items: refer to the specification sheets for each product.

### **Product list and ordering information**

Product Name	Grade	Storage	Product No.	PKG size
Albumin, Human, Recombinant expressed in Rice, Animal-Free	SP (For TC)	F	19597-01	1 g
		Г	19597-14	5 g
Insulin, Human, Recombinant expressed in Yeast, Animal-Free	SP (For TC)	F	12878-86	50 mg
ilisuilii, Huillalii, Necollibillalii expressed ili feast, Allillai-Free	3F (F01 1C)	r	12878-44	250 mg
		R	12879-34	100 mg
Transferrin, Human, Recombinant expressed in Rice, Animal-Free	SP (For TC)		12879-21	1 g
			12879-76	5 g

### Proteins

### (Powder)

» Quality control items: refer to the specification sheets for each product.

### **Product list and ordering information**

Product Name	Grade	Storage	Product No.	PKG size	
apo-Transferrin from Human	SP (For TC)	R	34401-84	100 mg	
			34401-55	500 mg	
Albumin, Bovine Serum, Low Endotoxin, pH5.2	SP	В	01861-84	10 g	
	(For biochemical research)	R	01861-97	100 mg	
Albumin, Bovine Serum, Fatty Acid Free, pH7.0	SP (For biochemical research)	R	08587-26	10 g	
			08587-42	25 g	
			08587-84	50 g	

# Reagents for cell dissociation

» Quality control items: pH, Sterility test, Mycoplasma test, Enzyme actinity (USP unit)

### **Product list and ordering information**

Product Name	Phenol Red	Buffer	Grade	Storage	Product No.	PKG size
0.5g/l-Trypsin/0.53mmol/l-EDTA Solution	-	HBSS(-)	SP (For TC)	F	35553-74	100 ml
0.5g/I-Trypsin/0.53mmol/I-EDTA Solution,	0	HBSS(-)	SP (For TC)	F	32778-34	100 ml
with Phenol Red			3F (F0FTC)	Г	32778-05	500 ml
2.5g/I-Trypsin/1mmol/I-EDTA Solution	-	HBSS(-)	SP (For TC)	F	35554-64	100 ml
2.5g/I-Trypsin/1mmol/I-EDTA Solution,	0	HBSS(-)	OD (F . TO)		32777-44	100 ml
with Phenol Red			SP (For TC)	F	32777-15	500 ml
5.0g/I-Trypsin/5.3mmol/I-EDTA Solution	-	HBSS(-)	SP (For TC)	F	35556-44	100 ml
2.5g/I-Trypsin Solution	-	HBSS(-)	SP (For TC)	F	35555-54	100 ml
25g/I-Trypsin Solution	-	Saline	SP (For TC)	F	18172-94	100 ml
0.2g/I-EDTA Solution	-	PBS(-)	SP (For TC)	R	14367-74	100 ml
0.5mmol/I-EDTA/PBS Solution	-	PBS(-)	SP (For TC)	RT	13567-84	100 ml
Accutase	0	PBS(-)	SP (For TC)	F	12679-54	100 ml
Accumax	-	PBS(-)	SP (For TC)	F	17087-54	100 ml

# **Antibiotics**

### **Product list and ordering information**

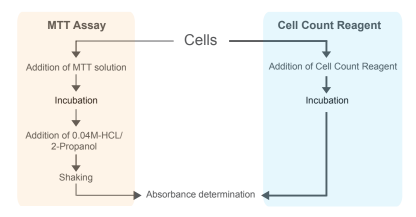
Product Name	Composition	Application	Grade	Storage	Product No.	PKG size
Actinomycin D Solution(1mg/ml)	1 mg/ml	Other	SP (For TC)	F	00393-41	1 ml
Antibiotic-Antimycotic Mixed Stock Solution(100x)	Penicillin 10,000 u/ml Streptomycin Potency: 10,000 µg/ml	Bacteria, Fungi, Yeast	SP (For TC)	F	02892-54	100 ml
Antibiotic-Antimycotic Mixed Stock Solution(100x)(Stabilized)	Amphotericin B Potency: 25 μg/ml		SP (For TC)	F	09366-44	100 ml
	-	Selection Antibiotics	SP (For TC)	RT	08973-01	1 g
G 418 Disulfate					08973-14	5 g
					08973-72	25 g
G 418 Disulfate Aqueous Solution	Potency: 50 mg/ml		SP (For TC)	R	09380-86	20 ml
G 418 Disulfate Aqueous Solution					09380-44	100 ml
C 419 Digulfoto	-		SP (For TC)	RT	08973-01	1 g
G 418 Disulfate					08973-14	5 g
G 418 Disulfate Aqueous Solution	Potency: 50 mg/ml		SP (For TC)	R	09380-86	20 ml
G 416 Disultate Aqueous Solution					09380-44	100 ml
Gentamicin Sulfate	-	Bacteria,	SP (For TC)	R	08975-81	1 g
Gentamicin Sunate					08975-94	5 g
Gentamicin Sulfate Solution(10mg/ml)	Potency: 10 mg/ml	Mycoplasma	SP (For TC)	R	16672-04	10 ml
Gentamicin Sulfate Solution(50mg/ml)	Potency: 50 mg/ml		SP (For TC)	R	11980-14	10 ml
	-	Selection Antibiotics	SP	R	07296-66	100 mg
Hygromycin B			(For biochemical research)		07296-11	1 g
					07296-24	5 g
Hygromycin B Solution	Potency: 50 mg/ml		SP (For TC)	R	09287-84	20 ml
Kanamycin Monosulfate	-	Bacteria, Mycoplasma	SP (For TC)	RT	08976-71	1 g
Ranamyon Monosulate					08976-84	5 g
Kanamycin Sulfate Solution(50mg/ml)	Potency: 50 mg/ml	Mycopiaema	SP (For TC)	R	11981-04	20 ml
Mitomycin C Solution(1mg/ml)	1 mg/ml	Other	SP (For TC)	F	20898-21	1 ml
Penicillin-Streptomycin Mixed Solution	Penicillin 10,000 u/ml		SP (For TC)	F	26253-84	100 ml
Penicillin-Streptomycin Mixed Solution (Stabilized)	Streptomycin Potency: 10,000 µg/ml		SP (For TC)	F	09367-34	100 ml
Penicillin-Streptomycin Mixed Solution	Penicillin 5,000 u/ml Streptomycin Potency: 5,000 μg/ml	Bacteria	SP (For TC)	F	26252-94	100 ml
Penicillin-Streptomycin-Glutamine Mixed Solution	Penicillin 10,000 u/ml Streptomycin Potency: 10,000 μg/ml L-Glutamine 29.2 mg/ml		SP (For TC)	F	06168-34	100 ml

# Cell Count Reagent SF, based on WST-8

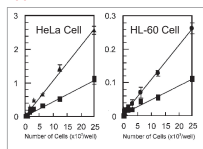
- » More sensitivity than other water-soluble tetrazolium salts, such as XTT and MTS
- » No radioisotope
- » Ready-to-use

### Comparison of Assay Procedure with MTT and Cell Count Reagent SF





#### **Application for Cell Proliferation Assay**

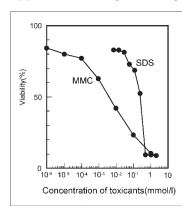


- 1. Prepare a cell suspension using an appropriate culture media, and dispense 100 µl of cell suspension into each well of a 96-well plate after counting cells.
- 2. Pre-incubate the plate in CO2 incubator.
- 3. Add 10 µl of Cell Count Reagent SF to each well.
- 4. Incubate the plate for 1-4 hours in the CO<sub>2</sub> incubator.
- 5. Measure the absorbance at 450 nm (calibration wavelength: 600 nm or more) by micro plate reader.

Incubation

: 5% CO<sub>2</sub>, 37°C, 1 hour ( $\blacksquare$ ), 2 hours ( $\blacktriangle$ ) Hela cells HL-60 : 5% CO<sub>2</sub>, 37°C, 1 hour (■), 3 hours (●)

#### **Application for Cytotoxicity Assay**



- 1. Prepare a cell suspension with 5,000 cells/well using an appropriate culture media, and dispense 100 µl of cell suspension into each well of a 96-well plate after counting cells.
- 2. Pre-incubate the plate in CO2 incubator for 24 hours.
- 3. Add 10 µl of a compoud prepared to appropriate concentration into each well.
- 4. Incubate the plate for 48 hours in the CO2 incubator.
- 5. Add 10  $\mu$ I of Cell Count Reagent SF to each well.
- 6. Incubate the plate for 1-4 hours in the incubator.
- 7. Measure the absorbance at 450 nm (calibration wavelength: 600 nm or more) by micro plate reader.

: Hela cells in DMEM (10% FCS) Cell

Compounds applied : MMC (Mitomycin C)

SDS (Sodium Dodecylsulfate)

5% CO2, 37°C, 48 hours / 5% CO2, 37°C, 2 hours Treatment / incubation period Wavelength

: 450 nm (reference: 650 nm)

#### References

M. Ishiyama, Y. Miyazono, K. Sasamoto, Y. Ohkura, K. Ueno, *Talanta*, 44, 1299 (1997)

H. Tominaga, M. Ishiyama, F. Ohseto, K. Sasamoto, T. Hamamoto, K. Suzuki and M. Watanabe, Anal. Commun, 36 (2), 47 (1999)

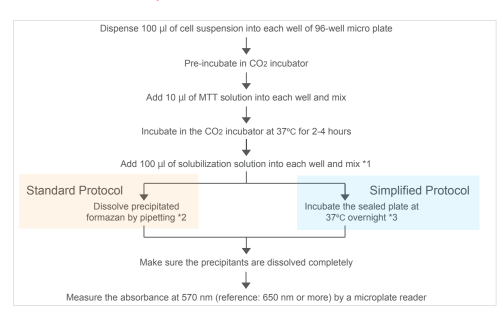
#### **Ordering Information**

<u> </u>				
	Product Name	Storage	Product No.	PKG Size
Call Count Desgent SE		В	07553-15	500 tests
Cell Count Reagent SF		K	07553-44	2500 tests

# MTT Cell Count Kit, based on reduction of MTT

- » No radioisotope
- » Ready-to-use

#### **Cell Proliferation Assay Procedure**



- \*1 Mix well Solubilization Solution and media as serum proteins might appear as precipitants.
- \*2 Aviod hard pipetting and shaking for a long time as that might help Solubilization Solution volatilize and affect the assay result.
- \*3 Make sure the plate is sealed completely. Alternatively, use a CO2 incubator at 37°C.

#### **Ordering Information**

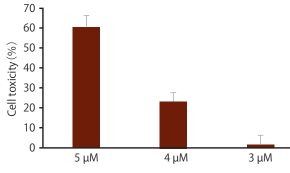
Product Name	Storage	Product No.	PKG Size
MTT Cell Count Kit	F	23506-80	1 KIT

# **LDH Cytotoxicity Assay Kit**

- » Economical packaging size (100 and 500-test sizes available)
- » Ready-to-use
- » Storable at 4°C for 3 months and at -20°C for 2 years without loss performance
- » By using cell suspension, enables unique assays with variety of other assays

#### **Application**

## Cell damage assay with melittin



Add represented concentration of melittin to 24-hour-cultured B16/BL6 cell (1 x 104 cells/well), then assay after 30-minute incubation. Substrate solution was directly added to each micro well. The higher concentration of melittin added, the stronger cell toxicity observed.

Date courtesy of Dr. Takuma Yamashita, Department of Biopharmaceutics and Drug Metabolism, Kyoto University

#### **Ordering Information**

	Product Name	Storage	Product No.	PKG Size
LDLI Cutatovicity Access Kit		_	18250-64	100 tests
LDH Cytotoxicity Assay Kit		Г	18250-35	500 tests

# **Trypan Blue Solution**

#### » Ready-to-use



## **Ordering Information**

	Product Name	Storage	Product No.	PKG Size
Trypan Blue Solution		RT	20577-34	100 ml

# Reagents for apoptosis research

## **Ordering Information**

Product Name	Grade	Storage	Product No.	PKG size
Annexin V-633 Apoptosis Detection Kit	SP	R	18146-44	100 tests
Alliexiii v-033 Apoptosis Detection Nit	(For fluorescence analysis)	IX	10140-44	100 tests
Annexin V-FITC Apoptosis Detection Kit	SP	R	15342-54	100 tests
Alliexiii V-FITC Apoptosis Detection Kit	(For fluorescence analysis)	K	15542-54	100 lesis
Actinomycin D Solution (1mg/ml)	SP (For TC)	F	00393-41	1 ml
Colcemid Solution (10µg/ml)	SP (For TC)	R	09356-74	10 ml
Mitomycin C Solution (1mg/ml)	SP (For TC)	F	20898-21	1 ml

# Reagents for alkaline phosphatase staining

## **Ordering Information**

Product Name	Grade	Storage	Product No.	PKG size
	SP			
BCIP-NBT Solution (Ready To Use)	(Reagents for alkaline phosphatase staining)	R	19880-84	100 ml
	priospriatase stairing)		09154-14	5 x 10 ml
4%-Paraformaldehyde Phosphate Buffer Solution	SP	R	09154-56	100 ml
	(For histochemistry research)		09154-85	500 ml

# Medium for Bacteria, Plusgrow II

Plusgrow II is high performance medium for bacteria that offers easy procedures for weighing, dissolving and autoclave treatments.

- » Higher fungus density than conventional products
- » High plasmid collection

# **Comparison with Conventional Products**

#### Bacteria growth test

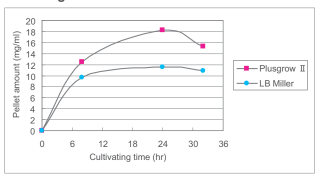


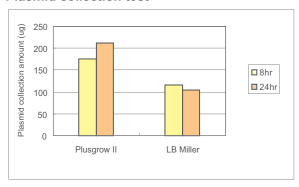


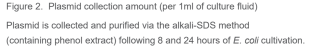
Figure 1. Bacteria growth curve

*E. coli* is first cultivated in ampicillin (50  $\mu$ g/ml) then added to the medium at 37°C and shaken. Then culture fluid is then centrifugally processed. Bacteria levels can then be evaluated by pellet amounts.

E. coli cell line : JM109
Plasmid : pGEM-3zf(+)

#### Plasmid collection test





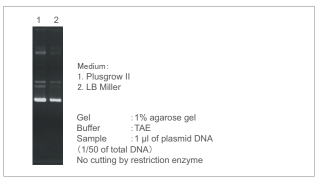


Figure 3. Electrophosis image of collection of plasmid (following 24 hours of cultivation)

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Plusgrow II (One package for 1L)	DT	08246-86	40 g
Dissolve $\rightarrow$ Autoclave at 121 °C for 15 minutes	RT	08246-44	10 x 40 g
Plusgrow II	DT	08202-04	100 g
Weigh out 40 g $\rightarrow$ Dissolve in 1L $\rightarrow$ Autoclave at 121 °C for 15 minutes	RT	08202-75	500 g

## **Related Products**

LB Agar, Lennox	RT		
22 / 1941, 201110/	1 ( )	20067-85	500 g
LB Agar, Miller	RT	20069-65	500 g
I.D. Droth Lanney	RT	20066-95	500 g
LB Broth, Lennox	KI	20066-24	2 kg
LB Broth, Miller	RT	20068-75	500 g
Agar, powder	RT	01028-85	500 g
Agar Purified, powder	RT	01162-15	500 g
Extract Yeast Dried	RT	15838-45	500 g
Tryptone	RT	35640-95	500 g

# **IPTG and X-gal Solutions**

# • 100mmol/l-Isopropyl-β-D-thiogalactopyranoside [IPTG] Solution

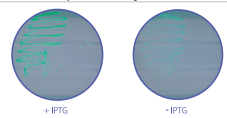
- » 0.22 µm filtrated ready-to-use solution
- » Sterilized
- » No need to adjust concentration
- » 1 ml package size allows for easy application



#### **Application**

Recombinant protein expressions are evaluated by the green fluorescent protein (GFP) expressing vector

#### **Induced expression by IPTG**



Numerous fluorescent proteins can be expressed by adding IPTG. In situations where IPTG is not applied, expressions occur at the basal level.

E. Coli : BL21 (DE3) pLysS (Novagen) : pQBI-T7-GFP (Q-Bio gene) Vector IPTG concentration: 1mM

: Transilluminator Detection

# • 5-Bromo-4-chloro-3-indolyl-β-D-galactoside [X-gal] Solution (20 mg/ml)

5-Bromo-4-chloro-3-indolyl-β-D-galactoside (X-Gal) is widely used for Blue-white selection.

- » Ready-to-use DMF solution
- » 1 ml package size allows for easy application



## **Ordering Information**

Product Name	Storage	Product No.	PKG Size
100mmol/l-Isopropyl-β-D-thiogalactopyranoside [IPTG] Solution	F	07496-91	10 x 1 ml
5-Bromo-4-chloro-3-indolyl-β-D-galactoside Solution(20 mg/ml)	F	03971-71	10 x 1 ml

#### Related product

Product Name	Storage	Product No.	PKG Size
		19742-36	100 mg
Isopropyl-β-D-thiogalactopyranoside [IPTG], Dioxane free		19742-81	1 g
		19742-94	10 g
5-Bromo-4-chloro-3-indolyl-α-D-galactoside [X-α-Gal]	R	02897-04	100 mg
		05627-86	10 mg
		05627-57	100 mg
5-Bromo-4-chloro-3-indolyl-β-D-galactoside [X-Gal]	R	05627-31	1 g
		05627-44	5 g
		05644-14	5 x 20 mg
5-Bromo-4-chloro-3-indolyl-β-D-qlucuronide Cyclohexylammonium Salt	F	05646-94	10 mg
5-biomo-4-cinoro-5-indoryi-p-b-glacuronide Cyclonexylammonium Sait	Г	05646-36	100 mg

# Gelling Agent for Plant Study: Gellan Gum

## » High Transparency

# Comparison Data with Agar Gel: Root growth observation

0.8% Agar Gel





0.2% Gellan Gum

These photos were taken on the 6th day.

Temperature : 27 °C Light Period : 13 hours Dark Period : 11 hours

Seeds of Komatsuna (Brassica rapa var. perviridis) were incubated in petri dishes containing a MS medium with either 0.8% agar or 0.2% Gellan Gum under sterile conditions. The seedlings were transferred to plant boxes containing the same medium 5 days later.

## **Ordering Information**

	Product Name	Storage	Product No.	PKG Size
Gellan Gum		RT	12389-96	50 g
		KI	12389-54	250 g

# Zymolyase<sup>™</sup> (from Arthrobacter Luteus)

Zymolyase<sup>TM</sup>, produced by a submerged culture of Arthrobacter luteus<sup>(1)</sup>, has strong lytic activity against living yeast cell walls<sup>(2),(3)</sup> to produce protoplast or spheroplast of various strains of yeast cells. An essential enzyme for the lytic activity of Zymolyase<sup>™</sup> is β-1,3-glucan laminaripentaohydrolase. It hydrolyzes linear glucose polymers with β-1,3-linkages and releases specifically laminaripentaose as the main and minimum product unit (4), (5), (10), (11). There are two preparations of Zymolyase<sup>™</sup>, Zymolyase<sup>™</sup>-20T and Zymolyase<sup>™</sup>-100T, having lytic activity of 20,000 units/g and 100,000 units/g respectively. Zymolyase<sup>TM</sup>-20T is ammonium sulfate precipitate while Zymolyase<sup>TM</sup>-100T is a further purified preparation by affinity chromatography<sup>(9)</sup>. Lytic activity varies depending on yeast strain, growth stage of yeast, or cultural conditions<sup>(6-8)</sup>. Further information related to Zymolyase<sup>™</sup> can be obtained in the reference section below<sup>(12-16)</sup>.

#### **Specifications**

Product Na	ame	Zymolyase <sup>™</sup> -20T	Zymolyase <sup>™</sup> -100T	
Form		Lyophilized Powder		
Purification		Ammonium Sulfate Precipitation	Affinity Chromatography	
Activity		20,000 units/g	100,000 units/g	
Essential enzyme		β-1,3-glucan lamina	aripentaohydrolase	
	β-1,3-glucanase	approx. 1.5 x 10 <sup>6</sup> units/g	approx. 1.0 x 10 <sup>7</sup> units/g	
Other activities contained(*1)	protease	approx. 1.0 x 10 <sup>4</sup> units/g	approx. 1.7 x 10 <sup>4</sup> units/g	
	mannanase	approx. 1.0 x 10 <sup>6</sup> units/g	approx. 6.0 x 10 <sup>4</sup> units/g	
Contaminants	Amylase, Xylanase, Phosphatase	Trace amount	Not detectable	
Optimum pH and Temp.		pH7.5, 35°C (for lysis of viable yeast cells) pH6.5, 45°C (for hydrolysis of yeast glucan)		
Stability	2°C	No loss of activity was fou	nd after storage for 1 year	
Lleet etability	30°C	70% of the lytic activity is lost after storage for 3 months	90% of the lytic activity is lost after storage for 3 months	
Heat stability 60°C		Lytic activity is lost on incubation for 5 minutes		
Specificity (Lytic Spectrum)		Debaryomyces, Eremothecium, Endomyces, Hansenula, Hanseniaspora, Kloeckera, kluyveromyces, hnikowia, Pichia, Pullularia, Torulopsis, Saccharomyces, Saccharomycopsis, Saccharomycodes, etc.		

<sup>(\*1)</sup> See reference, Kitamura, K., Kaneko, T., Yamamoto, Y., J. Gen. Appl. Microbiol., 18, 57 (1972) as to the definition of each enzyme units.

#### **Unit Definition**

One unit of lytic activity is defined as that amount which indicates 30% of decrease in absorbance at 800 nm (A800) of the reaction mixture under the following condition.

[Reaction Mixture]

Enzyme solution : 1 ml (0.05-0.1 mg/ml for Zymolyase<sup>™</sup>-20T)

(0.012-0.024 mg/ml for Zymolyase<sup>™</sup>-100T)

Brewer's yeast cell suspension : 3 ml (2 mg/ml) 1/15M Phosphate buffer : 5 ml (pH7.5)

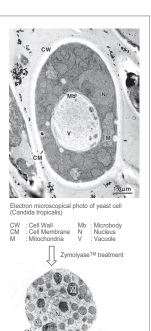
After incubation for 2 hours at 25°C with gentle shaking, A®0 of the mixture is determined. When 60% of A®0 decrease, equivalent to 2 units, is observed in the reaction system, the brewer's yeast cells are completely lysed, namely 1 unit of Zymolyase<sup>™</sup> lyses 3 mg dry weight of brewer's yeast.

#### Reference

- 1. Kaneko, T., Kitamura, K and Yamamoto, Y.: J. Gen. Appl. Microbiol., 15, 317 (1969)
- 2. Kitamura, K., Kaneko, T. and Yamamoto, Y.: Arch. Biochem. Biophys., 145, 402 (1971)
- 3. Kitamura, K., Kaneko, T. and Yamamoto, Y.: J. Hen. Appl. Microbiol., 18, 57 (1972)
- 4. Kitamura, K. and Yamamoto, Y.: Arch. Biochem. Biophys., 153, 403 (1972)
- 5. Kaneko, T., Kitamura, K. and Yamamoto, Y.: Agric. Biol. Chem., 37, 2295 (1973)
- 6. Kitamura, K., Kaneko, T. and Yamamoto, Y.: J. Gen Appl. Microbiol., 20, 323 (1974)
- 7. Kitamura, K. and Yamamoto,.: Agric. Biol. Chem., 45, 1761 (1981)
- 8. Katamura, K. and Tanabe, K.: *Agric, Biol. Chem.*, **46**, 553 (1982) 9. Katamura, K.: *J. Ferment. Technol.*, **60**, 257 (1982)
- 10. Kitamura, K.: Agric. Biol. Chem., 46, 963 (1982)
- 11. Kitamura, K.: Agric. Biol. Chem., 46, 2093 (1982)
- 12. Calza R. E., Schroeder A. L.: J. Gen. Microbiol., 129, 413 (1983)
- 13. lizuka Masaru, Torii Yasuhiko, Yamamoto Takehiko: Agric. Biol. Chem., 47 (12), 2267 (1983)
- 14. Shibata Nobuyuki, Kobayashi Hidemitsu, tojo Menehiro, Suzuki Shigeo: *Arch. Biochem. Biophys.*, **251** (2), 697 (1986)
- 15. lijima Y., Yanagi S. O.: Agric. Biol. Chem., 50 (7), 1855 (1986)
- 16. Herrero Enrique, Sanz Pascual. Sentandreu Rafael: J. Gen. Microbiol., 133 (10), 2895 (1987)

## **Ordering Information**

	Product Name	Storage	Product No.	PKG Size
Zymolyase <sup>™</sup> 20T		R	07663-91	1 g
Zymolyase <sup>™</sup> 100T		R	07665-55	500 mg



# **Cell Lysis Solution: Cell Lysis Buffer**

Cell Lysis Buffer is designed for protein extraction, similar to RIPA Buffer, and is available in two forms: a ready-to-use type and a 10x concentrated type. The 10x concentrated type is packaged separately with SDS solution, which may adversely affect immunoprecipitation.

- » Available for protein extraction in the same way as RIPA Buffer
- » Does not include OPE/NPE (e.g., NP-40), substances regulated under REACH
- » Available in two forms

#### Lineup

Available in two forms as below.

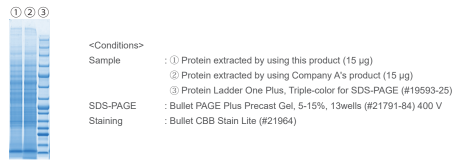
Product name	Cell Lysis Buffer	Cell Lysis Buffer (10x)
Product type	1x (ready to use)	10x
Protease inhibitor cocktail	Not included	Included
SDS	Included	Not included (10x SDS solution is packed separately)
Storage	Refrigerator	Freezer

#### Comparison

Protein extraction by using Cell Lysis Buffer or Company A's product

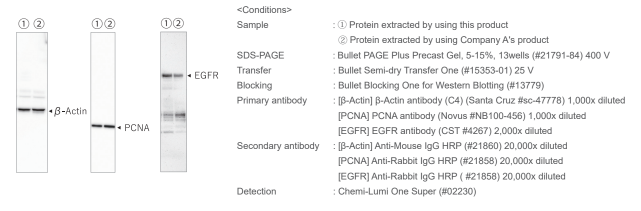
#### (CBB staining)

Protein was extracted from rat kidney by using this product or Company A's, then subjected to electrophoresis, and CBB staining. The result shows protein extraction efficiency of this product is equal to that of Thermo's RIPA Buffer.



## (Western blotting)

Protein was extracted from mouse liver by using this product or Company A's, then subjected to western blotting. The results show that this product is applicable to western blotting, similar to Company A's.

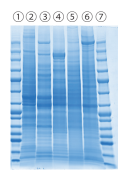


#### **Applications**

Protein extraction by using Cell Lysis Buffer (10x)

#### [CBB staining]

Protein was extracted from various mouse organs by using this product with SDS, subject to electrophoresis and CBB staining.



<Conditions>

SDS-PAGE

Stain

Sample : ①, ⑦ Protein Ladder One Plus, Triple-color for SDS-PAGE (#19593-25)

2 Protein extracted from brain (12 µg) 3 Protein extracted from heart (12 µg) 4 Protein extracted from liver (12 µg)

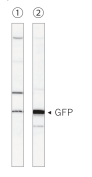
⑤ Protein extracted from kidney (12 μg)
⑥ Protein extracted from stomach (12 μg)

: Bullet PAGE Plus Precast Gel, 5-15%, 13wells (#21791-84) 400 V

: Bullet CBB Stain Lite (#21964)

# [Immunoprecipitation]

Protein was extracted from GFP-expressing HEK293 cells using this product without SDS and immunoprecipitated using rat anti-GFP antibody-conjugated agarose beads. GFP was detected by western blotting of the samples before and after immunoprecipitation. GFP in the samples before and after immunoprecipitation was detected by western blotting.



<Conditions>

Sample : ① Sample before immunoprecipitation (3 µL)

② Sample after immunoprecipitation (3 µL)

SDS-PAGE : Bullet PAGE Plus Precast Gel, 5-15%, 13wells (#21791-84) 400 V

Transfer : Bullet Semi-dry Transfer One (#15353-01) 25 V

Blocking : Bullet Blocking One for Western Blotting (#13779)

Primary antibody : Anti-GFP (Mouse IgG1-k) (#04363) 5,000x diluted

Secondary antibody : Anti-Mouse IgG HRP (#21860) 20,000x diluted

Secondary antibody . Anti-wouse 199 mit (#2 1000) 20,000x dilute

Detection : Chemi-Lumi One Super (#02230)

#### **Ordering Information**

	Product Name	Storage	Product No.	PKG Size
Cell Lysis Buffer		R	22352-04	100 ml
Cell Lysis Buffer (10x)		F	22353-81	1 set

#### **Related Products**

Product Name	Storage	Product No.	PKG Size
Protease Inhibitor Cocktail for General Use (100x)	F	04080-11	5 x 1 ml
Protease Inhibitor Cocktail (EDTA free) (100x)	F	03969-21	5 x 1 ml
Protease Inhibitor Cocktail for Use with Mammalian Cell and Tissue Extracts	F	25955-11	5 x 1 ml
Phosphatase Inhibitor Cocktail (EDTA free)	R	07575-51	1 ml
Phosphatase Inhibitor Cocktail	R	07574-61	1 ml

# **Protease Inhibitor Cocktail**

Inhibition of intra and extra cellular proteases is vital to purify and collect the expressed proteins. Saving trouble of finding adequate inhibitors, a wide range of protease is inhibited by the Protease Inhibitor Cocktail.

- » Contains inhibitors for a variety of protease
- » Available in 3 types: General use, Mammalian cell and tissue, and EDTA free



# **Composition of Each Protease Inhibitor Cocktail**

-				
Inhibitors	Target Protease	#04080-11	#03969-21	#25955-11
4-(2-Aminoethyl) benzenesulfonyl fluoride hydrochloride (AEBSF)	Serin protease	~	~	~
Aprotinin	Serin protease and Esterase	<b>~</b>	<b>~</b>	<b>~</b>
E-64	Cysteine protease	<b>~</b>	<b>~</b>	<b>~</b>
Leupeptin hemisulfate monohydrate	Cysteine protease and Trypsin-like protease	<b>~</b>	<b>~</b>	<b>~</b>
Disodium dihydrogen ethylenediaminetetraacetate dihydrate	Metalloprotease	<b>~</b>		
Bestatin	Aminopeptidase and Leucine aminopeptidase			<b>~</b>
Pepstatin A	Aspartic protease			~

## Reference

- 1. Okada, S. et al. The Journal of Cellular Physiology 226(2), 552-558 (2011)
- 2. Yang, JH. et al. The Journal of Biological Chemistry (2010)
- 3. Iyama, T. et al. Nucl. Acids Res. 38(14), 4834-4843 (2010)
- 4. Kimura, Y. et al. Cancer Research 70(2), 501-511 (2010)
- 5. Burnett, T. J. et al. J. Bacteriol 165, 139-145 (1986)
- 6. Hagiwara B et al. J. Biochem., 45, 185-194 (1958)

## **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Protease Inhibitor Cocktail for General Use (100x)	F	04080-11	5 x 1 ml
Protease Inhibitor Cocktail (EDTA free) (100x)	F	03969-21	5 x 1 ml
Protease Inhibitor Cocktail for Use with Mammalian Cell and Tissue Extracts	F	25955-11	5 x 1 ml

# **Phosphatase Inhibitor Cocktail**

Phosphatase Inhibitor Cocktail is a mixture of several inhibitors to protect valuable proteins from dephosphorylation. The product preserves phosphorylated proteins existing in small quantity in cells and tissues.

- » Contains 6 kinds of phosphatase inhibitors for different targets
- » 100 times concentrated stock solution
- » Compatible with protein assay
- » Ready-to-use



# **Composition of Each Phosphatase Inhibitor Cocktail**

Inhibitors	Target Phosphatase	#07575-51 EDTA free	#07574-61
Sodium orthovanadate (V)	Tyrosine phosphatase and Alkaline Phosphatase	~	<b>~</b>
Disodium molybdate (VI) dihydrate	Acid phosphatase	~	~
Sodium (+) -tartrate dihydrate	Acid phosphatase	~	~
Imidazole	Alkaline Phosphatase	~	~
Sodium fluoride	Acid phosphatase	~	~
b-Glycerophosphoric acid disodium salt	Serine-threonine phosphatase	~	~
tetra-Sodium ethylenediaminetetraacetate	Alkaline Phosphatase		~

<sup>\* 100</sup> times concentrated aqueous solution

#### **Comparison Data**

Figure 1.
The detection of phosphorylated proteins in Hela cells with Anti-p-Thr antibody

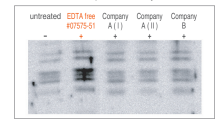
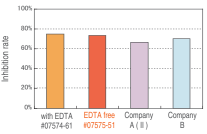
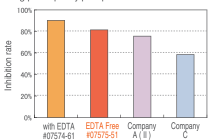


Figure 2.
The phosphatase inhibition efficiency assayed by fluorescence labeled p-Tyr peptide substrate



The phosphatase inhibition efficiency assayed using p-nitrophenylphosphoric acid



#### Reference

- 1. Yang, JH. et al. The Journal of Biological Chemistry (2010)
- 2. Selamat, W. et al. Neuroscience Letters 450(2), 163-166 (2009)
- 3. Saito, T. et al. Biochemical and Biophysical Research Communications 357(2), 371-376 (2007)
- 4. Murakami, Y. et al. J. Biochem., 141, 401-410 (2007)
- 5. Takenaga, M. et al. J. Cell Sci., 120, 2078-2090 (2007)

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Phosphatase Inhibitor Cocktail(EDTA free)	R	07575-51	1 ml
Phosphatase Inhibitor Cocktail	R	07574-61	1 ml

Immunohistochemistry

# **Determination of Protein Concentration: Protein Assay**

The protein assay is one of the most important key techniques in proteomics. To determine protein concentration, three products with spectrophotometric method are available as follows.

# **Comparison of Each Method**

Assay Method	Bradford	BCA	Lowry
Product Name	Protein Assay CBB Solution	Protein Assay BCA Kit	Protein Assay Lowry Kit
Linearity	1.6 1.2 0.8 0.8 0.5 1 1.5 2 Protein Concentration (mg/mL)	2 1.6 1.2 0.8 0.8 0.5 1 1.5 2 Protein Concentration (mg/mL)	2.4 2 1.6 1.6 1.2 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0
	+	+++	++
Convenience	+++	++	+
Absorbance	595 nm	562 nm	750 nm
Incompatible with	Detergents	Reducing Agents	Reducing Agents
	Condition: Mixing BSA (left: 0 mg/ml, righ	t: 1 mg/ml) and each substance described i	in the column left below
Incubate with Water			
Incubate with 0.1% SDS			
Incubate with 1 mM DTT			
Incubate with 0.1% SDS and 1 mM DTT			
Remarks	For protein samples containing detergents, BCA assay method or removal of detergents by CBB Clean Up Kit (Prod No. 11611) is helpful.	For protein samples containing reducing agents, BCA Kit with BCA Reductant Adaptable Reagent (Prod No. 21014) or the Bradford method is useful.	For protein samples containing reducing agents, the Bradford method is useful.

# **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Protein Assay CBB Solution (Ready To Use)	R	11617-71	1 L
Protein Assay BCA Kit	RT	06385-00	1 KIT
Protein Assay Lowry Kit	RT	29470-60	1 KIT

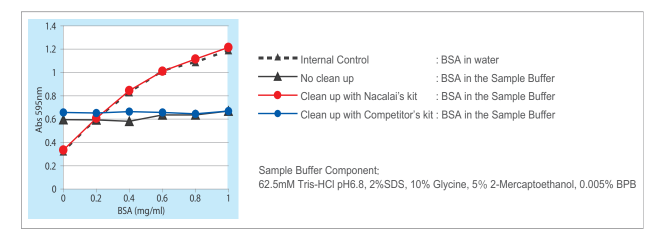
# **Protein Assay CBB Clean Up Kit**

- » Removal interfering substances within 5 minutes
- » Designed for the Bradford protein assay



#### Comparison data of determination of BSA concentration with competitor's clean up kit

Protein Assay CBB Clean Up Kit is helpful to get rid of reducing agents and detergents that cause interfering with the Bradford assay, and enables better quantitative assays compared to the competitor's clean up kit.



## Components

Reagent Name	Volume	Quantity
Solution A	2.5 ml	1 bottle
Solution B	2.5 ml	1 bottle
Solution C *Add 185 ml of ethanol (99.5%) into Solution C bottle, and mix it thoroughly	80 ml	2 bottles

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Protein Assay CBB Clean Up Kit	RT	11611-60	1 KIT
Protein Assay CBB Solution (Ready To Use)	R	11617-71	1 L
Albumin, Bovine Serum, Solution (2mg/ml) for Protein Assay	F	00653-31	10 x 1 ml

# Reductant adaptable reagent for protein assay BCA Kit

- » This product enables measurement of samples containing reducing agents
- » Quick and easy: Pretreatment takes 15 mins. and measurements are completed in the same well
- » Easy dispensing and homogenizing protocols

#### Comparison

Measurement of samples containing reducing agents and surfactants

	PBS	0.1% SDS	1 mM DTT	0.1% SDS 1 mM DTT	Effective for
BCA Kit					samples with both surfactants and reductants as well!
Product pre-treatment operation + BCA Kit					

<sup>\*</sup>Refer to the table of possible coexisting substances (instruction manual) for interfering substances that can be used in combination with reducing agents.

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Protein Assay BCA Reductant Adaptable Reagent	R	21014-80	1 KIT
Protein Assay BCA Kit	RT	06385-00	1 KIT
Albumin, Bovine Serum, Solution(2mg/ml) for Protein Assay	F	00653-31	10 x 1 ml

# COSMOGEL® Ig-Accept Protein G

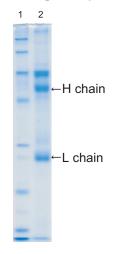
This product utilizes the binding of the Fc region of immunoglobulin (Ig) with Protein G, and is mainly used for the purification of IgG.

# **Specification**

Carrier Name	COSMOGEL® Ig-Accept Protein G
Ligand	Protein G
Particle size	50 - 150 μm
Matrix	4% Cross-linked agarose
Recommended Line Velocity	26 cm/h
Limit pressure	3.6 psi (0.25 bar)
Supply condition	50 vol% suspension (20 vol% ethanol solution)

# **Application**

# **COSMOGEL®** Ig-Accept Protein G



Sample : Mouse serum (ammonium sulfate precipitation fraction)

Elution : Glycine - Hydrochloric acid buffer (pH 2.8) Lane : 1. Protein Markers (10x) (#29458-24) in 1x

2. Sample purified by  $\mathsf{COSMOGEL}^{\otimes}$  Ig-Accept Protein G

## **Ordering information**

Product Name	Storage	Product No.	PKG Size
COSMOGEL® Ig-Accept Protein G	D	02198-64	5 ml
COSMOGEL 1g-Accept Protein G	K	02198-22	25 ml

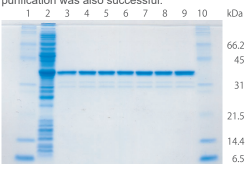
# **COSMOGEL® GST-Accept**

- » Superior durability
- » Superior cost performance

## **Application**

## 10 consecutive purifications of GST-tag fusion proteins

We performed 10 consecutive purifications of GST-His fusion proteins by using the same resin from *E. coli* lysate. The 10th purification was also successful.



Lane:

- 1. Use Protein Markers(10x)(#29458-24)1x
- 2. E. coli lysate
- 3. 1st elution fraction
- 4. 2nd elution fraction
- 5. 3rd elution fraction
- 6. 4th elution fraction
- 7. 6th elution fraction
- 8. 8th elution fraction
- 9. 10th elution fraction
- 10. Use Protein Markers(10x)(#29458-24)1x

Sample : E. coli (BL21(DE3)pLysS) lysate transformed with GST-His(pET-41b(+))

Binding buffer : Use Phosphate Buffered Saline(10x)(pH 7.4)(#27575-31)1x

Elution buffer : 10 mM Glutathione, 50 mM Tris-HCl, pH 8.0

## **Ordering information**

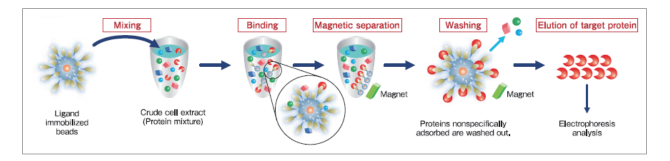
	Product Name	Storage	Product No.	PKG Size
			09277-56	5 ml
COSMOGEL® GST-Accept		R	09277-72	25 ml
			09277-14	100 ml

# High Performance Magnetic Nanoparticles: FG beads®

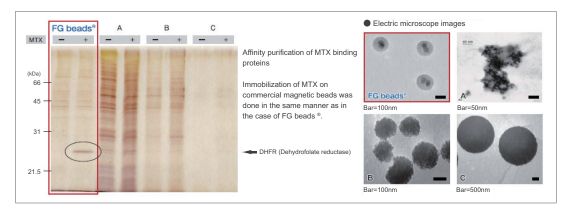
- » Excellent recovery of target proteins
- » Extremely low non-specific binding
- » High stability in organic solvents

The FG beads developed by Tokyo Institute of Technology consists of ferrite nanoparticles coated firmly with a polymer layer and its diameter is approx. 200 nm. The FG beads are used as carriers for affinity purification of target proteins.<sup>1)</sup>

#### **Purification Process**



# Comparison with Other Magnetic Beads 2)



- 1. S. Sakamoto et al., Chem. Rec. 9 (2009) 66
- 2. K. Nishio et al., Colloids Surfaces. B. 64 (2008) 162

#### **Ordering Information**

Product Name	Density	Storage	Product No.	PKG Size
Plain beads	20mg/ml	R	TAS8848N1010	0.5ml
Linker beads (Epoxy beads)	20mg/ml	R	TAS8848N1110	0.25ml
OH beads	20mg/ml	R	TAS8848N1120	0.25ml
NH2 beads	20mg/ml	R	TAS8848N1130	0.25ml
COOH beads	20mg/ml	R	TAS8848N1140	0.25ml
NHS beads	20mg/ml	F	TAS8848N1141	0.25ml
Ts beads	20mg/ml	R	TAS8848N1150	0.25ml
Azide beads	20mg/ml	R	TAS8848N1160	0.25ml
Alkyne beads	20mg/ml	R	TAS8848N1161	0.25ml
Streptavidin beads	20mg/ml	R	TAS8848N1170	0.25ml
NeutrAvidin beads	20mg/ml	R	TAS8848N1171	0.25ml
Protein A beads	20mg/ml	R	TAS8848N1172	0.25ml
Protein G beads	20mg/ml	R	TAS8848N1173	0.25ml

[Storage] RT = Room Temperature, R = Refrigerator, F = Freezer

FG beads® are produced by Tamagawa Seiki Co., Ltd.

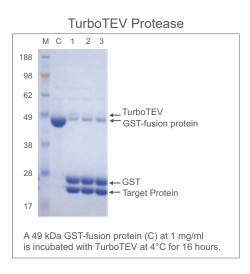
Product Name	Density	Storage	Product No.	PKG Size
HM-Streptavidin beads	20mg/ml	R	TAB8848N3170	0.25ml
HM-NeutrAvidin beads	20mg/ml	R	TAB8848N3171	0.25ml
HM-Protein A beads	20mg/ml	R	TAB8848N3172	0.25ml
HM-Protein G beads	20mg/ml	R	TAB8848N3173	0.25ml
Magnetic Stand (for 1.5 ml tube)		RT	TAB4899N12	1ea
Magnetic Stand (for 15 ml tube)		RT	TAB4899N20	1ea
Magnetic Stand (for 50 ml tube)		RT	TAB4899N30	1ea
Magnetic Stand (for PCR tube)		RT	TAB4899N41	1ea

FG beads® are produced by Tamagawa Seiki Co., Ltd.

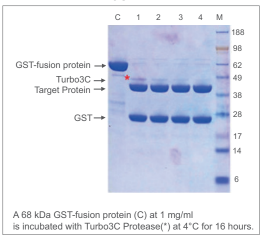
# **TurboTEV Protease & Turbo3C Protease**

- » Both GST and His tags to facilitate its removal from the digested protein sample
- » Activity over a broad temperature (4°C to 37°C) and pH (6.5 to 8.5) range

#### **Application**







GST-fusion protein (C) at 1 mg/ml is incubated with TurboTEV or Turbo3C Protease at a ratio of (1) 1:50, (2) 1:100, (3) 1:200 (w/w) in a buffer of 25 mM Tris-HCl, pH 8.0, 150 mM NaCl, 14 mM 2-mercaptoethanol at 4°C for 16 hours.

#### **Specification**

Molecular Weight	52 kDa	47 kDa		
Cleavage Site	Glu-Asn-Leu-Tyr-Phe-Gln↓Gly	Leu-Glu-Val-Leu-Phe-Gln↓Gly-Pro		
Enzymatic Activity	One unit cleaves >85% of 3 µg control substrate at 30°C	One unit cleaves >95% of 100 µg control substrate at		
	for 1 h	4°C for 16 h		
Concentration	20,000 units/ml 2,000 units/ml			
Cleavage Condition	A broad temperature (4°C to 37°C) and pH (6.5 to 8.5) range			
Formulation	25 mM Tris-HCl(pH8.0), 50 mM NaCl, 1 mM TCEP, 50% Glycerol			

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
	•	NU0102S	1,000 units (0.1 mg)
TurboTEV (TEV Protease) 2 mg/ml	F	NU0102M	10,000 units ( 1 mg)
		NU0102L	100,000 units ( 10 mg)
T. (1-20 (11D)/20 Dente ) 2 (1-1)	F	NU0101S	1,000 units ( 1 mg)
Turbo3C (HRV3C Protease) 2 mg/ml	F	NU0101M	10,000 units ( 10 mg)

[Storage] RT = Room Temperature, R = Refrigerator, F = Freezer

TurboTEV Protease & Turbo3C Protease are produced by Nacalai USA.

[Ultra-fast Precast Gel]

# **Bullet PAGE Plus Precast Gel**

- » Only 10 minutes with 400 V
- » Wide range of gel concentrations available
- » Compatible with conventional Laemmli running buffer and sample buffer
- » Surfactant-free, so can be used for nucleic acid analysis (refer to p11)

### **Performance Comparison**

This product demonstrates separation ability and band sharpness comparable to competitors' gels, despite a short running time of about 10 minutes.

Product Name	Bullet PAGE Plus	Company A	Company B
Gel %	5 - 20%	4 - 20 %	4 - 12%
Prod No.	21794-54	-	-
Running Time	10 m 07 s	29 m 35 s	48 m 32 s
Separation Image	M.W.(kDa)	M.W.(kDa)	M.W.(kDa)
	200→	200→	200→
	45→	45→	45→
	22→ 14.4→ 6.5→	22→ 14.4→ 6.5→	=
		350000	22→
			14.4→ 6.5→

<Condition>

Sample : Protein Markers(10x)(# 29458-24), 3µl

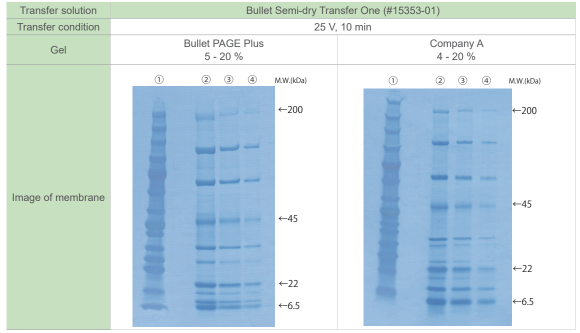
Gel Staining : CBB Stain One (#04543) Voltage Constant : Bullet PAGE Plus: 400V

Company A and Company B: 200V

<sup>\*</sup>Please use 25 mM Tris and 192 mM glycine buffer for nucleic acid electrophoresis

# **Blotting performance: Comparison of protein transfer efficiency**

This product shows a transfer efficiency equivalent to Company A's product.



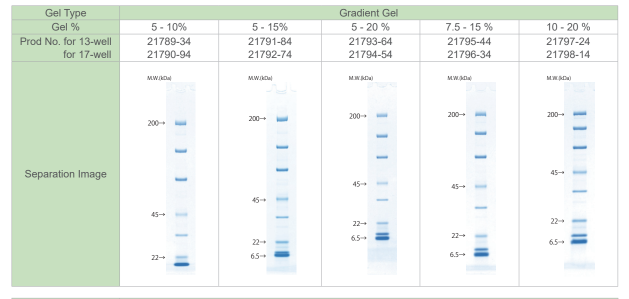
<Condition>

Sample : ① Protein Ladder One Plus, Triple-color for SDS-PAGE (#19593-25)

Dilute Protein Marker (10x) (#29458-24) to achieve @ 3x, @ 1x, and @ 1/3x, 2  $\mu$ L of each.

Staining : Stain the PVDF membrane with CBB Stain One (Ready To Use) (#04543)

## **Gel types**



Gel Type	Single-percentage gel					
Gel %	7.5%	10%	12.5%	15%		
Prod No. for 13-well	21799-04	21801-44	21807-84	21853-74		
for 17-well	21800-54	21806-94	21811-14	21854-64		
Separation Image	M.W.(kDa) 200→ 45→	M.W.β.Da)  200→  45→  65→	M.W.(kDa)  200→  45→  22→  6.5→	M.W.(kDa)  200→  45→  22→  6.5→		

# **Specification**

Glass Plate Size : W100 mm x H80 mm x T3.2 mm (made of glass)

Gel Size : W80 mm x H60 mm x T1.0 mm

Sample Well Configuration / Maximum Load Volume : 13-well / 40 µl, 17-well / 28 µl

Shelf Life : 9 months

# **Ordering Information**

Product name	Storage	Product number	PKG size
Bullet PAGE Plus Precast Gel, 5-10%, 13wells	R	21789-34	10 Sheets
Bullet PAGE Plus Precast Gel, 5-10%, 17wells	R	21790-94	10 Sheets
Bullet PAGE Plus Precast Gel, 5-15%, 13wells	R	21791-84	10 Sheets
Bullet PAGE Plus Precast Gel, 5-15%, 17wells	R	21792-74	10 Sheets
Bullet PAGE Plus Precast Gel, 5-20%, 13wells	R	21793-64	10 Sheets
Bullet PAGE Plus Precast Gel, 5-20%, 17wells	R	21794-54	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5-15%, 13wells	R	21795-44	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5-15%, 17wells	R	21796-34	10 Sheets
Bullet PAGE Plus Precast Gel, 10-20%, 13wells	R	21797-24	10 Sheets
Bullet PAGE Plus Precast Gel, 10-20%, 17wells	R	21798-14	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5%, 13wells	R	21799-04	10 Sheets
Bullet PAGE Plus Precast Gel, 7.5%, 17wells	R	21800-54	10 Sheets
Bullet PAGE Plus Precast Gel, 10%, 13wells	R	21801-44	10 Sheets
Bullet PAGE Plus Precast Gel, 10%, 17wells	R	21806-94	10 Sheets
Bullet PAGE Plus Precast Gel, 12.5%, 13wells	R	21807-84	10 Sheets
Bullet PAGE Plus Precast Gel, 12.5%, 17wells	R	21811-14	10 Sheets
Bullet PAGE Plus Precast Gel, 15%, 13wells	R	21853-74	10 Sheets
Bullet PAGE Plus Precast Gel, 15%, 17wells	R	21854-64	10 Sheets

# [General-use Precast Gel]

# **Extra PAGE One Precast Gel**

- » Superior resolution
- » Sharp bands
- » High transfer efficiency
- » Compatible with conventional Laemmli sample and running buffers

## **Performance Comparison**

Extra PAGE One Precast Gel offers higher resolution than other gels. Especially 5-20% gradient gel and 12.5% gel provide improved resolution for low molecular weight.

Gel		Gradi	ent Gel		Single-percentage Gel			
Product Name	Extra PAGE One	Company A	Extra PAGE One	Company B	Extra PAGE One	Company A		
Gel%	5 - 20%	4 - 20%	5 - 20%	5 - 20%	12.5%	12.5%		
Product No.	11955-54	-	11955-54	-	11965-24	-		
Migration patterns	M.W.(KDa)  200→  45→  22→ 6.5→	M.W.(KDa) 200→ 45→ 14→ 6.5→	M.W.(KDa) 200→ 45→ 22→ 6.5+14→					

<Condition>

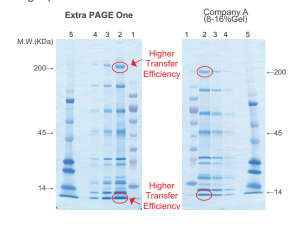
Sample : Protein Markers (#29458-24) 5ul. Staining : CBB Stain One (#04543)

Electrophoresis condition: Extra PAGE One Precast Gel: 300 V, approx. 50 min.

Company A: followed the manufacture's instructions

### **Comparison of Transfer Efficiency**

Extra PAGE One Precast Gel offers higher transfer efficiency than Company A for both high and low molecular weight proteins.



- <Sample>
- 1. Protein ladder One (Triple-color) (Product No. 09547)
- 2. 2 ul of Protein Markers (3x)3. 2 ul of Protein Markers (1x)
- 4. 2 ul of Protein Markers (1/3)
- 5. 10 ul of Bio-Rad Precision Plus Protein™, Kaleidoscope™
- <Transfer>

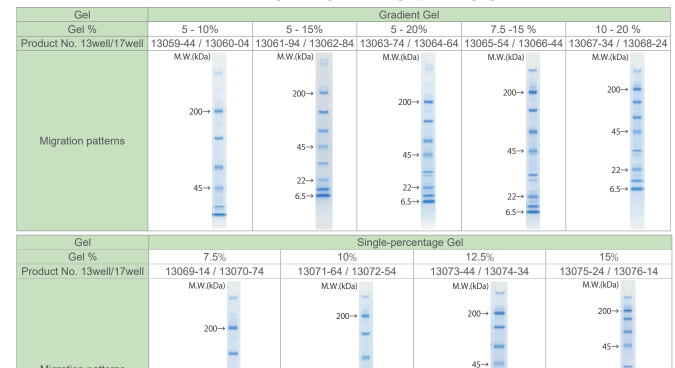
Semi-dry with PVDF membrane and Blotting Buffer Solution for Western Blotting (Product No.: 30650-31) (Transfer time 20 min with 10V constant current)

<Staining>

CBB Stain One (Product No.: 04543)

#### **Product Lines**

Extra PAGE One Precast Gel is available in gradient gels and single-percentage gels.



Note) Back ground may remain slightly when using Nacalai's Rapid Stain CBB Kit (Product No.: 30035-14) or other products. To reduce back ground, wash the gel with 10% acetic acid for 20 minutes.

45→

#### **Specifications**

Migration patterns

Cassette Size : 100 mm(W) x 100 mm(H) x 3.2 mm(T) (made of glass)

Gel Size : 80 mm(W) x 80 mm(H) x 1.0 mm(T)

45→

Well : 13 wells, 17 wells

Shelf Life Expire date: 9 months

Note) Use electrophoresis tank for plate size 100mm(W) x 100mm(H) x 3.2mm(T).

## **Ordering Information**

Product name	Storage	Product number	PKG size
Extra PAGE One Precast Gel, 5-10%, 13wells	R	13059-44	10 Sheets
Extra PAGE One Precast Gel, 5-10%, 17wells	R	13060-04	10 Sheets
Extra PAGE One Precast Gel, 5-15%, 13wells	R	13061-94	10 Sheets
Extra PAGE One Precast Gel, 5-15%, 17wells	R	13062-84	10 Sheets
Extra PAGE One Precast Gel, 5-20%, 13wells	R	13063-74	10 Sheets
Extra PAGE One Precast Gel, 5-20%, 17wells	R	13064-64	10 Sheets
Extra PAGE One Precast Gel, 7.5-15%, 13wells	R	13065-54	10 Sheets
Extra PAGE One Precast Gel, 7.5-15%, 17wells	R	13066-44	10 Sheets
Extra PAGE One Precast Gel, 10-20%, 13wells	R	13067-34	10 Sheets
Extra PAGE One Precast Gel, 10-20%, 17wells	R	13068-24	10 Sheets
Extra PAGE One Precast Gel, 7.5%, 13wells	R	13069-14	10 Sheets
Extra PAGE One Precast Gel, 7.5%, 17wells	R	13070-74	10 Sheets
Extra PAGE One Precast Gel, 10%, 13wells	R	13071-64	10 Sheets
Extra PAGE One Precast Gel, 10%, 17wells	R	13072-54	10 Sheets
Extra PAGE One Precast Gel, 12.5%, 13wells	R	13073-44	10 Sheets
Extra PAGE One Precast Gel, 12.5%, 17wells	R	13074-34	10 Sheets
Extra PAGE One Precast Gel, 15%, 13wells	R	13075-24	10 Sheets
Extra PAGE One Precast Gel, 15%, 17wells	R	13076-14	10 Sheets

22∹

 $6.5 \rightarrow$ 

22-

[Apparatus for Precast Gel]

# **Electrophoresis Tank for Precast Gel**

**Product Name** 

### **Specification**

Size : 143W x 84D x 140H (mm)

Required Buffer Volume: 550 ml

### **Ordering Information**

WEP-MN Vertical Electrophoresis Tank



1 Set

[Storage] RT = Room Temperature, R = Refrigerator, F = Freezer

WEP-MN

RT

Note: Since the adapter for fitting Bullet PAGE Plus includes this product, please inform a distributor if you plan to use this product for Extra PAGE One.

# [Gel Buffer for Hand-made Gel]

# WIDE RANGE Gel Preparation Buffer (4x) for PAGE

Gradient gels offer a much wider separation range of proteins than single percentage gels. However, casting gradient gels is more difficult and labor intensive. WIDE RANGE Gel Preparation Buffer offers a gradient gel-like separation on a single percentage gel by simply mixing it with acrylamide/ bisacrylamide gel casting solution. The gel can be used with the common sample buffers and running buffers. It is also suitable for standard staining methods including CBB and silver staining.

### » Simple casting procedure

WIDE RANGE Gel Preparation buffer is a 4x concentrated neutral pH buffer. It can be used for preparation of both stacking gel and separation gel by replacing the Tris-HCl buffer in Laemmli buffer system.



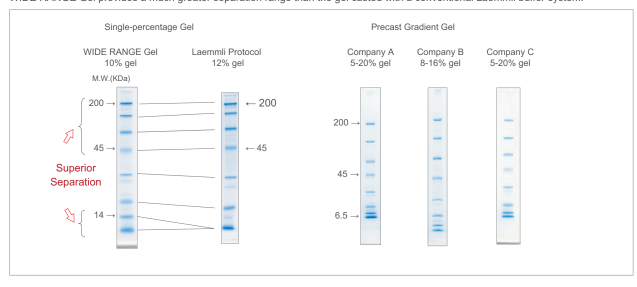
#### » Improved stability and strength

The incresed tensile strength allows easy handling even a low percentage gel. The neutral pH buffer improves the stability of gel resulting in a longer shelf life than the gel with Laemmli buffer system.

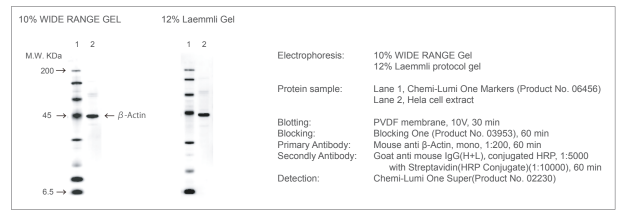


#### » A wide separation range

WIDE RANGE Gel provides a much greater separation range than the gel casted with a conventional Laemmli buffer system.



### **Applicable for Western Blotting**



#### **Ordering Information**

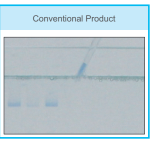
Product Name	Storage	Product No.	PKG Size
WIDE RANGE Gel Preparation Buffer (4x) for PAGE	R	07831-94	250 ml

# [Stacking Gel Buffer for Hand-made Gel]

# Stacking Gel Buffer Solution (4x) with Blue Color

# » Easy to see well locations due to coloring stacking gel







Wells are clearly confirmed on stacking gel prepared with Stacking Gel Buffer Solution with Dye.

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Stacking Gel Buffer Solution (4x) with Dye for SDS-PAGE	R	09268-34	100 ml

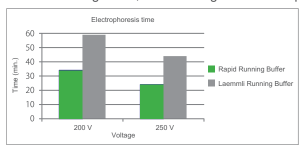
# [Fast running buffer for Hand-made Gel]

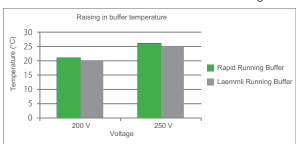
# **Rapid Running Buffer Solution**

- » Approximately 25 min. electrophoresis time with mini-gel at 250 V
- » Just replace the Laemmli running buffer with this product
- » High protein transfer efficiency to western blotting membrane

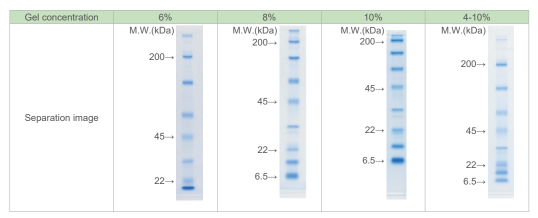
# Comparison of electrophoresis duration and buffer temperature rise

Running proteins with this product shortens the electrophoresis duration to about 60% compared to Laemmli running buffer, and its rising in buffer temperature is the almost same as Laemmli running buffer's.





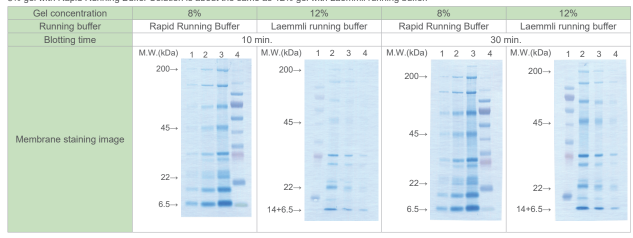
## Separation patterns of Laemmli gel with Rapid Running Buffer



## Comparison of protein transfer efficiency to western blotting membrane

Due to lower gel concentration when Rapid Running Buffer Solution is used<sup>\*1</sup>, its protein transfer efficiency to a membrane is higher than the Laemmli running buffer's.

"By casting a gel with 4% lower gel concentration than usual, its separation patterns can be made similar to the original's, e.g. separation patterns of 8% gel with Rapid Running Buffer Solution is about the same as 12% gel with Laemmli running buffer.



## **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Rapid Running Buffer Solution (20x) for SDS-PAGE	RT	12981-74	250 ml

# **Polyacrylamide Gel Casting Reagents**

# **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Acrylamides (monomer)			
And and deside (conserve) Durity 000/	RT	00809-14	100 g
Acrylamide (monomer), Purity, 99%	KI	00809-85	500 g
		06114-24	100 g
Acrylamide (monomer), Purity, 99%, Nuclease and Protease tested	RT	06114-95	500 g
	R R R R R R R	06114-11	1 kg
Crosslinking Agents			
N,N'-Methylenebisacrylamide, [BIS]	R	22402-02	25 g
N,N'-Methylenebisacrylamide, Purity, 99%, Nuclease and Protease tested	R	22407-52	25 g
Acrylamide/Bis Mixed Solutions			
30(w/v)%-Acrylamide/Bis Mixed Solution (37.5:1)	R	06144-05	500 ml
30(w/v)%-Acrylamide/Bis Mixed Solution (29:1)	R	06141-35	500 ml
40(w/v)%-Acrylamide/Bis Mixed Solution (37.5:1)	R	06121-95	500 ml
40(w/v)%-Acrylamide/Bis Mixed Solution (29:1)	R	06119-45	500 ml
Polymerization Initiators			
N,N,N',N'-Tetramethylethylenediamine [TEMED]	RT	33401-72	25 g
N,N,N,N-Tetrametryletriyleriediamine [TEMED]	KI	33401-14	100 g
Polymerization Promotors			
Ammonium Porovodiculfoto [ADS]	R	02627-21	1 g
Ammonium Peroxodisulfate [APS]	K	02627-34	10 g
10 (w/v)%-Ammonium Peroxodisulfate Solution	F	02634-34	10 ml
Gel Buffer Solutions			
Separating Gel Buffer Solution (4x) for SDS-PAGE, pH8.8 Filtrated by 0.45 µm filter Components : 1.5M-Tris-HCl, 0.4 (w/v)%-SDS	RT	30651-05	500 ml
Stacking Gel Buffer Solution (4x) with Dye for SDS-PAGE, pH6.8 Filtrated by 0.45 µm filter Components : 0.5M-Tris-HCl, 0.4(w/v)%-SDS, Dye	R	09268-34	100 ml
Stacking Gel Buffer Solution (4x) for SDS-PAGE, pH6.8 Filtrated by 0.45 µm filter	В	09267-44	100 ml
Components: 0.5M-Tris-HCl, 0.4 (w/v)%-SDS	R	32158-25	500 ml

# **Running Buffers**

# **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Pre-mixed Buffers			
Running Buffer Solution (10x) for SDS-PAGE, Tris-Glycine,	RT	30329-61	1 L
Filtrated by 0.45 µm filter Components: 0.25 mol/l-Tris, 1.92 mol/l-glycine, 10 g/l-SDS	KI	30329-74	5 L
Running Buffer Solution (10x) for PAGE,Tris-Glycine,	RT	30340-01	1 L
Filtrated by 0.45 µm filter Components: 0.25 mol/l-Tris, 1.92 mol/l-glycine	IXI	300-0-31	
Buffer Adjusting Reagents			
Tris(hydroxymethyl)aminomethane, Purity, 99%	RT	35410-34	100 g
Tris/hydray/mathyl\aminamathana		35434-76	100 g
Tris(hydroxymethyl)aminomethane,	RT	35434-05	500 g
Purity, 99.9%, Nuclease and Protease tested		30329-61 30329-74 30340-91 35410-34 35434-76	1 kg
Glycine	RT	17128-14	100 g
Chains Nucleans and Dratages tested	DT	17141-24	100 g
Glycine, Nuclease and Protease tested	RT	30329-61 30329-74 30340-91 35410-34 35434-76 35434-05 35434-21 17128-14 17141-24 17141-95 34713-62 34713-04 02437-24 31607-52 31607-94	500 g
Tribing (NI (Trib //bodessorms Abod) and Abod short a	DT	34713-62	25 g
Tricine {N-[Tris(hydroxymethyl)methyl]glycine}	RT	34713-04	100 g
Tricine {N-[Tris(hydroxymethyl)methyl]glycine} Nuclease and Protease tested	RT	02437-24	100 g
		31607-52	25 g
Sodium Lauryl Sulfate [Sodium Dodecyl Sulfate;SDS] Purity, 99%	RT	31607-94	100 g
		31607-65	500 g
Onditional actual College annual and Conditional Development College CDC1			

Sodium Lauryl Sulfate granular [Sodium Dodecyl Sulfate;SDS] Purity, 99%, Solids (granular)





	02873-62	25 g
RT	02873-04	100 g
	02873-75	500 g

10%-SDS Solution [10%-Sodium Lauryl Sulfate Solution]	RT	30562-04	100 ml

# Sample Buffer Solution for SDS-PAGE (6x)

- » Suitable for low concentration protein sample adjustment
- » No precipitation in the refrigerator
- » Two types of reagents (with and without reducing agent)

## **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Sample Buffers			
Sample Buffer Solution with Reducing Reagent (6x) for SDS-PAGE pH6.8 Filtrated by 0.45 µm filter, Components: 0.375M-Tris-HCl, 0.03(w/v)%-BPB, glycerin, anion surface acting agent and reducing agent	R	09499-14	5 ml
Sample Buffer Solution without Reducing Reagent (6x) for SDS-PAGE pH6.8 Filtrated by 0.45 µm filter, Components: 0.375M-Tris-HCI, 0.03(w/v)%-BPB, glycerin and anion surface acting agent	R	09500-64	5 ml
Sample Buffer Solution with 2-ME (2x) for SDS-PAGE pH6.8 Filtrated by 0.45 µm filter, Components: 0.125M-Tris-HCl, 4(w/v)%-SDS, 20(v/v)%-glycerin, 0.01(w/v)%-BPB, 10(v/v)%-2-ME	R	30566-22	25 ml
Sample Buffer Solution without 2-ME (2x) for SDS-PAGE pH6.8 Filtrated by 0.45 µm filter, Components: 0.125M-Tris-HCl, 4(w/v)%-SDS, 20(v/v)%-glycerin, 0.01(w/v)%-BPB	R	30567-12	25 ml
Reducing Agent			
2-Mercaptoethanol	RT	21418-42 21418-84 21418-55	25 g 100 g 500 g
Dithiothreitol	R	14112-36 14112-81 14112-94 14112-52	100 mg 1 g 5 g 25 g
1mol/l-Dithiothreitol Solution	F	14130-41	1 ml
Tris (2-carboxyethyl) phosphine Hydrochloride (TCEP)	R	07277-61	1 g
Tracking Dyes			
Bromophenol Blue	RT	05808-61 05808-32	1 g 25 g
Others			
Glycerol Nuclease and Protease tested	RT	17045-94 17045-65	100 ml 500 ml

# **Molecular Weight Markers**

# Protein Ladder One Plus

- » Excellent visibility with triple color
- » Sharp bands
- » Consists of 14 bands
- » Low lot-to-lot variation



Extra PAGE One Precast GeI, 5-20%, 13wells(#13063-74) Load volume 5  $\mu$ L

#### Notes

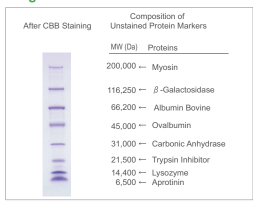
- There is a possibility of different molecular weight result depending on gel type (concentration, gradient, buffer, etc.). It may not be suitable for accurate molecular weight measurement.
- The product is ready-to-use type. Sample preparation, such as heating and reduction, is not required.

otein
kinds
position
15% (v/v)
2% (w/v)
0.2 mM
3.6 M
20 mM

# Unstained Protein Markers (10x)

» Contains 8 kinds of protein (M.W. 6,500 - 200,000 Da)

#### **Image**



#### Composition

50(v/v)% Glycerol
0.3 M NaCl
0.1 M DTT, 2 mM EDTA • 2Na
3 mM NaN<sub>3</sub>
10 mM Tris-HCl (pH 7.0)

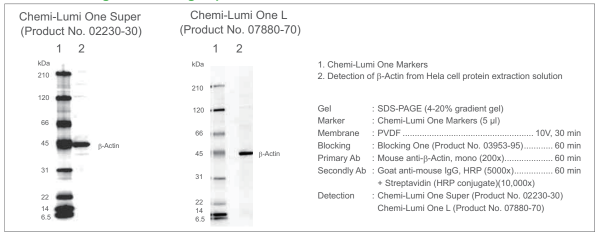
## Chemi-Lumi One Markers Kit

Chemi-Lumi One Markers Kit consists of biotinylated proteins and HRP-conjugated streptavidin. Each band can be visualized on a western blotting by the same chemiluminescent reagents for the target protein.

- » Contains 8 biotinylated proteins as molecular weight markers (M.W. 6,500 200,000 Da)
- » Includes HRP-conjugated strepavidin to detect biotinylated proteins



### Western Blotting for Detecting of β-Actin



#### Components

Chemi-Lumi One Markers consists of 8 biotinylated proteins, 50  $\mu$ l: 1 tube Streptavidin (HRP conjugate), 250  $\mu$ l: 1 tube

Note: The molecular weight of Chemi-Lumi One Markers may slightly differ from unmodified proteins because of biotinylation.

#### **Ordering Information**

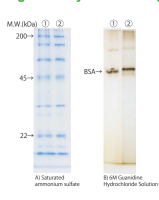
Product Name	Storage	Product No.	PKG Size
Protein Ladder One Plus, Triple-color for SDS-PAGE	F	19593-25	500 μL
Protein Markers(M.W. 6,500~200,000)(10x) for SDS-PAGE	F	29458-24	200 μL
Chemi-Lumi One Markers Kit	F	06456-70	1 KIT

# **PAGE Clean Up Kit**

This product is designed for sample preparation in two-dimensional electrophoresis and SDS-PAGE. This product is a combination of acetone, trichloroacetic acid, and coprecipitate optimized for protein precipitation. By resuspending the precipitated proteins in sample lysate according to the intended use, it is possible to concentrate the sample protein and remove interfering substances in the buffer.

- » Suitable for concentrating protein samples and removing analytical interferences
- » Precipitation is possible even in the subsistence of high concentrations of urea, guanidine, and SDS
- » Easier than dialysis and small volume processing at the  $\mu l$  level

#### Removing electrolyte interfering substances



<Conditions>

[Sample]

(A) 10x dilution of Protein Markers(10x) #29458-24 (standard concentration)

(B) BSA (100ng: non-reducing)

[Precipitation treatment]

 $\ensuremath{\textcircled{1}}$  without this product

② with this product

[Electrophoresis]

SDS-PAGE (12% gel)

[Staining]

(A) CBB Stain One #04543 (CBB staining)

(B) Sil-Best Stain One #06865-81 (silver staining)

When a sample containing components that affect the electrophoretic image was treated with this product, the electrophoretic image was improved.

Note) Yield may vary depending on the composition of the sample solution, protein concentration, and other factors. Please use after preliminary examination

## **Ordering Information**

Product Name	Storage	Product No.	PKG Size
PAGE Clean Up Kit	R	06441-50	1 KIT

# **CBB Stain Solution**

CBB staining, Coomassie Briliant Blue staining, is a popular method for detectiong proteins in polyacrylamide gel. We offer four types of CBB staining solution.

## **Selection of CBB staining solution**

	Bullet CBB Stain Lite (#21964)	CBB Stain One Super (#11642)	CBB Stain One (#04543)	Rapid Stain CBB Kit (#30035)
Dye Type	CBB-G250	CBB-R250	CBB-G250	CBB-R250
Component	Single bottle (Ready-to-use) w/o acetic acid and methanol	Single bottle (Ready-to-use) w/o acetic acid and methanol	Single bottle (Ready-to-use) w/o acetic acid and methanol	Two bottles w/ acetic acid and w/o methanol
Gel rinsing	Unnecessary	Required 3 times for 5min.	Required 3 times for 5min.	Unnecessary
Staining Period	15 min.	30 min.	60 min.	20 min.
Destaining Period	Unnecessary	More than 1 hr.	More than 1 hr.	More than 1 hr.
Sensitivity	Sensitivity Up to tens of ng proteins			

# Bullet CBB Stain Lite

#### » In one step, complete staining in 15 minutes

No need for gel pre-staining washing

Clear staining image without the need for a destaining step

» Free from strong irritants like acetic acid and methanol

# Observable staining image in the staining solution



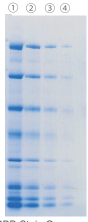
<Conditions>

Sample : E. coli was prepared in Sample Buffer Solution with 2-ME(2x) for SDS-PAGE #30566-22

SDS-PAGE : Bullet PAGE Plus Precast Gel, 5-20%, 17wells (#21794-54) 400 V (constant voltage), for 10 minutes

CBB stain : Bullet CBB Stain Lite (#21964) at room temperature for 15 minutes

- CBB Stain One
- » Single bottle
- » CBB G-250 type
- » Free from acetic acid and methanol



CBB Stain One (#04543)

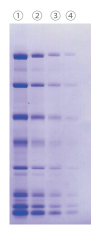
Staining: 60 mins. Destaining: Over night

<Conditions>

Sample SDS-PAGE Washing: 3 times x 5 mins.

CBB Stain One Super

- » Single bottle
- » CBB R-250 type
- » Free from acetic acid and methanol



CBB Stain One Super

(#11642)

Washing: 3 times x 5 mins.

Staining: 30 mins. Destaining: Over night

: Diluted Protein Markers (10x)(#29458-24) into ① 3x, ② 1x, ③ 1/3x, and ④ 1/10x, then added 2  $\mu L$ : Extra PAGE One Precast Gel, 5-20%, 13wells (#13063-74) 400V (constant voltage), for 30 mins.

## **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Bullet CBB Stain Lite	R	21964-24	50 ml
	K	21964-95	500 ml
CBB Stain One Super (Ready-to-use)	RT	11642-31	1 L
CBB Stain One (Ready-to-use)	DT	04543-51	1 L
	RT	04543-64	5 L
Rapid Stain CBB Kit (Coomassie R-250)	RT	30035-14	1 set (for 2 L)

#### **Related Products**

Product Name	Storage	Product No.	PKG Size
Coomassie Brilliant Blue G-250	RT	09409-42	25 g
Coomassie Brilliant Blue R-250	RT	09408-52	25 g
Ponceau S	RT	28322-72	25 g

# Silver Staining Kit

Silver staining method is a high sensitive method for detecting nucleic acids and proteins in polyacrylamide gel. We offer two types of silver staining kits, each having unique features for your experimental needs.

## **Selection of Silver Staining Kit**

	Sil-Best Stain One	Sil-Best Stain-Neo
2-Dimensions	Excellent	Poor
SDS-PAGE	Good	Good
Nucleic Acid	Poor	Good
Step	12	6
Staining time	80 min.	60 min.

# **Comparison of Each Procedure**



### **Comparison of Each Staining Image**



## Sil-Best Stain One

Sil-Best Stain One is based on the silver staining method for protein detection. Its composition does not contain glutaraldehyde affects a result of mass spectrography.

- » High sensitivity and low background
- » No glutaraldehyde

# More visible protein spot numbers than Competitors' silver staining kit

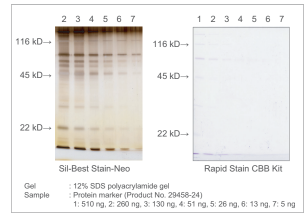


## Sil-Best Stain-Neo

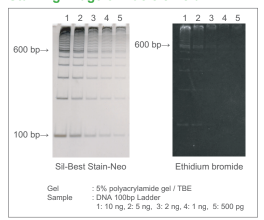
Sil-Best Stain-Neo is a highly sensitive method for detecting nucleic acids and proteins in polyacrylamide gel. It is 50-100 fold more sensitive than coomassie brilliant blue and ethidium bromide.

## » Only 6 steps within up to 1 hour

# **Staining Image of Protein**



## Staining Image of Nucleic Acid



## **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Sil-Best Stain One	R	06865-81	1 set
Sil-Best Stain-Neo for Protein and Nucleic Acid/PAGE	R	05773-11	1 set

#### **Related Products**

Product Name	Storage	Product No.	PKG Size
Dispotray-SS	RT	19386-74	40 pieces
Dispotray-S (for minigel staining)	RT	16526-82	25 pieces
Dispotray-M	RT	16551-84	20 pieces

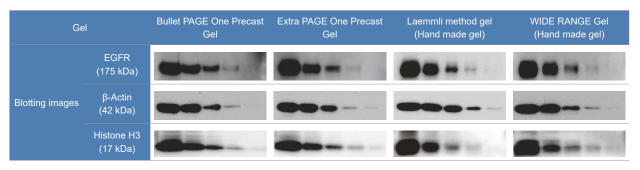
## [Ultra-fast Blotting Buffer]

## **Bullet Semi-dry Transfer One**

- » Transfer can be completed at high speed
- » Efficient transfer a wide range of molecular weights
- » No need for gel equilibration process
- » Ready-to-use

#### **Application 1**

After electrophoresis on various gels, high speed transfer was performed by using this product. EGFR, β-Actin, and Histone H3 were detected through western blotting. Proteins ranging from low molecular weight (Histone H3) to high molecular weight (EGFR) have been successfully transferred by using this product and detected quantitatively.



(Conditions)

Samples : HeLa cell suspension (2 times dilution starting from 10 ug)

SDS-PAGE : Bullet PAGE One Precast Gel, 5-15%, 17wells (#13080-44) 400 V

Extra PAGE One Precast Gel, 5-15%, 17wells (#13062-84) 300 V Laemmli method gel (EGFR; 6%, β-Actin; 12%, Histone H3; 15%) 200 V WIDE RANGE Gel\*(EGFR; 6%, β-Actin; 10%, Histone H3; 12%) 200 V \*Casted with WIDE RANGE Gel Preparation Buffer(4x) for PAGE (#07831-94)

: Bullet Semi-dry Transfer One (#15353-01) 25 V 10 min PVDF (poresize; 0.45 µm)

Transfer

Blocking : Bullet Blocking One for Western Blotting (#13779) RT 5 min

: EGF Receptor (D38B1) XP Rabbit mAb (CST #4267) 2,000x diluted Primary antibody (EGFR)

by Signal Enhancer HIKARI for Western Blotting and ELISA (#02270-81) Solution ART 60 min

: β-Actin (C4) (Santa Cruz #sc-47778) 5,000x diluted (β-Actin)

by Bullet ImmunoReaction Buffer (#18439) RT 30min

(Histone H3): Histone H3 (D1H2) XP Rabbit mAb (CST #4499) 2,000x diluted

by Bullet Blocking One for Western Blotting (#13779) RT 60 min

Secondary antibody (EGFR) : Goat anti-Rabbit IgG (H+L) Secondary Antibody [HRP](Pre-adsorbed) (Novus #NB7187) 50,000x diluted

by Signal Enhancer HIKARI for Western Blotting and ELISA (#02270-81) Solution B RT 60 min

(β-Actin) : Goat anti-Rat, Mouse IgG (H+L) Secondary Antibody [HRP](Pre-adsorbed) (Novus #NB7574) 80,000x diluted

by Bullet ImmunoReaction Buffer (#18439-85) RT 30min

(Histone H3): Goat anti-Rabbit IgG (H+L) Secondary Antibody [HRP](Pre-adsorbed) (Novus #NB7187) 30,000x diluted

by Bullet Blocking One for Western Blotting (#13779) RT 60 min

Detection ChemiDoc Touch MP 3x3 binning (Bio-rad) (AUTO mode)

> (EGFR) : Chemi-Lumi One Super (#02230) (β-Actin) : Chemi-Lumi One L (#07880) (Histone H3) : Chemi-Lumi One Super(#02230)

#### Comparison with conventional transfer method

After electrophoresis on the Bullet PAGE Plus Precast Gel, rapid transfer (semi-dry) and conventional transfer (semi-dry and tank) were processed by this product, then detected GAPDH through western blotting. This product enables rapid transfer in just 10 minutes, achieving detection results comparable to the traditional method.

	Conditions	Blotting images
High-speed transfer	Bullet Semi-dry Transfer One(#15353-01) 25 V 10 mins Semi-dry transfer (no need for equilibration)	
Conventional	Semi-dry Blotting Buffer Solution for Western Blotting 10 V for 45 min Semi-dry transfer (15 mins. equilibration performed)	
transfer	Towbin method composition transfer buffer* 100 V tank transfer for 60 mins (15 mins equilibration performed) *25 mM Tris, 192 mM Glycine, 20%MeOH	

(Conditions)

Samples : HeLa cell suspension (2 times dilution started from 4 ug)

Transfer : Conditions listed above

Primary antibody : GAPDH Antibody (Novus #NB300-322) 10,000x diluted

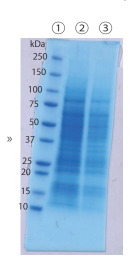
by Bullet ImmunoReaction Buffer (#18439) RT 30 min

Secondary antibody : Goat anti-Rabbit IgG (H+L) Secondary Antibody [HRP](Pre-adsorbed) (Novus #NB7187) 200,000x diluted

Detection : Exposure time 3 mins

#### **Application 2**

After running on Mini-PROTEAN TGX Gels, high-speed transfer was performed by this product and transfer was confirmed by CBB staining.



[Conditions]

Sample:

① Precision Plus Protein™ All Blue Prestained Protein Standards (Bio-rad #1610373)

2 293T Cell Suspension

3 293T Cell Suspension (2x diluted by 2)

SDS-PAGE : 4-20% Mini-PROTEAN TGX Gels (Bio-rad #4561095)
Transfer : Bullet Semi-dry Transfer One (#15353-01) 25 V for 10 min

CBB Staining: Bullet CBB Stain One(Ready To Use) (#13542)

Data provided by a University researcher.

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Bullet Semi-dry Transfer One	RT	15353-01	1 L

#### [Blocking Buffer]

## **Blocking One and Blocking One P**

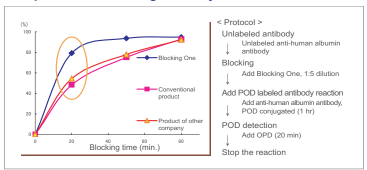
Blocking is indispensable in immunoassays to block non-specific binding reactions. As Blocking One contains high molecular weight compounds, casein and bovine serum protein, it is superior to conventional blocking solutions.

Blocking One-P is an exclusive blocking solution, free of phosphate group and endogenous phosphatase for phosphorylated protein detection. The performance is superior compared with conventional blocking solutions such as 1% BSA.

The preservative in our blocking solutions do not affect the enzyme activity of peroxidase (POD) or alkaline phosphatase (ALP). Only simple refrigerator storage is necessary, even after opening the bottle.

- » In many assays, a reduction of incubation time for blocking can be achieved
- » Simple storage in a refrigerator even after opening the bottle

#### **Comparison of Blocking Efficincy**

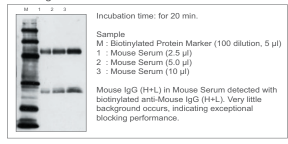


The figure left shows the relationship between the reaction time and the effect of blocking in microplate assay.

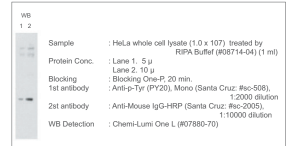
In comparison with other products, Blocking One offers the strongest blocking efficiency along with faster blocking treatment time.

#### **Western Blotting**

#### Blocking One



#### Blocking One-P



#### **Comparison with Conventional Blocking Agents**

	Composition	Treatment Time	Blocking Efficacy	Phospho-specific antibody applications
Bullet Blocking One	- Amphiphilic compound - High molecular weight compounds - BSA - Casein - refer to p75	5 min.	+++	+
Blocking One	- High molecular weight compounds - BSA - Casein	20-30 min.	+++	+
Blocking One-P	- High molecular weight compounds - BSA	20-30 min.	+++	+++
Skim milk	- Casein	1 hour	+	-
1% BSA	- BSA	1 hour	+	++

#### **Ordering Information**

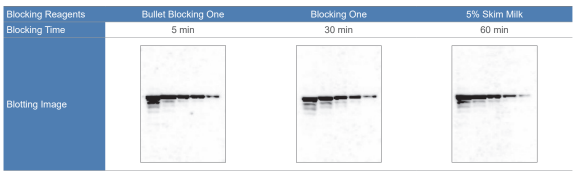
Product Name	Storage	Product No.	PKG Size
Blocking One	R	03953-95	500 ml
Blocking One-P	R	05999-84	200 ml

#### [Ultra-fast Blocking Buffer]

## **Bullet Blocking One for Western Blotting**

- » Fast blocking in 5 mins
- » Ready-to-use
- » Simple storage in a refrigerator even after opening the bottle

#### Comparison Data with Blocking One (our standard blocking reagent) and 5% skim milk



(Conditions)

Sample : 20 µg of HeLa cell extract, 5 serial two-fold dilution series

SDS-PAGE : Bullet PAGE One 5-15% (Product No. 13080) with SDS Running Buffer (Product No. 30329) at 400 V for 10 min.

Blotting : Semi-dry Blotting Buffer Solution (Product No. 30650) at 10 V for 30 min

Washing : 0.1% t-TBS (Product No. 12750)
Blocking : Refer to the figure above

1st antibody : Anti-Vimentin (C-20) (Rabbit) (Product No. SC-7557-R) diluted 1:2,000, 1 hr. at RT 2nd antibody : Anti-Rabbit IgG-HRP (Product No. SC-2004) diluted 1:100,000, 1 hr. at RT

Detection : Chemi-Lumi One (Product No. 11644)

Detector : LAS-3000 (High mode), 15 min. exposure time

#### Comparison Data: Blocking efficiency in 5 minutes

The original Blocking One, the competitors' ready-to-use blocking reagents and the conventional blocking reagents did not show enough blocking efficiency in 5 mins, while Bullet Blocking One performed well.

5 min. Blocking Time	Blocking Reagents	Manufacturers' Recomm	nended Blocking Time
• • • • •	Bullet Blocking One		
	Blocking One	• • • • • •	30 min.
OF THE STATE OF TH	Company A	• • • • • •	60 min.
o o Bill	Company B		30 min.
	Company C		60 min.
	5% Skim Milk in 0.05 Tween <sup>®</sup> 20-TBS	• • • • • •	60 min.
000	3% BSA in 0.05% Tween <sup>®</sup> 20-TBS	• • • • • •	60 min.

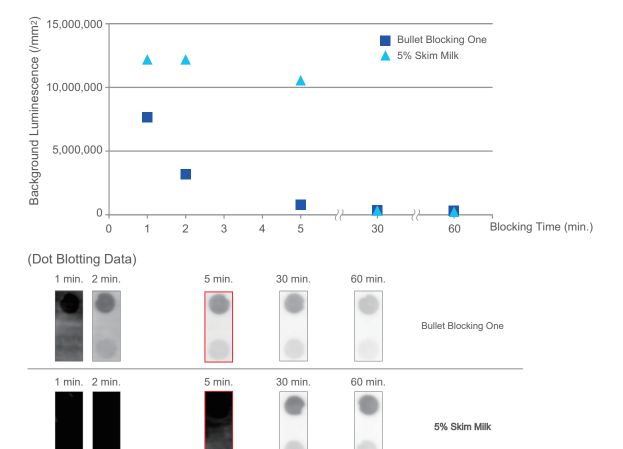
#### (Conditions)

PVDF membranes dot-blotted with mouse serum were washed with TBS. Blocking was performed using each reagent above. Anti-mouse IgG (Product No.SC-2005) (1:5,000 in 0.01% t-TBS) was applied, and the membranes were washed with 0.05% t-TBS. After reaction with Chemi-Lumi One Super (Product No. 02230), detection was performed using LAS-3000 (High mode) with 90 sec. exposure time.

#### Comparison of Blocking Time and Efficiency with 5% Skim Milk

Bullet Blocking One performed better than 5% skim milk in blocking time and efficiency.

(Analysis of Background Noise on Dot Blotting Data)



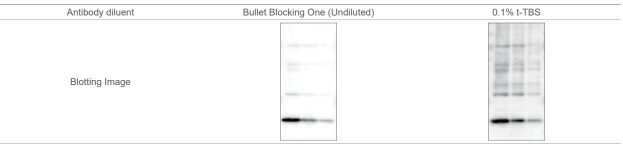
#### (Conditions)

PVDF membranes dot-blotted with mouse serum were washed with TBS. Blocking was performed using each reagent above. Anti-mouse IgG (Product No. SC-2005) (1:5,000 in 0.01% t-TBS) was applied, and the membranes were washed with 0.05% t-TBS. After reaction with Chemi-Lumi One Super (Product No. 02230), detection was performed using LAS-3000 (High mode) with 90 sec. exposure time. Data analysis was done with Multi Gauge.

Western Blotting

#### Use as antibody diluent

Diluting antibody with Bullet Blocking One resulted in less non-specific binding compared to using t-TBS as diluent.



(Conditions)

Sample : 10  $\mu g$  of HeLa cell extraction, 3 serial two-fold dilution series

: Bullet PAGE One 5-15% (Product No. 13080) with SDS Running Buffer (Product No. 30329) at 400 V for 12 min. SDS-PAGE

: Semi-dry Blotting Buffer Solution (Product No. 30650) at 10 V for 30 min. Blotting

Washing : 0.1% t-TBS (Product No. 12750) : Bullet Blocking One, 5 min. Blocking

: Anti-Cox4 (D-20) (Goat) (Product No. SC-69359) diluted 1:500, 1 hr. at RT 1st antibody 2nd antibody : Anti-Goat IgG-HRP (Product No. SC-2350) diluted 1:5,000, 1 hr. at RT Detection : Chemi-Lumi One Super (Product No. 02230), 1 min. reaction time

Detector : LAS-3000 (High mode), 10 min. exposure time

(Please note)

For antibody dilution, use Bullet Blocking One undiluted, or dilute up to 20x with TBS or PBS containing 0.05-0.1% detergent, such as Tween 20. Since the appropriate dilution ratio depends on antibody conditions, such as type and concentration, pretest is required.

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
		13779-56	50 ml
Bullet Blocking One for Western Blotting	R	13779-14	200 ml
		13779-01	1 L

#### [Ultra-fast Antibody Dilution]

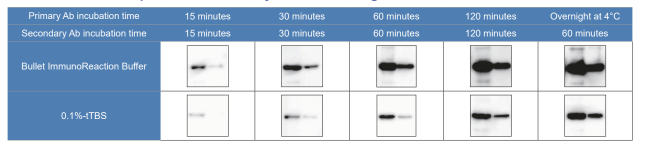
## **Bullet ImmunoReaction Buffer**

» Saving-time : Cut in half antigen-antibody reaction time

» Simple : Replace tTBS, tPBS, skim milk or BSA with Bullet ImmunoReaction Buffer

» Ready-to-use : Do not need to dilute

#### Reference data: Comparison of sensitivity related to Ab-Ag reaction time with 0.1% tTBS



(Conditions)

Sample : Hela cell extract; (left) 2 µg, (right) 0.4 µg

Blocking : Incubated for 5 minutes with Bullet Blocking One for Western Blotting manufactured by Nacalai, #13779

Primary antibody : β-Actin antibody manufactured by Santa Cruz, #sc-47778, x 1,000 dilution

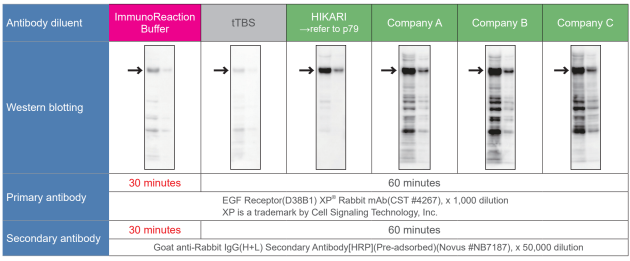
Secondary antibody : Goat anti-mouse IgG(H+L) secondary antibody manufactured by Novous, NB7574, x 40,000 dilution

Detection : Chemi-Lumi One Super manufactured by Nacalai, #02230, 5-minute exposure time

Detector : LAS-300 High mode

#### Transition to saving-time protocol

If a signal enhancer reagent, such as HIKARI, is use, speeding up the process with this product is not recommended, as it may decrease sensitivity and increase non-specific reactions.



#### (Conditions)

Sample : Hela cell extract, (left) 10 µg and (right) 2 µg

Blocking : Incubated for 5 minutes with Bullet Blocking One

Detection : Chemi-Lumi One Super, #02230; 3-minute exposure time

. Chemi-Lumi One Super, #02250, 5-minute exposure

Detector : LAS-300 High mode

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Bullet ImmunoReaction Buffer	R	18439-85	500 ml

### [High-performance Antibody Dilution]

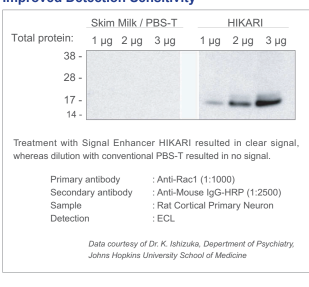
# Signal Enhancer HIKARI for Western blotting and ELISA

Dilute your antibodies with Signal Enhancer HIKARI instead of conventional diluents such as PBS-t or TBS-t before performing your next western blotting detection protocol and witness a remarkable increase in the ability to detect the protein of interest and to eliminate undesired background. Signal Enhancer HIKARI was developed to resolve the problems of low sensitivity and high background often encountered during procedures such as western blotting and ELISA.

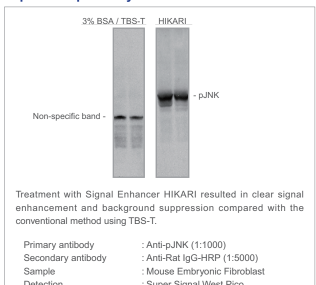
- » Enhances antigen-antibody reactions
- » Removes background
- » Works with any substrate and membrane
- » Ready-to-use reagent



#### Improved Detection Sensitivity



#### Improved Specificity



Detection : Super Signal West Pico

> Data courtesy of Dr. S. Matsuzawa, Signal Transduction, NCI Cancer Center, Burnham Institute for Medical Research

#### Referenes

- 1. Feng-Ming Yang et al. FEBS 276, 425-436 (2009)
- 2. Jian-Bin Wang et al. The Journal of Cell Science 122(12), 2024-2033 (2009)
- 3. Chunwei Huang et al. Reproductive Toxicology 27, 103-110 (2009)
- 4. Sawako Yamashiro et al. The Journal of Cell Science 121 (Pt 23), 3867-3877 (2008)

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Signal Enhancer HIKARI for Western Blotting and ELISA  Kit contents: Solution A for Primary Antibody  Solution B for Secondary Antibody	R	02267-41 02270-81	1 set ( 50 ml each) 1 set (250 ml each)
Signal Enhancer HIKARI for Western Blotting and ELISA Solution A	R	02272-74	250 ml
Signal Enhancer HIKARI for Western Blotting and ELISA Solution B	R	02297-64	250 ml

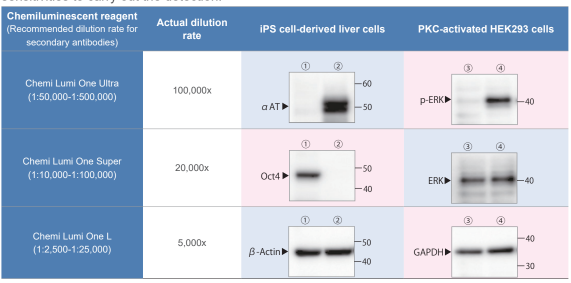
## HRP-conjugated secondary antibody

- » Affinity-purified form
- » Cross-absorbed form\*

\*Anti-Mouse IgG: Cross-absorbed with Human IgG, Rat IgG, and Rabbit IgG Anti-Rabbit IgG: Cross-absorbed with Human IgG, Rat IgG, and Mouse IgG

#### **Application data**

We detected target proteins in liver cells derived from induced pluripotent stem (iPS) cells, as well as in cells treated for Protein Kinase C activation. These proteins are related to cellular differentiation and phosphorylation processes. We employed three different chemiluminescent reagents with varying sensitivities to carry out the detection.



Blue: Anti-Mouse IgG(#21860) Pink: Anti-Rabbit IgG(#21858)

(Conditions)

Sample: ① Control (I iPS cells 10  $\mu$ g) ② iPS cell-derived liver cells 10  $\mu$ g ③ Control (I HEK293 cells 10  $\mu$ g) ④ PKC-activated HEK293 cells 10  $\mu$ g Primary antibody: Diluted with Bullet ImmunoReaction Buffer (#18439-85) and incubated for 30 mins at room temperature.

[ $\alpha$ AT] (Proteintech #66135-1-Ig), [Oct4] (Abcam #ab19857), [ $\beta$ -Actin] (MBL #M177-3), [ $\rho$ ERK] CST #4370S [ERK] (Santa Cruz #sc-514302), [GAPDH] (Novus Biologicals #NB300-322)

Secondary antibody: Diluted with Bullet ImmunoReaction Buffer (#18439-85) and incubated for 30 mins at room temperature.

#### Comparison of detection sensitivity by western blotting

We detected samples that were serially diluted two-fold, from 16  $\mu$ g/well using Anti-Mouse IgG and Anti-Rabbit IgG antibodies sourced from various companies at a 20,000-fold dilution. Our product demonstrated equal or higher sensitivity compared to products from other companies.



(Conditions)

 $Reference\ (https://www.nacalai.co.jp/products/entry/d005021.html)$ 

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Anti-Mouse IgG(Goat), HRP-conjugated, Pre-absorbed	В	21860-74	100 µL
	R	21860-61	1 ml
Anti-Rabbit IgG(Goat), HRP-conjugated, Pre-absorbed	В	21858-24	100 µL
	R	21858-11	1 ml

#### [Detection]

## **Chemiluminescent Western Blotting Substrates**

Chemi-Lumi One is a series of high sensitive luminol-based chemiluminescence assay Kits for western blotting. Three types of chemiluminescent substrates are available for western blotting detection with horseradish peroxidase enzyme (HRP).

Product Name	Chemi-Lumi One L	Chemi-Lumi One Super	Chemi-Lumi One Ultra
Product No.	07880	02230	11644
Lower Detection Limit	Low-picogram	Mid-femtogram	Low-femtogram
Required Working Solution	Approx. 0.125 ml / cm <sup>2</sup>	Approx. 0.1 ml / cm <sup>2</sup>	Approx. 0.1 ml / cm <sup>2</sup>
Suggested Antibody Dilution Ratio	Primary: 1:1,000-1:5,000 Secondary: 1:20,000-1:100,000	Primary: 1:1,000-1:20,000 Secondary: 1:20,000-1:200,000	Primary: 1:5,000-1:100,000 Secondary: 1:100,000-1:500,000
Reaction Period	1 min.	1 min.	5 min.
Comparable to	ECL (Cytiva) ECL Start (Cytiva) SuperSignal Pico (Thermo)	ECL Prime (Cytiva) SuperSignal Dura (Thermo) SuperSignal PLUS (Thermo) Clarity (Bio-Rad)	ECL Select (Cytiva) SuperSignal Femto (Thermo) Clarity Max (Bio-Rad)
Sensitivity	General Use	High Sensitivity	Ultra high Sensitivity
<pre><conditions> Antigen</conditions></pre>	Conc. of IgG HRP-linked 900 Chemi-Lumi One L  Chemi-Lumi One Chemi-Lumi One	1111	Femtogram

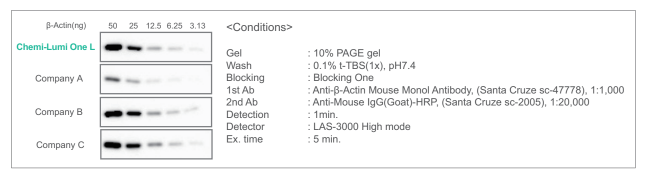
#### Chemi-Lumi One L

- » Suitable for optimization of target proteins
- » Reasonable price
- » Detect wide range of protein concentration

# The state of the s

#### Comparison of sensitivity with competitors

Chemi-Lumi One L offers similar sensitivity to competitors' and higher sensitivity than Company A's products.

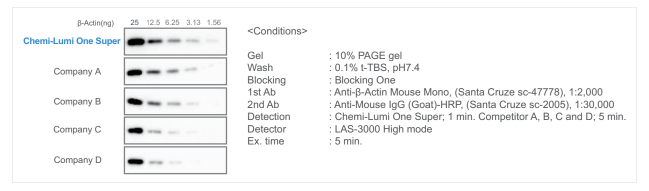


#### Chemi-Lumi One Super

- » Extendable of exposure time
- » Detects proteins at mid-femtogram level with low background
- » Rapid substrate processing of blot

#### Comparison of sensitivity with competitors

Chemi-Lumi One Super offers the highest sensitivity out of competitors' substrates even though its exposure time is 1 minute, while others require 5 minutes.



#### Chemi-Lumi One Ultra

- » The most sensitive in Chemi-Lumi One series
- » Longer signal duration
- » Wider range of experimental conditions due to low background



#### Comparison of sensitivity with competitors

Western blot of identical transferrin samples. The membranes were incubated with substrate that was prepared according to the manufacturers' instructions. The membranes were exposed to film for 2 minutes.



<Conditions>

Gel: 10% acrylamidegel

Wash : 0.1% t-TBS(x1), pH7.4 Detector : LAS-3000 High mode

Blocking : Blocking One Expose time : 3 min.

1st Ab : Anti- $\beta$ -Actin Mouse Monoclonal Antibody, (Santa Cruze sc-47778), 1:300 2nd Ab : Anti-Mouse IgG (Goat), HRP Conjugated, (Santa Cruze sc-2005), 1:1,000

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Chemi-Lumi One L, Luminol 250 ml and Peroxide 250 ml Sufficient substrate for 4,000 cm² of blotting membrane	R	07880-70	1 KIT
Chemi-Lumi One Super, Luminol 50 ml and Peroxide 50 ml Sufficient substrate for 1,000 cm² of blotting membrane	R	02230-30	1 KIT
Chemi-Lumi One Ultra, Luminol 50 ml and Peroxide 50 ml Sufficient substrate for 1,000 cm² of blotting membrane	RT	11644-40	1 KIT

Detection period : 5 min.

## **Colorimetric Western Blotting Substrates**

Colorimetric detection enables detection of a target protein on a membrane by a simple procedure without usage of detection equipment. Depending on the enzyme type conjugated to the antibody, some detection kits are available.

#### TMB Solution for western blotting

TMB (tetramethyl benzidine) solution is used with horseradish peroxidase (HRP)-based western blotting procedure, but not applicable for ELISA or immunohistochemistry.

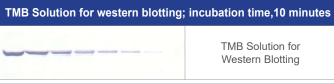
» Convenience : No CCD-camera-equipped imager or X-ray film needed

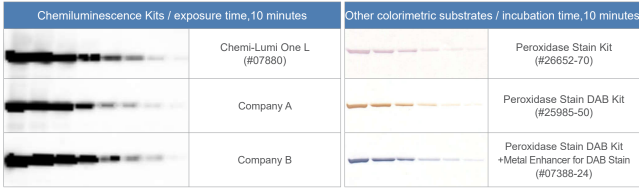
» Sensitivity : Low picogram level

» Safety : Does not contain DAB or o-Dianisidine

» Ready-to-use : Does not need to dilute

Reference data: Comparison with other picogram-level chemiluminescence kits and colorimetric substrates





Sample : Hela cell extraction, 2 times dilution started from 12 ug

Primary antibody : Anti-β-actin mAb manufactured by MBL, #177-3, x 5,000 dilution

Secondary antibody : Anti-mouse IgG, HRP-linked whole Ab Sheep manufactured by GE, #NA931, x 10,000 dilution

Detector for chemiluminescence : LAS-3000 Super mode

## BCIP-NBT Solution (Ready-To-Use)

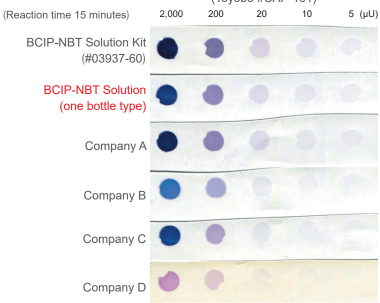
A ready-to-use solution of chromogenic substrate of alkaline phosphatase (AP). It can be used in a wide range of applications, such as detection of alkaline phosphatase (AP) bound by antigen-antibody reaction and endogenous alkaline phosphatase (AP) of stem cells and osteoblasts, in blotting and tissue or cell staining.

- » Ready-to-use
- » Can be used in wide range of applications such as immunochemical staining or endogenous AP staining
- » Can be used in situ hybridization (ISH), as it is tested for DNase and RNase

#### **Comparison: Detection sensitivity by dot blotting**

Compare with other brands, BCIP-NBT Solution has equal or better sensitivity.





#### **Ordering Informattion**

Product Name	Storage	Product No.	PKG Size
TMB Solution for Western Blotting	R	18186-24	200 ml
BCIP-NBT Solution (Ready To Use)	R	19880-84	100 ml

#### **Related products**

Product Name	Storage	Product No.	PKG Size
Peroxidase Stain DAB Kit (Brown Stain)	R	25985-50	1 KIT
Metal Enhancer for DAB Stain	RT	07388-24	100 ml

## **WB Stripping Solution**

WB Stripping Solution removes antibodies from blotting membrane, enabling subsequent detections with different antibodies on the very same blot. After the first antigen-antibody reaction and following chemiluminescent visualization, the antibodies can be removed by the WB Stripping Solution. A second antigen-antibody reaction can be conducted on the same blot. The same blot can be probed 2-5 times if chemiluminescent detection is employed.

» No heating : Reaction at room temperature

» No odor : Does not contain 2-mercaptoethanol

» Fast : Stripping time 5-15 minutes» Ready-to-use : One solution in one bottle

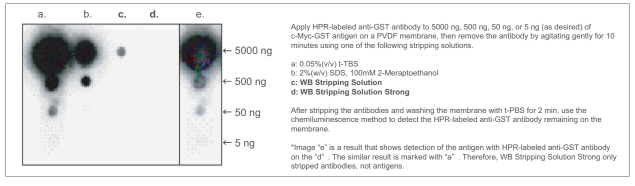




#### **Applications**



#### Comparison of WB Stripping Solution and WB Stripping Solution Strong



#### **Ordering Informattion**

Product Name	Storage	Product No.	PKG Size
WB Stripping Solution	RT	05364-55	500 ml
WB Stripping Solution Strong	R	05677-65	500 ml
WB Stripping Solution Trial Set (WB Stripping Solution: 40 ml, WB Stripping Solution Strong: 40 ml)	R	05680-21	1 set

## **Epitope Tag Antibody**

A family of epitope tag antibodies available for the detection and purification of the recombinant proteins. Most of our tag antibodies are highly specific mouse and rat monoclonal antibodies.

#### Anti-GFP (Rat IgG2a), Mono (GF090R)

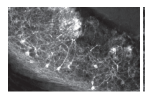
Clone : GF090R Isotype : IgG2a (Rat) Product form : Liquid

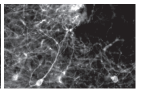
Immunogen : His-GFP (full length) fusion protein

Application : Immunohistochemstry 1:1000-1:2000

Western Blotting 1:1000-1:2000 ELISA 1:2000-1:20000

#### **Immunohistochemistry**





Sample : Mouse brain (nerve cell)

Primary antibody : Anti-GFP(Rat IgG2a), Mono (1:1000) RT, O/N

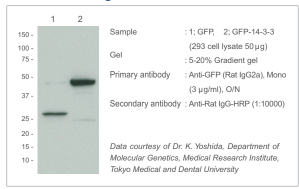
Secondary antibody: Anti-Rat IgG-Cy3 (1:300) RT, 1 hr

Blocking 5% Normal goat serum/0.2% TritonX-100 in PBS

Fixing method : 4% Paraformaldehyde

Data courtesy of Dr. Y. Yoshihara, RIKEN Brain Science Institute

#### **Western Blotting**



#### Reference

- 1. Nakamura, M. et al. Molecular Vision 16, 425-437 (2010)
- 2. Nishide, K. et al. PLoS ONE 4(8), e6869 (2009)
- 3. Nagao, M. et al. The Journal of Cell Biology 183(7), 1243-1257 (2008)
- 4. Ono, K. et al. Development Biology 320(2), 356-468 (2008)
- 5. Nakashiba T. et al. Science 319(5867), 1260-1264 (2008)
- Batista-Brito R et al. The Journal of Neuroscience 28(15), 3966-3975 (2008)
- 7. Esumi S. et al. Neuroscience Research **60**(4), 439-451 (2008)
- 8. Fogarty M. et al. The Journal of Neuroscience 27(41), 10935-10946 (2007)
- 9. Sasamura T. et al. Development 134, 1347-1356 (2007)
- 10. Sato Y. et al. The Journal of Neuroscience 27(7), 1606-1615 (2007)
- 11. Yamada M. et al. Stem Cells 25(3), 562-570 (2007)
- 12. Togashi H. et al. The Journal of Cell Biology 174, 141-151 (2006)
- 13. Ogata M. et al. Molecular and Cell Biology 26(24), 9220-9231 (2006)
- 14. Sato Y. et al. The Journal of Neuroscience 25(20), 4889-4897 (2005)

#### Anti-GFP (Mouse IgG1-k), Mono (GF200)

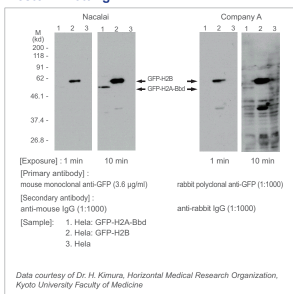
Clone : GF200 Isotype : IgG1-k (Mouse)

Product form : Liquid

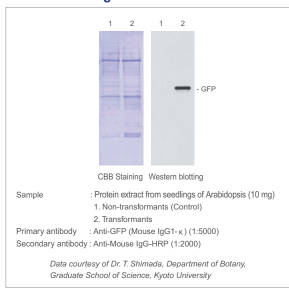
Immunogen: His-GFP (full length) fusion proteinApplication: Western Blotting1:1000-1:2000

ELISA 1:2000-1:20000

#### Western Blotting



#### **Western Blotting**



#### Anti-c-Myc (Mouse IgG1-k), Mono (MC045)

Clone : MC045

Isotype : IgG1-k (Mouse)

Product form : Liquid

Immunogen : c-Myc synthetic peptide [EQKLISEEDL]

conjugated with KLH

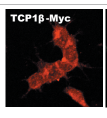
Application : Western Blotting 1:1000-1:2000

 Immunoprecipitation
 1:400-1:1000

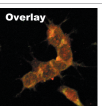
 Immunocytochemstry
 1:400-1:1000

 ELISA
 1:2000-1:20000

#### **Immunocytochemistry**







Sample SH-SY5Y cell in which c-Myc-tagged TCP1β is expressed

Primary antibody : Anti-c-Myc (Mouse IgG1-κ) (2.5 μg/ml)

Anti-Protein X

Secondary antibody : Anti-Fluor 546-conjugated antibody (1:400)

Anti-Fluor 488-conjugated antibody (1:400)

Data courtesy of RIKEN Brain Science Institute

#### Anti-6xHis (Mouse IgG1a-k), Mono (HI192)

Clone : HI192

Isotype : IgG1-k (Mouse)

Product form : Liquid

Immunogen : 6xHis synthetic peptide [HHHHHH]

conjugated with KLH

Application : Western Blotting 1:1000-1:2000

ELISA 1:2000-1:20000

#### Western Blotting



Sample:

Lysate from Sf9 cells in which His-tagged Drosophila PTEN is expressed

Filter : FluoroTrans [PALL]

Blocking : 5% non-fat dry milk/PBS (30 min)

Primary antibody : Anti-6xHis (Mouse IgG1ĸ), Monoclonal (HI192)

Wash : 0.25% Tween-20/TBS (10 min x 3)

Secondary antibody: Anti-Mouse IgG-HRP (1:1000) /2% BSA/0.25% Tween-20

/TBS (30 min)

Wash : 0.25% Tween-20/TBS (10 min x 5)
Detection : Luminol Reagent (Santa Cruz)

Data courtesy of Dr. A. Maehama, The Tokyo Metropolitan Institute of Medical Science (RINSHOKEN)

#### **Ordering Information**

Product Name	Application	Storage	Product No.	PKG Size
Anti-c-Myc (Mouse IgG1-k), Monoclonal (MC045), AS	WB, IP,	R	04362-76	50 μg
Anti-c-Myc (Mouse 1961-k), Monocional (Moo45), AS	ICC, ELISA	K	04362-34	200 μg
Anti-c-Myc (Mouse IgG1-κ), Monoclonal (MC045), AS, Agarose Conjugate	IP	R	04145-55	500 μg
Anti-GFP (Mouse IgG1-k), Monoclonal (GF200), AS	WB, ELISA	R	04363-66	50 μg
		K	04363-24	200 µg
Anti CER (Ret IgC2e) Managional (CE000R) CC	WB, IHC	В	04404-26	50 μg
Anti-GFP (Rat IgG2a), Monoclonal (GF090R), CC	ELISA	R	04404-84	200 µg
Anti-GFP (Rat IgG2a), Monoclonal(GF090R), CC, Agarose Conjugate	IP	R	06083-05	500 μg
Anti-6xHis (Mouse IgG1-k), Monoclonal (HI192), AS	WB FLICA	В	04428-26	50 μg
	WB, ELISA	R	04428-84	200 µg

## 4% - Paraformaldehyde Phosphate Buffer Solution

We offer a 10% formalin neutral buffer solution, which contains about 4% formaldehyde, designed for use as a general fixation buffer in histological specimen preparations. Since this product is based on commonly available formalin, methanol is used as a stabilizer.

If a methanol free formalin solution is required, paraformaldehyde is used instead of formaldehyde. However, paraformaldehyde is extremely toxic and can cause injury if scattered. To deal with this hazard, additional work, such as making the solution alkaline with heating when dissolving the paraformaldehyde, is required. Our product is available in two volume types: 500 ml and a 5 x 10 ml package set.



- » Small unit volume
- » Enables to immerse histological specimens directly into the solution (5 x 10 ml)
- » Low cost for waste
- » Ready-to-use
- » Storable in refrigerator

#### **Usage examples**

#### Hematoxylin-eosin staining



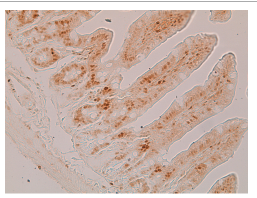
Sample Fixation method : Mouse small intestine

: Immersion fixation with this product

(over night at 4°C)

Deparaffinization : Limonene and ethanol

#### Immunohistological staining



Sample

Mouse small intestine

Fixation method : Immersion fixation with this product

(overnight at 4°C)

Deparaffinization

: Limonene and ethanol

Primary antibody : Anti-PCNA (FL-261) (rabbit)
Staining : Peroxidase Stain DAB Kit (bro

: Peroxidase Stain DAB Kit (brown stain) (Product No. 25985-50)

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
40/ Desertance Idebude Discoule to Duffer Colution	R	09154-14	5 x 10 ml
4% - Paraformaldehyde Phosphate Buffer Solution		09154-85	500 ml

## HistoVT One (10x, pH 7.0)

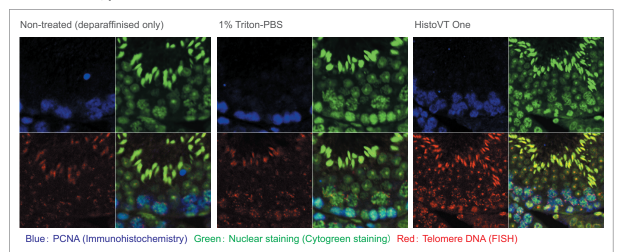
HistoVT One is an antigen retrieval solution for immunohistochemistry and *in situ* hybridization. This product can unmask antigenic sites without damage to antigen from formalin-fixed, frozen or paraffinembedded tissue sections.

- » Enhancing antigen-antibody reaction
- » Usable with frozen or paraffin-embedded tissue section
- » High reproducibility

#### **Application 1**

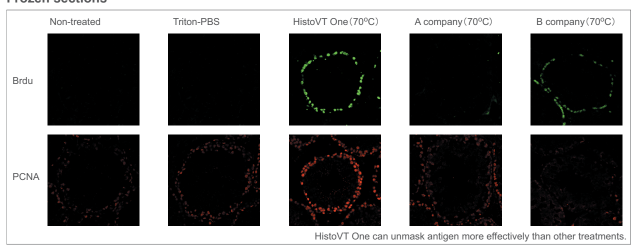
Formalin-fixed, paraffin-embedded tissue sections





#### **Application 2**

#### Frozen sections



Data courtesy of RIKEN Brain Science Institute, Brain Development Research Group

#### **Ordering Information**

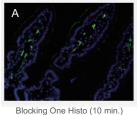
Product Name	Storage	Product No.	PKG Size
HistoVT One (10x, pH 7.0)	RT	06380-05	500 ml

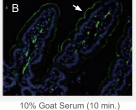
## **Blocking One Histo**

Blocking One Histo is a blocking solution to prevent non-specific binding of antibodies in immunohistochemistry (IHC). The product is designed for immunohistochemistry application based on Blocking One (refer to Western Blotting Section).

- » Eye-drop bottle
- » Can be used for immunofluorescence staining
- » The preservative does not affect the activity of alkaline phosphatase (AP) or horseradish peroxidase (HRP)

#### Comparison of blocking efficiency with 10% goat serum (Immunofluorescence)





Antigen Retrieval Primary antibody

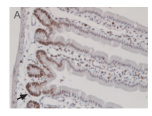
Mouse small intestine (Paraffin-embedded section) Histo VT One, 90°C, 20 min.

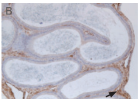
Anti-Vimentin rabbit polyclonal antibody (Santa Cruz: #sc-7557R)

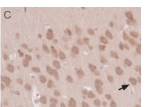
: CFTM 488A Goat anti-Rabbit IgG(H+L), F(ab')<sub>2</sub> Fragment (Biotium: #20013)

In both panels, mouse small intestine tissue section was stained with secondary antibody conjugated with CF™ 488A (green) and counter stained with DAPI (blue). In the panel B with 10% Goat Serum, the stained white arrow along the lines of shape of small intestine show non-specific staining. Blocking One Histo is more effective at reducing non-specific background staining than normal serum.

#### **Applications**







- A: Mouse small intestine (PCNA) x5
- B: Mouse epididymis (Vimentin) x25
- C: Mouse brain (GluR) x100

Primary antibody

Detection

- Histo VT One, Room temp., 10 min. A: Anti-PCNA rabbit pAb (Santa Cruz: #sc-7907)
- B: Anti-Vimentin rabbit pAb (Santa Cruz: #sc-7557R) C: Anti-GluR-1 goat pAb (Santa Cruz: #sc-7608)
- B: Goat anti-rabbit IgG (H+L), biotin conjugated (Vector, #BA-1000)
  B: Goat anti-rabbit IgG (H+L), biotin conjugated (Vector, #BA-1000) Secondary antibody

  - C: Bovine anti-goat IgG (H+L), biotin conjugated (Santa Cruz: #sc-2347) Streptavidin Biotin Complex Peroxidase Kit (Product No. 30462)

  - Peroxidase Stain DAB Kit (Brown Stain) (Product No. 25985)

Blocking treatment of each tissue section had been performed by Blocking One Histo. Mouse small intestine (panel A) was stained with anti-PCNA and DAB (3,3'-Diamino Benzidine) to stain nuclear (black arrow), Mouse epididymis (panel B) was stained with anti-Vimentin and DAB to stain muscle (black arrow), Mouse brain (panel C) was stained with anti-GluR and DAB to stain membrane proteins (black arrow) and counter stained with hematoxylin.

#### **Ordering Information**

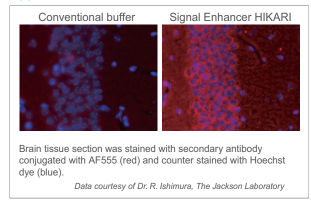
Product Name	Storage	Product No.	PKG Size
Blocking One Histo	R	06349-64	50 ml

## Signal Enhancer HIKARI for Immunostain

Signal Enhancer HIKARI for Immunostain was developed to resolve the problems of low sensitivity and high background often encountered during immunostain procedures such as immunohistochemistry (IHC) and immunocytochemistry. Dilute your antibodies with Signal Enhancer HIKARI for Immunostain instead of conventional diluents such as PBS or TBS before performing your next IHC experiment and witness a remarkable increase in the ability to detect the protein of interest and to eliminate unwanted background.

- » Enhances signals
- » Reduces background
- » Ready-to-use reagent
- » Works with any detection system
- \* The kit can also be used in combination with sensitizing systems such as the ABC or polymer complex method.

#### **Applications**

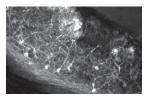


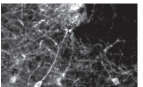
#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Signal Enhancer HIKARI for Immunostain Trial Set	R	02363-71	1 set (5 ml each)
Signal Enhancer HIKARI for Immunostain Solution A	R	02373-54	20 ml
Signal Enhancer HIKARI for Immunostain Solution B	R	02375-34	20 ml

## Anti-GFP (Rat IgG2a), Monoclonal (GF090R)

- » Immunohistochemical application
- » Rat monoclonal antibody





Sample : Mouse brain (nerve cell)

Primary antibody : Anti-GFP(Rat IgG2a), Mono (1:1000) RT, O/N

Secondary antibody: Anti-Rat IgG-Cy3 (1:300) RT, 1 hr

Blocking 5% Normal goat serum/0.2% TritonX-100 in PBS

Fixing method : 4% Paraformaldehyde

Data courtesy of Dr. Y. Yoshihara, RIKEN Brain Science Institute

#### **References for Immunostaining**

- 1. Takeshi Sasamura et al. Development 134, 1347-1356 (2007)
- 2. Takashi Inoue et al. The Journal of Neuroscience, **27**(20), 5461–5473 (2007)
- 3. Anoop Kumar G. Velikkakath et al. Mol Biol Cell. 23(5), 896-909 (2012)
- 4. Eisuke Itakura et al. Molecular Biology of the Cell, 19, 5360-5372,
- 5. Keith N. Brown, et al. Science, 334, 480 (2011)
- Anna N. Rubin et al. The Journal of Neuroscience, 30(36), 12050-12062 (2010)
- 7. Shinsuke Nakao et al. J Cell Biol., 182(2), 395-410 (2008)
- 8. Maiko Ogata et al. Mol Cell Biol., 26(24), 9220-9231 (2006)
- 9. Matthew Fogarty et al. The Journal of Neuroscience, **27**(41), 10935-10946 (2007)
- 10. Takuya Sato et al. Nature Communications, 2 (472)
- 11. Toshiaki Nakashiba et al. Science, **319**(5867), 1260-4 (2008)
- 12. Hiromi Takanaga et al. Stem Cells, 27(1), 165-74 (2009)
- 13. Akinori Yamasaki et al. Mol. Biol. Cell , 17 (11), 4876-4887 (2006)
- Natsumi Ageta-Ishihara et al. The Journal of Neuroscience, 29(43), 13720-13729 (2009)
- 15. Naoyuki Asada et al. Journal of Neuroscience, 30(26), 8852-8865 (2010)
- 16. Shizue Ohsawa et al. Dev Cell, 20(3), 315-28 (2011).

#### **Ordering Information**

For more informations about Epitope Tag Antibody, please refer to p87.

Product Name	Storage	Product No.	PKG Size
Anti-GFP(Rat IgG2a), Monoclonal(GF090R), CC	R	04404-26	50 μg
Anti-GFP(Rat IgG2a), Monoclonal(GF090R), CC, Agarose Conjugate	R	06083-05	500 μg

## **Streptavidin Biotin Complex Peroxidase Kit**

Streptavidin Biotin Complex Peroxidase Kit includes reagents for the "Avidin-Biotin Complex, ABC technique", a highly sensitive method for immunoblotting, immunohistochemistry, ELISA and *in situ* hybridization.



#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Streptavidin Biotin Complex Peroxidase Kit	R	30462-30	1 KIT

# High Sensitivity Peroxidase DAB Stain

#### Peroxidase DAB Stain Kit and Metal Enhancer

Peroxidase Stain DAB Kit (Brown Stain) is used to detect horseradish peroxidase (HRP) activity and stain them brown in immunoblotting, immunohistochemistry and *in situ* hybridization.

Metal Enhancer for DAB Stain (Product No. 07388-24) is used to stain peroxidase purplish gray with Peroxidase Stain DAB Kit (Brown Stain)(Product No. 25985-50) in immunoblotting, immunohistochemistry and *in situ* hybridization. The sensitivity of Metal Enahcer for DAB Stain used with Peroxidase Stain DAB Kit (Brown Stain) is about two times higher than the product with Peroxidase Stain DAB Kit (Brown Stain) alone.

- » High sensitivity
- » Metal Enhancer for DAB Stain stains peroxidase purplish gray (Just change the solution mix from water to Metal Enhancer for DAB)
- » RNase, DNase free, applicable to in situ hybridization
- » Eye drop bottle





#### Procedure of combination Peroxidase Stain DAB Kit and Metal Enhancer Solution

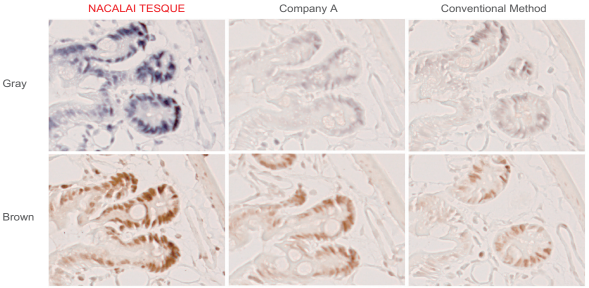


Change the adjusting solution of Peroxidase Stain DAB Kit (Brown Stain) from water to Metal Enhancer for DAB.

#### **Application**

#### Immunohistostaining of mouse small intestines with anti-PCNA antibody (Serial membranes)

The sensitivity achieved when the Peroxidase Stain DAB Kit (Brown Stain) is used alone is higher than the competitors' products. However, when used in conjunction with Metal Enhancer for DAB stain, the sensitivity of Peroxidase Stain DAB Kit (Brown Stain) is about two times higher than what can be achieved by the Peroxidase Stain DAB Kit (Brown Stain) alone.



Reaction Time: 7 min.

NACALAI TESQUE : (Gray) Peroxidase Stain DAB Kit (Brown Stain) + Metal Enhancer for DAB

(Brown) Peroxidase Stain DAB Kit (Brown Stain)

Vector : (Gray) Kit (witth attached nickel solution)

(Brown) Kit (without attached nickel solution)

Basic method : (Gray) 0.6mg/ml DAB, 0.03%H2O2, 50mM Tris-HCl Buffer pH7.6, 0.03%NiCl2

(Brown) 0.6mg/ml DAB, 0.03%H $^2$ O $^2$ , 50mM Tris-HCl Buffer pH7.6

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Peroxidase Stain DAB Kit (Brown Stain)	R	25985-50	1 KIT
Metal Enhancer for DAB Stain	RT	07388-24	100 ml

## **Mounting Medium for Fluorescent Staining**

Fluoro-KEEPER Antifade Reagent is a non-hardening mounting medium with a unique antifade reagent. It suppresses rapid photobleaching during fluorescence microscopy observation. The coverslipped slide with nail polish or other sealants can be stable for several weeks. There are two types of products available, with DAPI [4',6-Diamidino-2-phenylindole] and without DAPI, which counterstains nucleus blue.

- » Inhibits photobleaching of various fluorescent dyes
- » Easy to use with eye-drop bottle

#### **Application 1 (Fluoro-KEEPER with DAPI)**

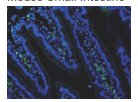
#### Fluorescent microscopy experiments

Fluoro-KEEPER with DAPI offers nuclear staining along with mounting.

Mounting Medium : Fluoro-KEEPER with DAPI for 30 min. at room temperature protecting from light.

Microscopy : Olympus BX-50-34-FLA1

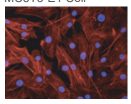
#### Mouse Small Intestine



Antigen retrieval : HistoVT One (#06380)
Blocking : Blocking One Histo (#06349)

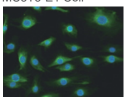
1st Ab : Anti-Vimentin Rabbit Polyclonal Antibody (Santa Cruz #sc-7557R)
2nd Ab : CFTM 488A Goat Anti-Rabbit IgG (H+L), Fragment (Biotium #20013)

#### MC3T3-E1 Cell



Blocking : Blocking One Histo (#06349)
Rhodamine-conjugated phalloidin (Cytoskeleton #PHDR1)

#### MC3T3-E1 Cell



Blocking : Blocking One Histo (#06349)

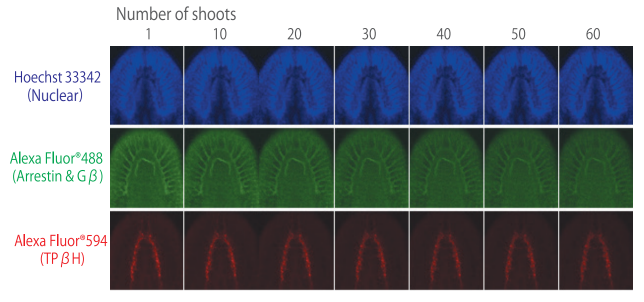
1st Ab : Anti-Vimentin Rabbit Polyclonal Antibod (Santa Cruz #sc-7557R) 2nd Ab : Cy®2 Goat Anti-Rabbit IgG (H+L)(GENETEX #GTX26940)

#### **Application 2 (Fluoro-KEEPER without DAPI)**

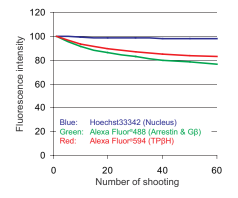
#### Fluorescent microscopy experiments: Planarian

Fluorescence intensities are shown as percentages of initial intensities remaining during repeated frame capture up to 60 times. The images were acquired by Olympus FV10. The samples mounted in the Fluoro-KEEPER Antifade Reagent were clearly detected after 60 times of frame capture.

#### ■ Stain Image



#### ■ Fluorescence intensity



#### Nuclear

Hoechst 33342 Arrestin and G Protein  $\beta$  Subunit(G $\beta$ )

Art Ab. Mayor Anti-planarian

1st Ab: Mouse Anti-planarian Arrestin Mouse Anti-planarian Gβ

2nd Ab: Alexa Fluor® 488 Goat Anti-mouse IgG

Tryptophan  $\beta$  Hydroxylase(TP $\beta$ H)

1st Ab: Rabbit Anti-planarian TPβH

2nd Ab: Alexa Fluor® 594 Goat Anti-rabbit IgG

Samples were mounted in the Fluoro-KEEPER Antifade Reagent

Data courtesy of Agata Lab, Department of Biophysics, Kyoto University

#### Comparison of antifade effectiveness with different fluorescent dyes

Fluoro-KEEPER Antifade Reagent offers increased resistance to photobleaching of various fluorescent dyes.

with DAPI with DAPI

Fluorescence Dye	Fluoro- KEEPER	Control	Fluoro- KEEPER	Control
Hoechst 33258	98	96	_	_
Hoechst 33342	100	90	_	_
DAPI	99	93	_	_
Propidium Iodide	95	67	_	_
Fluorescein	97	49	96	49
Alexa Fluor® 488	93	86	96	91
CF <sup>™</sup> 488	93	82	91	82
Cy® 2	99	83	98	81
Rhodamine	72	51	78	41
Alexa Fluor® 555	98	81	97	87
CF <sup>™</sup> 555	98	85	97	85
Cy® 3	89	71	86	66

Cells stained by each fluorescent dye were mounted in Fluoro-KEEPER Antifade Reagent, 85% Glycerol containing PBS as a control. Samples were illuminated for 60 seconds. Each number indicates fluorescence intensity as percentage of initial intensity after 60 seconds exposure photobleaching.

Control Condition: 85% Glycerol-PBS w/o DAPI 85% Glycerol-PBS w/ DAPI

Fluorescent Microscopy: Olympus BX-50-34-

FLA1

Exposure time: 60 seconds.

#### **Ordering Information**

Product Name	Storage	Product No.	PKG Size
Fluoro-KEEPER Antifade Reagent, Non-Hardening Type	R	12593-64	2 x 5 ml
Fluoro-KEEPER Antifade Reagent, Non-Hardening Type with DAPI	R	12745-74	2 x 5 ml

## NACALAI TESQUE, INC.

Nijo Karasuma, Nakagyo-ku, Kyoto 604-0855 JAPAN

TEL : +81 (0)75 251 1730 E-mail : info.intl@nacalai.com





#### ATTENTION:

Nacalai Tesque makes no representation or warranty as to whether the products and/or their uses infringe any patent of any third party, nor shall Nacalai Tesque be liable for infringement of any such patent.

#### Warranties and Disclaimers:

Nacalai Tesque warrants that its products shall conform to the description of such products as provided by Nacalai Tesque through its catalog, analytical data or other literature. Nacalai Tesque makes no other warranty, express or implied, as to the fitness of these products for any particular purpose. Nacalai Tesque shall not in any event be liable for incidental or consequential damages that may result from any use or failure of the products.

For more information on products and pricing, please contact your local distributor.