

Instructions Code No. 27408

Code No. 27408 Rat a2, 6-Sialyltransferase (E41 Form) Assay Kit - IBL

INTRODUCTION

The histopathological picture of Alzheimer's disease is characterized by senile plaques and neurofibiliary tangles, and because the senile plaques form first, they are considered the initial lesion. Senile plaques are known to be formed by accumulation of β -amyloid peptide (A β). A β peptide is produced by the cleavage of amyloid precursor protein (APP) by two types of proteolytic enzymes. The first cleavage is performed by β -secretase (BACE1), and the second γ -secretase. It is thought that their inhibitors may be capable of serving as safe drugs for the treatment of Alzheimer's disease.

In recent years a glycosyltransferase involved in the biosynthesis of sugar chains (rat $\alpha 2$, 6-sialyltransferase) has also been shown to be cleaved by BACE1. The cleavage site was identified at the same time, and as a result it was demonstrated that in rats it produces cleavaged-type rat $\alpha 2$, 6-sialyltransferase (E41 Form). This is an EIA kit for quantitative determination of rat $\alpha 2$, 6-sialyltransferase (E41 Form).

PRINCIPLE

This kit is a solid phase sandwich ELISA using 2 kinds of high specific antibodies. Tetra Methyl Benzidine (TMB) is used as a coloring agent (Chromogen). The strength of coloring is in proportional to the quantities of Rat $\alpha 2$, 6-sialyltransferase (E41 Form)

MEASUREMENT RANGE

0.31 ~ 20 ng/mL

INTENDED USE

The IBL's Rat α 2, 6-Sialyltransferase (E41 Form) Assay Kit is a complete kit for the quantitative determination of Rat α 2, 6-sialyltransferase (E41 Form) in EDTA-plasma, serum, cell culture media.

KIT COMPONENT

- : Anti-a2, 6-sialyltransferase (C) Rabbit IgG Affinity Purify Precoated plate 1 96Well x 1
- 2 Labeled antibody Conc.
- (30 X) HRP conjugated Anti-Rat α2, 6-sialyltransferase (E41) Rabbit IgG Fab' Affinity purify 0.4mL x 1 Standard : Recombinant Rat α2, 6-sialyltransferase (E41 Form) 0.5mL x 2 30mL x 1 EIA buffer Solution for Labeled antibody Chromogen : TMB solution 12mL x 1 5 6 15mL x 1 Stop solution 1N H₂SO₄ 12mL x 1 Wash buffer Conc. : (40 X) 0.05% Tween20 in phosphate buffer 8 50mL x 1

OPERATION MANUAL

1. Materials needed but not supplied

 Plate reader (450nm) 	 Micropipette and tip
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- · Graduated cylinder and beaker · Deionized water Incubator(37°C±1°C)
 - · Graph paper (log/log)
- Paper towel
- Tube for dilution of Standard · Washing bottle for precoated plate
- · Disposable test tube for "2, Labeled antibody Conc." and "6, Chromogen"

2. Preparation

- 1) Preparation of wash buffer "8, Wash buffer Conc." is a concentrated (40 X) buffer. Adjust the temperature of "8, Wash buffer Conc." to room temperature and then, mix it gently and completely before use. Dilute 50mL of "8, Wash buffer Conc." with 1,950mL of deionized water and mix it. This is the wash buffer for use. This prepared wash buffer shall be stored in refrigerator and used within 2 weeks after dilution.
- Preparation of Labeled antibody
 - "2, Labeled antibody Conc." is a concentrated (30 X). Dilute "2, Labeled antibody Conc." with "5, Solution for Labeled antibody" in 30 times according to required quantity into a disposable test tube. Use this resulting solution as Labeled antibody. Example)

In case you use one slit (8 well), the required quantity of Labeled antibody is 800 μ L. (Dilute 30 μ L of "2, Labeled antibody Conc." with 870 μ L of "5, Solution for Labeled antibody" and mix it. And use the resulting solution by 100 µL in each well.)

This operation should be done just before the application of Labeled antibody. The remaining "2, Labeled antibody Conc." should be stored at 4°C in firmly

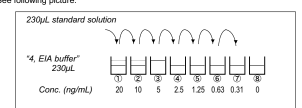
sealed vial.

- Preparation of Standard Put just $\underline{0.5 \text{ mL}}$ of "4, EIA buffer" to the vial of "3, Standard" and mix it gently and completely. This solution is 40 ng/mL Rat α 2, 6-sialyltransferase (E41 Form) standard.
- Dilution of Standard 4)
- Prepare 8 tubes for dilution of "3, Standard". Put 230 µL each of "4, EIA buffer" into the tube.

Specify the following concentration of each tube.

Tube-1	20 ng/mL	
Tube-2	10 ng/mL	
Tube-3	5 ng/mL	
Tube-4	2.5 ng/mL	
Tube-5	1.25 ng/mL	
Tube-6	0.63 ng/mL	
Tube-7	0.31 ng/mL	
Tube-8	0 ng/mL	(Test Sample Blank)

Put 230 µL of Standard solution into tube-1 and mix it gently. Then, put 230 µL of tube-1 mixture into tube-2. Dilute two times standard solution in series to set up 7 points of diluted standard between 20 ng/mL and 0.31 ng/mL. Tube-8 is the test sample blank as 0 ng/mL See following picture.



Dilution of test sample 5)

The test sample should be diluted with "4, EIA buffer" more than 4 times If the concentration of Rat α2, 6-sialyltransferase (E41 Form) in samples may not be estimated in advance, the pre-assay with several different dilutions will be recommended to determine the proper dilution of samples.

3. Measurement procedure

All reagents shall be brought to room temperature approximately 30 minutes before use. Then mix it gently and completely before use. Make sure of no change in quality of the reagents. Standard of simultaneously with the measurement of test samples. Standard curve shall be prepared

	Test Sample	Standard	Test Sample Blank	Reagent Blank
Reagents	Test sample 100 μL	Diluted standard (Tube 1~7) 100 μL	EIA buffer (Tube-8) 100 μL	EIA buffer 100 μL
	Incubation for	60 minutes at 37	°C with plate lid	
		Washing 7 times	5	
Labeled Antibody	100 µL	100 µL	100 µL	-
	Incubation for 30 minutes at 37°C with plate lid			
	Washing 9 times			
Chromogen	100 µL	100 µL	100 µL	100 µL
Incubation for 30 minutes at room temperature (shielded)				
Stop solution	100 µL	100 µL	100 µL	100 µL
Read the plate at 450nm against a Reagent Blank within 30 minutes after addition of Stop solution.				

- 1) Determine wells for reagent blank. Put 100 μL each of "4, EIA buffer" into the wells
- 2) Determine wells for test sample blank, test sample and diluted standard. Then, put 100 µL each of test sample blank (tube-8), test sample and dilutions of standard (tube-1-7) into the appropriate wells.
- 3) Incubate the precoated plate for 60 minutes at 37°C after covering it with plate lid.
- Wash each well of the precoated plate vigorously with wash buffer using the washing bottle. Then, fill each well with wash buffer and leave the precoated plate laid for 15 30 seconds. Remove wash buffer completely from the precoated plate by snapping. This procedure must be repeated more than 7 times. Then, remove the remaining liquid from all wells completely by snapping the precoated plate onto paper towel. 4)
 - In case of using a plate washer, after 4 times washing with plate washer, washing with above washing bottle must be repeated 3 times.
- Pipette 100 μL of labeled antibody solution into the wells of test samples, diluted standard and test sample blank. 5) 6)
- Incubate the precoated plate for 30 minutes at 37°C after covering it with plate lid.
- Wash the precoated plate 9 times in the same manner as 4).
- "6, Chromogen" should be taken the required quantity into a disposable test tube. Then, pipette 100 μL from the test tube into the wells. Please do not 8) return the rest of test tube to "6, Chromogen" bottle due to avoid contamination.
- Incubate the precoated plate for 30 minutes at room temperature in the dark. 9)
- The liquid will turn blue by the addition of "6, Chromogen". Pipette 100 μ L of "7, Stop solution" into the wells. Mix the liquid by tapping the side of precoated plate. The liquid will turn yellow by the addition of "7, Stop 10) ; solution
- Remove any dirt or drop of water on the bottom of the precoated plate and 11) confirm there is no bubble on the surface of the liquid. Then, run the plate reader and conduct measurement at 450nm against a reagent blank. The measurement shall be done within 30 minutes after the addition of "7, Stop solution".

SPECIAL ATTENTION

Test samples should be measured soon after the collection. For the storage 1) of test samples, store them frozen and do not repeat freeze/thaw cycles. Thaw the test samples at low temperature and mix them completely before measurement

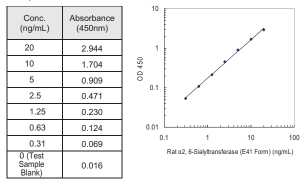


- Test samples should be diluted with "4, EIA buffer", if the need arises
- Duplicate measurement of test samples and standard is recommended. 3)
- 4) Use test samples in neutral pH range. The contaminations of organic solvent may affect the measurement.
- 5) Use only wash buffer contained in this kit for washing the precoated plate.
- Insufficient washing may lead to the failure in measurement. Remove the wash buffer completely by tapping the precoated plate on paper 6)
- towel. Do not wipe wells with paper towel.
- "6, Chromogen" should be stored in the dark due to its sensitivity against light. "6, Chromogen" should be avoided contact with metals. 7)
- Measurement should be done within 30 minutes after addition of "7, Stop 8) solution"

CALCULATION OF TEST RESULT

Subtract the absorbance of test sample blank from all data, including standards and unknown samples before plotting. Plot the subtracted absorbance of the standards against the standard concentration on log-log graph paper. Draw the best smooth curve through these points to construct the standard curve. Read the concentration for unknown samples from the standard curve.

Example of standard curve



The typical standard curve is shown above. This curve can not be used to derive test results. Please run a standard curve for each assay.

PERFORMANCE CHARACTERISTICS

1. Titer Assay (Samples with standard added are used.)

Specimen	Titer (X)	Measurement Value (ng/mL)	Theoretical Value (ng/mL)	%
10% FCS	8	4.63	5.00	92.6
added	16	2.42	2.50	96.8
RPMI-1640	32	1.21	1.25	96.8
Rat Serum (Wistar)	16	8.65	9.56	90.5
	32	4.68	4.91	95.3
	64	2.54	2.63	96.8
Rat Plasma (EDTA) (Wistar)	16	7.39	8.22	89.9
	32	4.15	4.55	91.2
	64	2.22	2.41	92.3

2. Added Recovery Assay

Specimen	Theoretical Value (ng/mL)	Measurement Value (ng/mL)	%
10% FCS added RPMI-1640 (x8)	5.00	4.78	95.6
	2.50	2.35	94.0
	1.25	1.11	88.8
Rat Serum (Wistar) (x16)	6.36	6.50	100.6
	6.04	6.27	100.5
	5.89	6.10	101.6
Rat Plasma (EDTA) (Wistar) (x16)	5.64	5.67	100.6
	5.32	5.35	100.5
	5.17	5.25	101.6

3. Intra- Assay

Measurement Value (ng/mL)	SD value	CV value (%)	n
7.43	0.23	3.1	24
1.85	0.08	4.3	24
0.45	0.02	4.4	24

4. Inter - Assay

Measurement Value (ng/mL)	SD value	CV value (%)	n
7.60	0.32	4.2	38
1.89	0.10	5.3	38
0.46	0.02	4.7	38

5. Sensitivity

0.02 na/mL

The sensitivity for this kit was determined using the guidelines under the National Committee for Clinical Laboratory Standards (NCCLS) Evaluation Protocols. (National Committee for Clinical Laboratory Standards Evaluation Protocols, SC1, (1989) Villanova, PA: NCCLS.

PRECAUTION FOR INTENDED USE AND/OR HANDLING

- 1. All reagents should be stored at 2 8 °C. All reagents shall be brought to room temperature approximately 30 minutes before use.
- 2
- "3, Standard" is lyophilized products. Be careful to open this vial. "7, Stop solution" is a strong acid substance. Therefore, be careful not to have your skin and clothes contact "7, Stop solution" and pay attention to the 3. disposal of "7, Stop solution".
- Dispose used materials after rinsing them with large quantity of water. Precipitation may occur in "2, Labeled antibody Conc.", however, there is no 5. problem in the performance.
- 6. Wash hands after handling reagents.
- Do not mix the reagents with the reagents from a different lot or different kit. Do not use the expired reagents. 8.
- 9. This kit is for research purpose only. Do not use for clinical diagnosis.

STORAGE AND THE TERM OF VALIDITY

Storage Condition : 2 - 8 °C The expiry date is specified on outer box.

REFERENCE

- 1. Kitazume S. Saido TC. Hashimoto Y. Identification of a novel substrate of Alzheimer's beta-secretase: beta-secretase dependent cleavage of alpha 2,6 sialyttransferase. Seikagaku. 2002 Sep; 74(9): 1180-1183. Kitazume-Kawaguchi S, Dohmae N, Takio K, Tsuji S, Colley KJ.
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Version 1.2





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