



For research use only

Urinary Diacetylspermidine ELISA Kit

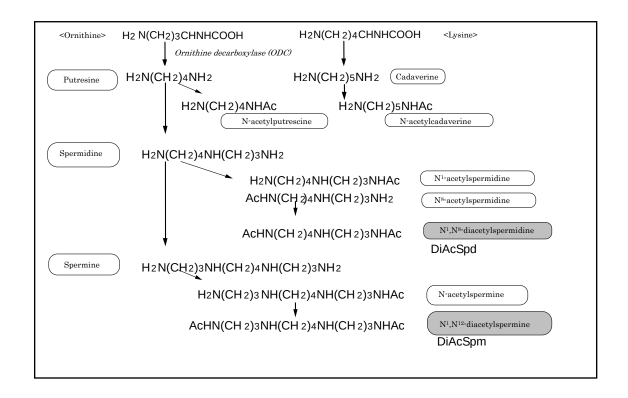
Polyamines are generally believed to function both in protein synthesis and DNA synthesis leading to control cell proliferation. In 1971, Russel firstly reported that total amount of urinary polyamines elevated in cancer patients. And quantitative kits of urinary polyamines were already developed and utilized as a general biochemical examination.

Recently two diacetyl-derivatives, N1, N12-diacetylspermine and N1, N8-diacetylspermidine, were found to be excreted in urine and form 0.6% and 1.4% of total polyamines respectively.

Comparing urine of diseased person with urine of healthy person, some reports suggested the possibility that diacetyl-derivatives correlate to the status of disease more closely than total amount of polyamines.

Our kit is convenient to quantify amount of urinary diacetylspermidine by using ELISA method. This kit is only for research use, not for diagnosis.

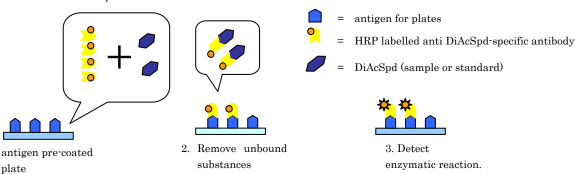
- Highly sensitive and specific
- •Strip type well, antigen pre-coated microplate
- Assay range: 9.375∼600 nM







1. Incubate with sample.



[Kit Contents]

(1)	Antigen coated microtiter plate, 96 wells	1 plate	
(2)	Diacetylspermidine standard	$250~\mu L$	$\times 2$
(3)	Reference standard and sample diluent	$15~\mathrm{mL}$	× 1
(4)	Antibody diluent	20 mL	× 1
(5)	HRP- anti Diacetylspermidine antibody concentrate (×100)	$60~\mu L$	× 1
(6)	OPD (o-phenylendiamine) tablets	2 tablets	
(7)	Substrate solution	30 mL	× 1
(8)	Stop solution	$15~\mathrm{mL}$	× 1
(9)	Wash buffer concentrate (×20)	30 mL	× 1
(10)	Dilution plate	1 plate	

[Equipments to be supplied by the user]

- (1) A microplate reader
- (2) Micropipets
- (3) A microplate washer

[Assay Method]

- (1) Preparation of working solution
 - (1) Wash solution

Make sure that wash buffer concentrate does not contain any crystallized material prior to use. Working solution is prepared by dilution 30 mL of wash buffer concentrate with 570 mL of distilled deionized water. For convenience this solution can be kept at 2-8 °C up to 14 days.

② Diacetylspermidine Standard

Prepare 6 standards by serial dilution of diacetyl spermidine standard concentrate (600 nM) as follows

We recommend a polypropylene tube for preparation of standard solution. A glass or polystyrene tube may cause non-specific adsorption of diacetylspermidine, so that you may not get reliable results.

		300	150	75.0	37.5	18.75	9.375	(nM)
Standard solution 600 nM	(μL)	100 ~	100	100	100	100	100	
Reference standard and sample diluent	(μL)	100 _	ر 100 مرا	ر 100 م	ر ₁₀₀	ر ₁₀₀ ک	100	



③ HRP-anti Diacetylspermidine antibody (×100) Dilute 40 µL of Anti Diacetylspermidine antibody concentrate (×100) with 4 mL of Dilution solution for 96 well reaction. Diluted antibody should not be stored.

4 Coloring solution

Add one OPD tablet to 13 mL of Substrate buffer to reconstitute the coloring solution just before use. This solution should not be stored.

(2) Preparation of urine sample

- ① Collect urine in sampling tube on demand. Add 0.1% Na₂N₃ at final concentration.
- 2 After centrifugation at 1500 rpm for 5 min, dilute the resulted supernatant over 4 times with Sample diluent.
- Measure the amount of creatinine in remaining diluted supernatant for compensation.

*Prepared urine sample should be kept below -30 °C if necessary.

(3) Assay procedure

① Pre-reaction

Prepare standard control wells containing 70 µL of anti Diacetylspermidine antibody solution and 70 µL of 6 standards (600, 300, 150, 75.0, 37.5, 18.75, 9.375 nM) in dilution plate. Likewise prepare experimental wells containing 70 µL of anti Diacetylspermidine antibody solution and 70 µL of prepared urinary sample in the same plate. After settlement, incubate at room temperature for 30 minutes.

* Above reaction volumes can be applied for double measurements of primary reaction. In the case of single measurement, reduce to 40 µL of each solution.

Preparation of reaction plate

- ②-1 Add wash solution 300 µL to each well and wait another 30 minutes.
- 2-2 Discard the wash solution from the wells completely and wash with 300 µL wash solution.

Repeat this step another 2 times

Primary reaction

- Apply 50 uL/well × 2 (In the case of measuring double wells) pre-reaction solution (See ①) and incubate for 1 hour.
- After the incubation, discard the reaction solution and wash with 300 µL wash solution. Repeat this step another 2 times.

Apply 100 µL Coloring solution to each well and incubate for 10 minutes at room temperature.

⑤ Stop reaction

Apply 100 µl of Stop solution to stop the enzymatic reaction

Read absorbance

Read absorbance of 490 nm or 492 nm with a microplate reader.

Measure concentration

Measure the Diacetylspermidine concentration using standard curve.

- If actual measurements of sample exceed over 600 nM, dilute those urine samples to evaluate within the range of 18.75~600 nM.
- Concentration of diacetylspermine needs to be calculated from actual measurements by





consideration of dilution ratio.

* For the comparison of clinical data, actual measurements need compensation with the concentration of urinary creatinine (nmoL/g · cre).

Calculation method of clinical data

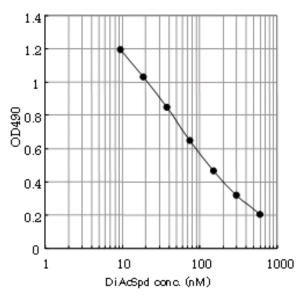
The concentrations of excrement in urine depend on the urine volume which changes by the amount of water take or environment in both renal disease patients and healthy persons. In general, the measurements of urinary excrement need to be corrected by the urinary creatinine because the amount depends on that of muscle and the measurement is able to convert into the amount per 1g creatinine. Therefore, correct actual measurement of Diacetylspermidine concentration (nM) in urine with creatinine concentration (mg/dL), as follows,

Diacetylspermidine concentration (nM)

Data correction : nmoL/g · cre = $\frac{}{\text{Creatinine concentration (mg/dL)}} \times 100$

(Standard curve)





[Reproducibility]

Domain of standard curve: 9.375~600 nM

Minimum measurement range for detection: 18.75 nM

Minimum dilution number of urine sample : $\times 2$ Minimum sensitivity for detection : 37.5 nM

Within-run (n=20, 2 concentration) : CV(%) = 5.76, 2.58 Between-run (n=20, 2 concentration) : CV(%) = 6.76, 4.25

Recovery test: In the recovery study, recoveries between 96% and 108% were obtained for 2, 4, 8 times dilutions of the sample urine

Coexistent substance : No influence to Hemoglobin 4500 mg/dL \cdot Bilirubin 180 mg/dL \cdot Glucose 1000 mg/dL \cdot Ascorbic acid 40 mg/dL





[Usage notes]

- ① The Reagents should be stored at recommended temperature, -30 °C.
- ② Do not use the reagents which is expired the date of usage.
- ③ Urine sample should be diluted more than 4 times with Dilution solution.
- ④ Do not leave the standard and Antibody for long time under room temperature.
- ⑤ The glassware for making coloring solution should be clean.
- 6 Since OPD (o-phenylendiamine) is harmful, handle with care.
- © Since Stop solution 1 M H₂SO₄, is strong acid, handle with care.
- The kit is constructed with well-adjusted combination in each lot. Replaced combination among different lots may cause unexpected results.
- 9 This kit is only for research use. Do not use for medicinal or any other purposes.
- When using the reagents, take care to avoid them from touching to skin, mucous membrane, clothes, and getting into eyes.
- ① If the reagents happen to get into eyes or mouth, wash out them and consult a doctor if you need.
- ② After using the kit, wash your hands very carefully.
- If you find that the packages of the reagents are broken or something wrong, do not use them.
- When you store the reagents, make sure to avoid them from vaporizing, falling down.
- (5) After using the reagents, the packages should be discarded under the established rule.
- (6) We do not guarantee the quality of the packages and accompaniments if not used according this direction.

[Storage]

All reagents: -30 °C





[References]

- $1) \ \ {\hbox{Russell DH:Increased polyamine concentration in the urine of human cancer patients}}.$
 - Nature New Biol 233:144-145,1971
- 2) Hiramatsu K, Sugimoto M, Kamei S, Hoshino M, Kinoshita K, Iwasaki K, and Kawakita M:Determination of amounts of polyamines excreted in urine;demonstration of N1,N8-diacetylspermidine and N1,N12diacetylspermine as components commonly occurring in normal human urine.
 - J. Biochem., 117:107-112,1995
- 3) Sugimoto M, Hiramatsu K, Kamei S, Kinoshita K, Hoshino M, Iwasaki K, and Kawakita M: Significance of urinary N1,N8-diacetylspermidine and N1,N12- diacetylspermine as indicators of neoplastic diseases.
 - J. Cancer Res. Clin. Oncol.,121:317-319,1995
- 4) Hiramatsu K, Sugimoto M, Kamei S, Hoshino M, Kinoshita K, Iwasaki K, and Kawakita M:Diagnostic and prognostic usefulness of N1,N8- diacetylspermidine and N1,N12- diacetylspermine in urine as novel markers of malignancy.
 - J. Cancer Res. Clin. Oncol., 123:539-545,1997



This product is generated from GANP® mice.

Manufacturer

Previous manufacturer



Medicinal Chemistry Pharmaceutical Co., Ltd.

Trans Genic Inc.

Kobe Research Institute

7-1-14 Minatojimaminami-machi, Chuo-ku, Kobe, Japan 650-0047

Telephone: +81-78-945-7075 FAX:+81-78-306-0694

URL:https://soyaku.co.jp/english/ tech-kobe@soyaku.co.jp