

# TriCat ELISA



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**DRG International, Inc**., USA 841 Mountain Ave., Springfield, NJ 07081 Phone: (973) 564-7555, Fax: (973) 564-7556 Website: www.drg-international.com E-mail: corp@drg-international.com Please use only the valid version of the Instructions for Use provided with the kit. Verwenden Sie nur die jeweils gültige, im Testkit enthaltene, Gebrauchsanweisung. Si prega di usare la versione valida delle istruzioni per l'uso a disposizione con il kit. Por favor, use sólo la versión válida de las instrucciones de uso que se suministran con el kit. Utilisez seulement la version valide des Instructions d'utilisation fournies avec le kit. Por favor, usar a versão válida das instruções de utilização fornecidas com o kit.

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SYMBOLS USED
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#### 1 INTENDED USE

Manual and automated enzyme immunoassay for the *in-vitro-diagnostic* quantitative determination of adrenalin (epinephrine), noradrenalin (norepinephrine) and dopamine in human plasma and urine.

#### 2 SUMMARY AND EXPLANATION

The catecholamines adrenalin, noradrenalin and dopamine are synthesized in the adrenal medulla, the sympathetic nervous system and in the brain. They influence virtually all tissues and are involved together with other hormonal and neuronal systems in the regulation of a wide variety of physiological processes.

As catecholamines and their metabolites metanephrine and normetanephrine are secreted in increasing amounts in a number of diseases, they may be used for diagnostic purposes.

In this context, diagnosis and the follow-up of tumor diseases of the nervous system are of special importance. This applies primarily to the pheochromocytoma, but also the neuroblastoma and the ganglioneuroma.

Because of the extraction step at the beginning of the assay, the customer is able to use all kinds of animal species material. It works for rats, mice and others. The chemical structure of the catecholamines is identical in all animals.

#### 3 TEST PRINCIPLE

Solid phase enzyme-linked immunosorbent assay (ELISA) based on the sandwich principle. The wells are coated with a goat anti rabbit antibody. The added liquid antibody, directed towards an epitope of an antigen molecule binds to the plate within the incubation time. The antigen of the sample is incubated in the coated well with enzyme conjugated second antibody (E-Ab), directed towards a different region of the antigen molecule. After the substrate reaction the intensity of the developed color is proportional to the amount of the antigen. Results of samples can be determined directly using the standard curve.

#### 4 WARNINGS AND PRECAUTIONS

- 1. For *in-vitro diagnostic* use only. For professional use only.
- 2. Before starting the assay, read the instructions completely and carefully. Use the valid version of the package insert provided with the kit. Be sure that everything is understood.
- 3. In case of severe damage of the kit package please contact DRG or your supplier in written form, latest one week after receiving the kit. Do not use damaged components in test runs, but keep safe for complaint related issues.
- 4. Obey lot number and expiry date. Do not mix reagents of different lots. Do not use expired reagents.
- 5. Follow good laboratory practice and safety guidelines. Wear lab coats, disposable latex gloves and protective glasses where necessary.
- 6. Reagents of this kit containing hazardous material may cause eye and skin irritations. See MATERIALS SUPPLIED and labels for details. Material Safety Data Sheets for this product are available upon request.
- 7. Chemicals and prepared or used reagents have to be treated as hazardous waste according to national biohazard and safety guidelines or regulations.
- 8. The cleaning staff should be guided by the professionals regarding potential hazards and handling.
- 9. Avoid contact with Stop solution. It may cause skin irritations and burns.
- 10. All reagents of this kit containing human serum or plasma have been tested and were found negative for anti-HIV I/II, HBsAg and anti-HCV. However, a presence of these or other infectious agents cannot be excluded absolutely. For this reason, reagents should be treated as potential biohazards in use and for disposal.

#### 5 STORAGE AND STABILITY

The kit is shipped at ambient temperature and should be stored at 2 °C - 8 °C. Keep away from heat or direct sunlight. The storage and stability of specimens and prepared reagents is stated in the corresponding chapters.

The microtiter strips are stable up to the indicated expiry after the kit is broken. Make sure that the broken bag is tightly closed when stored at 2 °C - 8 °C.

#### 6 SPECIMEN COLLECTION AND STORAGE



The in-vivo catecholamine and metanephrines release is influenced by several foods and drugs. Vitamin B, coffee and bananas, alpha-methyldopa, MAO and COMT inhibitors as well as medications related to hypertension should be discontinued for at least 72 h prior to specimen collection.

#### Plasma (EDTA)



The blood sample should be stored at 2 °C - 8 °C until centrifuged to separate the plasma within 2 hours after blood collection.

The usual precautions for venipuncture should be observed. It is important to preserve the chemical integrity of a blood specimen from the moment it is collected until it is assayed. Do not use grossly hemolytic, icteric or grossly lipemic specimens. Samples appearing turbid should be centrifuged before testing to remove any particulate material.

Storage:	2 °C - 8 °C	≤ -20 °C (Aliquots)	Keep away from heat or direct sunlight.		
Stability:	6 hours	1 month	Avoid repeated freeze-thaw cycles. Ship samples frozen.		

#### Urine

It is possible to use spontaneous as well as 24 h urine. The total volume of urine excreted during a 24 h period should be collected and mixed in a single bottle containing 10-15 mL of 6 N HCl as preservative. Determine total volume for calculation of results. **Mix and centrifuge samples before use in the assay.** 

Storage:	≤ -20 °C (Aliquots)	Keep away from heat or direct sunlight.
Stability:	6 months	Avoid repeated freeze-thaw cycles.

#### 7 MATERIALS SUPPLIED

The reagents provided with this kit are sufficient for 96 extractions in single determinations in the sample preparation (extraction): 88 patient samples, 6 standards and 2 controls. Each extract is sufficient for a single determination for adrenalin, noradrenalin and dopamine immunoassay.

The microtiter plate can be used for all three analytes: Adrenalin, Noradrenalin and Dopamine.

Quantity	Symbol	Component
3 x 12 x 8	MTP	Microtiter Plate
		Break apart strips. Coated with anti-rabbit IgG (goat, polyclonal).
1 x 6 x 2.5 mL CAL A-F		Standard A-F        Adrenalin: 0; 1.5; 5.0; 15; 50; 150 ng/mL (0; 8; 27; 82; 273; 819 nmol/L)        Noradrenalin: 0; 5.0; 15; 50; 150; 500 ng/mL (0; 30; 89; 296; 887; 2955 nmol/L)        Dopamine: 0; 60; 180; 585; 2300; 11470 ng/mL (0; 392; 1175; 3819; 15014; 74876 nmol/L)        Ready to use. Contains: [-] Adrenalin, [-] Noradrenalin, [-] Dopamine (biologically active), and 0.1 M HCI.
1 x 2 x 2.5 mL	CONTROL 1+2	Control 1+2 Ready to use. Contains: [-] Adrenalin, [-] Noradrenalin, [-] Dopamine (biologically active), 0.1 M HCI. Exact concentrations see vial labels or QC certficate.
3 x 400 µL	ENZCONJ CONC	Enzyme Conjugate Concentrate (50x) Contains: Streptavidin alkaline phosphatase, Tris buffer, stabilizers.
4 x	EXTRPLATE	<b>Extraction Plate</b> (Macrotiter Plate) 24 wells each. Coated with boronate affinity gel.
2 x 60 mL	EXTRBUF	Extraction Buffer Pink colored. Ready to use. Contains: 0.016 % NaN <sub>3</sub> .
6 x 1.25 mL	COMTLYO	<b>COMT lyophilized</b> Contains: Catechol-O-methyltransferase (porcine liver), NaN <sub>3</sub> .
6 x 1.25 mL	COENZ	<b>Coenzyme Solution</b> Ready to use. Contains: S-Adenosyl-L-Methionine, stabilizers.
3 x 3 mL	ENZBUF	<b>Enzyme Buffer</b> Ready to use. Contains: Tris buffer, HCI, stabilizers.
1 x 100 mL	RELEASEBUF	Release Buffer Yellow Colored. Ready to use. Contains: 0.1 M HCl, indicator.

Quantity	Symbol	Component
2 x 3.0 mL	ACYLREAG	Acylation Reagent Ready to use. Contains: dimethylformamide, Ethanol. Caution! Toxic. Highly flammable.
2 x 100 mL	WASHBUF CONC	Wash Buffer Concentrate (10x) Contains: Tris buffer, HCl, Tween, 0.2 % NaN <sub>3</sub> .
2 x 2 mL	COMT ADD	<b>COMT Additive</b> Contains: human plasma, stabilizers, 0.01 % Thimerosal.
1 x 8.0 mL	ANTISERUM AD	Adrenalin Antiserum Green colored. Ready to use. Contains: antibodies against Adrenalin (rabbit), Buffer, stabilizers.
1 x 8.0 mL	ANTISERUM NAD	<b>Noradrenalin Antiserum</b> Blue colored. Ready to use. Contains: antibodies against Noradrenalin (rabbit), Buffer, stabilizers.
1 x 8.0 mL	ANTISERUM DO	<b>Dopamine Antiserum</b> Violet colored Ready to use. Contains: antibodies against Dopamine (rabbit), Buffer, stabilizers.
3 x 25 mL	PNPP SUBS	<b>PNPP Substrate Solution</b> Ready to use. Contains: p-nitrophenyl phosphate (PNPP).
3 x 15 mL	PNPP STOP	PNPP Stop Solution Ready to use. Contains: 1 M NaOH, 0.25 M EDTA.
9 x	FOIL	Adhesive Foil

# 8 MATERIALS REQUIRED BUT NOT SUPPLIED

- 1. Micropipettes (Multipette Eppendorf or similar devices, < 3% CV). Volume: 10; 10-100; 100-1000 μL
- 2. Orbital shaker (200-900 rpm) (e.g. EAS 2/4, SLT)
- 3. Vortex mixer
- 4. 8-Channel Micropipettor with reagent reservoirs
- 5. Wash bottle, automated or semi-automated microtiter plate washing system
- 6. Microtiter plate reader capable of reading absorbance at 405 nm (reference wavelength 600-650 nm)
- 7. Bidistilled or deionised water
- 8. Paper towels, pipette tips and timer
- 9. Disposable tubes for sample dilution
- 10. 0.1 M HCl, for sample dilution (Urine)

#### 9 PROCEDURE NOTES

- 1. Any improper handling of samples or modification of the test procedure may influence the results. The indicated pipetting volumes, incubation times, temperatures and pretreatment steps have to be performed strictly according to the instructions. Use calibrated pipettes and devices only.
- 2. Once the test has been started, all steps should be completed without interruption. Make sure that required reagents, materials and devices are prepared ready at the appropriate time. Allow all reagents and specimens to reach room temperature (18 °C 25 °C) and gently swirl each vial of liquid reagent and sample before use. Mix reagents without foaming.
- 3. Avoid contamination of reagents, pipettes and wells/tubes. Use new disposable plastic pipette tips for each component and specimen. Do not interchange caps. Always cap not used vials. Do not reuse wells/tubes or reagents.
- 4. It is advised to determine samples in duplicate to be able to identify potential pipetting errors.
- 5. Use a pipetting scheme to verify an appropriate plate layout. A pipetting scheme covering both sample pretreatment and assay is available upon request.
- 6. Incubation time affects results. All wells should be handled in the same order and time sequences. It is recommended to use an 8-channel Micropipettor for pipetting of solutions in all wells.
- 7. Microplate washing is important. Improperly washed wells will give erroneous results. It is recommended to use a multichannel pipette or an automatic microplate washing system. Do not allow the wells to dry between incubations. Do not scratch coated wells during rinsing and aspiration. Rinse and fill all reagents with care. While rinsing, check that all wells are filled evenly with Wash Buffer, and that there are no residues in the wells.
- 8. Humidity affects the coated wells/tubes. Do not open the pouch until it reaches room temperature. Unused wells/tubes should be returned immediately to the resealed pouch including the desiccant.

#### 10 MANUAL PROCEDURE

#### 10.1 PRE-TEST SETUP INSTRUCTIONS

The contents of the kit for 3 x 96 determinations can be divided into 2 separate runs.

Visible amounts of gel can be separated from surface of extraction plate during extraction.



This has no influence on test results.

Air contamination by peroxygen containing disinfectants for cleaning of surfaces or equipment used as powder or as solutions, e.g. VIRKON<sup>®</sup> must be avoided in any case. They will strongly disturb assay performance. VIRKON<sup>®</sup> is a trademark of DuPont.

#### 10.1.1 Dilution of Samples

Samples suspected to contain concentrations above the highest standard have to be diluted as follows:

Sample	to be diluted	with	Remarks
Plasma	> highest standard	bidist. water	prior to extraction step
Urine	> highest standard	0.1 N HCI	prior to extraction step

#### 10.1.2 Extraction of Samples, Standards and Controls (Extraction Plate) (manual version)

- 1. Pipette **20 μL** of each **Standard, Control and urine sample** and **500 μL** of each **plasma sample** into the respective wells of the extraction plate. Add **500 μL** of **bidist. water** to all wells except for the **plasma samples** to correct differences of volumes.
- 2. Pipette **1000 µL** of **Extraction Buffer** into each well.
- Cover plate with adhesive foil. Extract 30 min at RT (18 °C 25 °C) on an orbital shaker (600–900 rpm).
  During extraction the surface of the liquid should wet the adhesive foil, but the liquid level should not exceed 2/3 of the well. Splashing does not affect results.
- 4. Remove adhesive foil. Immediately empty plate and eliminate residual fluid on a paper towel.
- 5. Pipette 2 mL of bidist water into each well.
- 6. Cover plate with new adhesive foil.
  - Shake 5 min at RT (18 °C 25 °C) on an orbital shaker (600–900 rpm). Splashing does not affect results.
- 7. Remove adhesive foil. Immediately empty plate and eliminate residual fluid on a paper towel. Remove fluid completely.
- 8. Pipette **150 μL** of **Extraction Buffer** into each well. To each well add **50 μL** of **Acylation Reagent**. Mix immediately after pipetting.
- 9. Extract 20 min at RT (18 °C 25 °C) (without adhesive foil) on an orbital shaker (400–600 rpm).
- 10. Immediately empty plate and eliminate residual fluid on a paper towel. Remove fluid completely.
- 11. Pipette 2 mL of bidist. water into each well.
- 12. Cover plate with new adhesive foil. **Shake 5 min** at **RT (18 °C 25 °C)** on an orbital shaker (600-900 rpm). Splashing does not affect results.
- 13. Remove adhesive foil. Immediately empty plate and eliminate residual fluid on a paper towel. Remove fluid completely.
- 14. Pipette 300 µL of Release Buffer into each well.

15. Shake 30 min at RT (18 °C - 25 °C) (without adhesive foil) on an orbital shaker (400–600 rpm).

Prepared samples should be assayed the same day. If this is not possible, you can store the extraction plate covered with adhesive foil at 2 °C - 8 °C over night.

#### Important for measurement of **Dopamine**

Dilution of extracted **standard**, **controls** must be performed prior to pipetting into wells of Microtiterplate in extra tubes. Predilute **urine samples** as well.

Therefore, dilute all extracted Standards, Controls and urine samples 1:51 with Release Buffer in disposable tubes. (i.e. 10  $\mu$ L extracted samples + 500  $\mu$ L Release Buffer).

Extracted plasma samples does not require this predilution.

# 10.1.3 Preparation of concentrated components

The volumes stated below are for one run with 3 x 6 strips (3 x 48 determinations)

Dilute / dissolve	Component	with	Diluent	Relation	Remarks	Storage	Stability	
75 mL	WASHBUF CONC	675 mL	bidist. water	1:10	Mix vigorously.	2 °C - 8 °C	4 weeks	
300 µL	ENZCONJ CONC	15 mL	WASHBUF (diluted)	1:51	Prepare freshly and use only once. Mix without foaming.	18 °C - 25 °C	5 hours	

## 10.2 TEST PROCEDURE (manual version)

#### 10.2.1 Preparation of COMT Enzyme Solution

#### **I** The COMT Enzyme Solution should be freshly prepared directly before use.

Dissolve each kit component of lyophilized COMT in 1.25 mL bidist. water and mix the dissolved COMT.\*

Then **pipette 1.25 mL** of **Coenzyme Solution** followed by **1.25 mL** of **Enzyme Buffer** and **0.40 mL COMT Additive** to the mixed COMT vials to give a final volume of 4.15 mL of COMT Enzyme Solution per vial.

Pool three (3) vials for 48 determinations of adrenalin and 48 determinations of noradrenalin and 48 determinations of dopamine. Solution may be turbid. Mix without foaming. The COMT solution is stable at room temperature for 1 hour.

\* If only an aliquot of the COMT solution is needed, the rest of the COMT solution should be frozen immediately in aliquots at -20 °C. The COMT solution is stable under these conditions for 1-2 months.

#### 10.2.2 Enzymatic Derivatization of Samples, Standards and Controls (Microtiter Plate)

If pipetting with *positive displacement*, give the residual fluid from the tip of the pipette back to the corresponding wells of the extraction plate, otherwise the extracts may not be sufficient for the determination of the other analytes.

It is useful to hold the extraction plate in a sloping position.

Before use of the Microtiter plates, define and label the wells for Adrenalin, Noradrenalin and Dopamine.

# 10.2.2.a For research use of tissue homogenates and cell culture supernatants a general recommendation can be given:

Working with cell culture supernatants depends on the matrix as well as concentrations expected:

According to the urine protocol (extraction of at least 20 µL supernatant) a sensitivity for adrenalin of 0.3 ng/mL, for noradrenaline of 0.6 ng/mL and for dopamine of 5 ng/mL for diluted sample can be expected. In case of a matrix with addition of serum (FCS), plasma protocol (extraction of 500 µL supernatant) can be used with the sensitivities corresponding to the plasma protocol (see 16. PERFORMANCE).

For tissue homogenates no perchloric acid should be used for homogenization. For further details ask DRG.

#### 10.2.2.b Adrenalin for urine and plasma

- 1. Pipette **75** µL of freshly prepared **COMT Enzyme Solution** into each well of the **Microtiter Plate**. Shake plate briefly.
- 2. Pipette **100 µL** of each <u>extracted</u> **Standard, Control and sample** into the respective wells. During this step hold the pipette tips directly into the COMT solution. Color changes to pink. Shake briefly.
- 3. Pipette 50 µL of Adrenalin Antiserum (green colored) into each well.
- 4. Cover plate with adhesive foil. Incubate 120 min at RT (18 °C 25 °C) on an orbital shaker (400-600 rpm).

# 10.2.2.c Noradrenalin for urine and plasma

- 1. Pipette **25 μL** of freshly prepared **COMT Enzyme Solution** into each well of the **Microtiter Plate**. Shake plate briefly.
- 2. Pipette **25** µL of each <u>extracted</u> **Standard, Control and sample** into the respective wells. During this step hold the pipette tips directly into the COMT solution. Color changes to pink. Shake plate briefly.
- 3. Pipette **50 µL** of **Noradrenalin Antiserum** (blue colored) into each well.
- 4. Cover plate with adhesive foil. Incubate 120 min at RT (18 °C 25 °C) on an orbital shaker (400-600 rpm).

#### 10.2.2.d Dopamine for urine and plasma

- 1. Pipette **75 μL** of freshly prepared **COMT Enzyme Solution** into each well of the **Microtiter Plate**. Shake briefly.
- 2. <u>For urine</u>: Pipette **100** µL of 1:51 <u>prediluted extracted</u> **Standard, Control and urine sample** into the respective wells.

During this step hold the pipette tips directly into the COMT solution. Color changes to pink. Shake briefly.

<u>For plasma</u>: Pipette **100**  $\mu$ L of 1:51 <u>prediluted extracted</u> **Standard, Control** and <u>undiluted extracted</u> **plasma sample** into the respective wells.

During this step hold the pipette tips directly into the COMT solution. Color changes to pink. Shake briefly.

- 3. Pipette **50 µL** of **Dopamine Antiserum** (violet colored) into each well.
- 4. Cover plate with adhesive foil. **Incubate 120 min** at **RT (18 °C 25 °C)** on an orbital shaker (400–600 rpm).

#### 10.2.3 ELISA

The following procedure must be performed for Adrenalin, Noradrenalin and Dopamine.

- 1. Remove adhesive foil. Discard incubation solution. Wash plate **6 x** with **250-300 μL** of diluted **Wash Buffer.** Remove excess solution by tapping the inverted plate on a paper towel.
- 2. Pipette **100 µL** of freshly prepared **Enzyme Conjugate** into each well.
- 3. Cover plate with new adhesive foil. **Incubate 60 min** at **RT (18 °C 25 °C)** on an orbital shaker (400-600 rpm).
- 4. Remove adhesive foil. Discard incubation solution. Wash plate **6 x** with **250-300 μL** of diluted **Wash Buffer.** Remove excess solution by tapping the inverted plate on a paper towel.
- 5. For adding of Substrate and Stop Solution use, if available, an 8-channel Micropipettor. Pipetting should be carried out in the same time intervals for Substrate and Stop Solution. Use positive displacement and avoid formation of air bubbles.
- 6. Pipette **200 µL** of **PNPP Substrate Solution** into each well.
- 7. Incubate 40 min at RT (18 °C 25 °C) (without adhesive foil) on an orbital shaker (400–600 rpm).
- 8. Stop the substrate reaction by adding **50 µL** of **PNPP Stop Solution** into each well. Briefly mix contents by gently shaking the plate.
- 9. **Measure** optical density with a photometer at **405 nm** (Reference-wavelength: 620-650 nm) within **60 min** after pipetting of the Stop Solution. No air bubbles should be visible.

#### 11 AUTOMATED PROCEDURE

#### 11.1 PRE-TEST SETUP INSTRUCTIONS (automated version)

The contents of the kit for 3 x 96 determinations can be divided into 2 separate runs.



Visible amounts of gel can be separated from surface of extraction plate during extraction. This has no influence on test results.

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Air contamination by peroxygen containing disinfectants for cleaning of surfaces or equipment used as powder or as solutions, e.g. VIRKON<sup>®</sup> must be avoided in any case. They will strongly disturb assay performance. VIRKON<sup>®</sup> is a trademark of DuPont.

#### 11.1.1 Dilution of Samples

Samples suspected to contain concentrations above the highest standard have to be diluted as follows:

Sample	to be diluted	with	Remarks	
Plasma	> highest standard	bidist. water	prior to extraction step	
Urine	> highest standard	0.1 N HCI	prior to extraction step	

#### 11.1.2 Extraction of Samples, Standards and Controls (Extraction Plate) (automated version)

- 1. Pipette **30 μL** of each **Standard, Control and urine sample** and **750 μL** of each **plasma sample** into the respective wells of the extraction plate. Add **750 μL** of **bidist. water** to all wells except for the **plasma samples** to correct differences of volumes.
- 2. Pipette **1000 µL** of **Extraction Buffer** into each well.
- Cover plate with adhesive foil. Extract 30 min at RT (18 °C 25 °C) on an orbital shaker (600 900 rpm).
  During extraction the surface of the liquid should wet the adhesive foil, but the liquid level should not exceed 2/3 of the well. Splashing does not affect results.
- 4. Remove adhesive foil. Immediately empty plate and eliminate residual fluid on a paper towel.
- 5. Pipette 2 mL of bidist water into each well.
- 6. Cover plate with new adhesive foil.
  - Shake 5 min at RT (18 °C 25 °C) on an orbital shaker (600–900 rpm). Splashing does not affect results.
- 7. Remove adhesive foil. Immediately empty plate and eliminate residual fluid on a paper towel. Remove fluid completely.
- 8. Pipette **150 μL** of **Extraction Buffer** into each well. To each well add **50 μL** of **Acylation Reagent**. Mix immediately after pipetting.
- 9. Extract 20 min at RT (18 °C 25 °C) (without adhesive foil) on an orbital shaker (400–600 rpm).
- 10. Immediately empty plate and eliminate residual fluid on a paper towel. Remove fluid completely.
- 11. Pipette 2 mL of bidist. water into each well.
- 12. Cover plate with new adhesive foil. Shake 5 min at RT (18 °C 25 °C) on an orbital shaker (600 900 rpm). Splashing does not affect results.
- 13. Remove adhesive foil. Immediately empty plate and eliminate residual fluid on a paper towel. Remove fluid completely.
- 14. Pipette 450 µL of Release Buffer into each well.
- 15. Shake 30 min at RT (18 °C 25 °C) (without adhesive foil) on an orbital shaker (400–600 rpm).

Prepared samples should be assayed the same day. If this is not possible, you can store the extraction plate covered with adhesive foil at 2 °C - 8 °C over night.

Important for measurement of **Dopamine** Dilution of extracted **standard**, **controls** must be performed prior to pipetting into wells of Microtiterplate in extra tubes. Predilute **urine samples** as well. Therefore, dilute all extracted Standards, Controls and urine samples 1:51 with Release Buffer in disposable tubes. (i.e. 10 µL extracted samples + 500 µL Release Buffer). Extracted **plasma samples** does not require this predilution.

# 11.1.3 Preparation of concentrated components

The volumes stated below are for one run with 3 x 6 strips (3 x 48 determinations)

Dilute / dissolve	Component	with	Diluent	Relation	Remarks	Storage	Stability
75 mL	WASHBUF CONC	675 mL	bidist. water	1:10	Mix vigorously.	2 °C - 8 °C	4 weeks
380 µL	ENZCONJ CONC	19 mL	WASHBUF (diluted)	1:51	Prepare freshly and use only once. Mix without foaming.	18 °C - 25 °C	5 hours

#### 11.2 TEST PROCEDURE (automated version)

#### 11.2.1 Preparation of COMT Enzyme Solution

#### $! \ge$ The COMT Enzyme Solution should be freshly prepared directly before use.

Dissolve each kit component of lyophilized COMT in 1.25 mL bidist. water and mix the dissolved COMT.\*

Then **pipette 1.25 mL** of **Coenzyme Solution** followed by **1.25 mL** of **Enzyme Buffer** and **0.40 mL COMT Additive** to the mixed COMT vials to give a final volume of 4.15 mL of COMT Enzyme Solution per vial. Pool three (3) vials for 48 determinations of adrenalin and 48 determinations of noradrenalin and 48 determinations of dopamine. Solution may be turbid. Mix without foaming. The COMT solution is stable at room temperature for 1 hour.

\* If only an aliquot of the COMT solution is needed, the rest of the COMT solution should be frozen immediately in aliquots at -20 °C. The COMT solution is stable under these conditions for 1-2 months.

# 11.2.1.a For research use of tissue homogenates and cell culture supernatants a general recommendation can be given:

Working with cell culture supernatants depends on the matrix as well as concentrations expected:

According to the urine protocol (extraction of at least 20 µL supernatant) a sensitivity for adrenalin of 0.3 ng/mL, for noradrenaline of 0.6 ng/mL and for dopamine of 5 ng/mL for diluted sample can be expected.

In case of a matrix with addition of serum (FCS), plasma protocol (extraction of 500 µL supernatant) can be used with the sensitivities corresponding to the plasma protocol (see 16. PERFORMANCE).

For tissue homogenates no perchloric acid should be used for homogenization. For further details ask DRG.

#### 11.2.2 Enzymatic Derivatization of Samples, Standards and Controls (Microtiter Plate)

#### 11.2.2.a Adrenalin for urine and plasma

- 1. Pipette **75 μL** of freshly prepared **COMT Enzyme Solution** into each well of the **Microtiter Plate**. Shake plate 1 min.
- 2. Pipette **100 µL** of each <u>extracted</u> **Standard, Control and sample** into the respective wells. Shake plate 1 min.
- 3. Pipette **50 µL** of **Adrenalin Antiserum** (green colored) into each well.
- 4. Cover plate. Incubate 120 min at RT (18 °C 25 °C) on an orbital shaker (400–600 rpm).

#### 11.2.2.b Noradrenalin for urine and plasma

- 1. Pipette **25 μL** of freshly prepared **COMT Enzyme Solution** into each well of the **Microtiter Plate**. Shake plate 1 min.
- 2. Pipette **25 μL** of each <u>extracted</u> **Standard, Control and sample** into the respective wells. Shake plate 1 min.
- 3. Pipette 50 µL of Noradrenalin Antiserum (blue colored) into each well.
- 4. Cover plate. Incubate 120 min at RT (18 °C 25 °C) on an orbital shaker (400–600 rpm).

#### 11.2.2.c Dopamine for urine and plasma

- 1. Pipette **75 μL** of freshly prepared **COMT Enzyme Solution** into each well of the **Microtiter Plate**. Shake briefly.
- 2. For urine: Pipette 100 µL of 1:51 prediluted extracted Standard, Control and urine sample into the respective wells.

During this step hold the pipette tips directly into the COMT solution. Color changes to pink. Shake briefly.

<u>For plasma</u>: Pipette **100 μL** of 1:51 <u>prediluted extracted</u> Standard, Control and <u>undiluted extracted</u> plasma sample into the respective wells.

During this step hold the pipette tips directly into the COMT solution. Color changes to pink. Shake plate 1 min.

- 3. Pipette **50 µL** of **Dopamine Antiserum** (violet colored) into each well.
- 4. Cover plate. Incubate 120 min at RT (18 °C 25 °C) on an orbital shaker (400–600 rpm).

#### 11.2.3 ELISA

The following procedure must be performed for Adrenalin, Noradrenalin and Dopamine.

- 1. Discard incubation solution. Wash plate **6 x** with **250-300 µL** of diluted **Wash Buffer**.
- 2. Pipette **100 µL** of **Enzyme Conjugate** into each well.
- 3. Cover plate. Incubate 60 min at RT (18 °C 25 °C) on an orbital shaker (400–600 rpm).
- 4. Discard incubation solution. Wash plate **6 x** with **250-300 µL** of diluted **Wash Buffer**.
- 5. Pipetting should be carried out in the same time intervals for Substrate and Stop Solution.
- 6. Pipette 200 µL of PNPP Substrate Solution into each well.
- 7. Incubate 40 min at RT (18 °C 25 °C) on an orbital shaker (400–600 rpm). If temperature in automat exceeds 25°C, shorten incubation time to 30 min to avoid signal overflow.
- 8. Stop the substrate reaction by adding **50 µL** of **PNPP Stop Solution** into each well. Briefly mix contents by gently shaking the plate.
- 9. **Measure** optical density with a photometer at **405 nm** (Reference-wavelength: 620-650 nm) within **60 min** after pipetting of the Stop Solution.

#### **12 QUALITY CONTROL**

The test results are only valid if the test has been performed following the instructions. Moreover the user must strictly adhere to the rules of GLP (Good Laboratory Practice) or comparable standards/laws. User and/or laboratory must have a validated system to get diagnosis according to GLP.

All kit controls must be found within the acceptable ranges as stated on the labels and the QC certificate. If the criteria are not met, the run is not valid and should be repeated. Each laboratory should use known samples as further controls. It is recommended to participate at appropriate quality assessment trials.

In case of any deviation the following technical issues should be proven: Expiration dates of (prepared) reagents, storage conditions, pipettes, devices, incubation conditions and washing methods.

# **13 CALCULATION OF RESULTS**

The obtained OD of the standards (y-axis, linear) are plotted against their concentration (x-axis, logarithmic) either on semi-logarithmic graph paper or using an automated method. A good fit is provided with cubic spline, 4 Parameter Logisitcs or Logit-Log.

For the calculation of the standard curve, apply each signal of the standards.

The concentrations for adrenaline, noradrenaline and dopamine of the kit Controls and of the urine samples in ng/mL can be read directly from the corresponding standard curve.

The results for <u>Adrenaline and Noradrenaline</u> in plasma samples in ng/mL have to be divided by 25. This correction factor responds to the difference in the volume required during the extraction step (20  $\mu$ L of standards vs 500  $\mu$ L plasma for manual version and 30  $\mu$ L standards vs 750  $\mu$ L plasma for the automated version).

The results for **Dopamine** in plasma samples have to be divided by 1275. This correction factor responds to the difference above mentioned during the extraction procedure and to the 1:51 predilution of the standards. To convert from ng/mL to pg/mL please multiply by 1000.

In case of diluted samples the values have to be multiplied with the corresponding dilution factor.

Samples showing concentrations above the highest standard have to be diluted as described in PRE-TEST SETUP INSTRUCTIONS and reassayed.

Calculate the 24 h excretion for each urine sample:

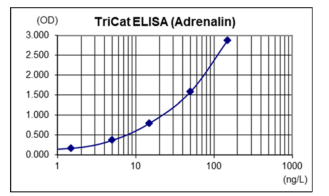
 $\mu g/24 h = \mu g/L x L/24 h$ 

Conversion:

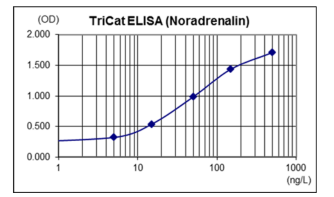
1000 pg/mL = 1 ng/mL Adrenalin ( $\mu$ g/L) x 5.458 = nmol/L Noradrenalin ( $\mu$ g/L) x 5.911 = nmol/L Dopamine ( $\mu$ g/L) x 6.528 = nmol/L

#### **Typical Calibration Curve**

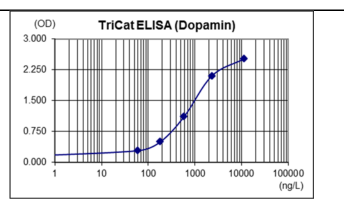
(Example. Do not use for calculation!)							
Standard	Adrenalin (ng/mL)	$OD_Mean$	OD/OD <sub>max</sub> (%)				
А	0.0	0.088	3.1				
В	1.5	0.167	5.8				
С	5.0	0.368	12.8				
D	15	0.796	27.6				
E	50	1.579	54.8				
F	150	2.881	100				



Standard	Noradrenali n (ng/mL)	$OD_Mean$	OD/OD <sub>max</sub> (%)
A	0	0.223	0.0
В	5	0.322	6.7
С	15	0.539	21.3
D	50	0.984	51.2
E	150	1.438	81.8
F	500	1.708	100



Standard	Dopamine	$OD_Mean$	OD/OD <sub>max</sub>
	(ng/mL)		(%)
А	0	0.135	0
В	60	0.287	6.4
С	180	0.500	15.3
D	585	1.113	41.0
E	2300	2.105	82.5
F	11470	2.522	100



# 14 EXPECTED VALUES

The results themselves should not be the only reason for any therapeutical consequences. They have to be correlated to other clinical observations and diagnostic tests.

Apparently healthy subjects show the following values: (5 % - 95 % percentile)

It is recommended that each laboratory establishes its own range of normal values.

	Urine		Plasma	
	µg/d	nmol/d	pg/mL	nmol/L
Adrenalin	< 20	< 110	< 125	< 0.68
Noradrenalin	< 90	< 535	< 600	< 3.55
Dopamine	< 600	< 3917	< 100	< 0.65

# **15 LIMITATIONS OF THE PROCEDURE**

Specimen collection and storage have a significant effect on the test results. See SPECIMEN COLLECTION AND STORAGE for details.

For cross-reactivities, see PERFORMANCE.

The following blood components do not have a significant effect (+/-20% of expected) on the test results up to the below stated concentrations:

Hemoglobin	2.0 mg/mL		
Bilirubin	1.0 mg/mL		
Triglyceride	91 mg/mL		

#### 16 PERFORMANCE

Analytical Specificity (Cross Reactivity)	Substance	Adrenalin	Noradrenalin	Dopamine			
	Adrenalin	100	< 0.02	< 0.05			
	Noradrenalin	< 0.4	100	< 0.05	•		
	Dopamine	< 0.1	< 0.02	100	Cross-reactivity of other substances tested < 0.5 %		
	Metanephrine	< 0.1	< 0.002	< 0.05			
	Normetanephrine	< 0.1	< 0.005	< 0.05	100		
	3-MT	< 0.1	< 0.002	< 0.05			
	L-DOPA	< 0.1	< 0.001	< 0.05			
	A dranalin	Urine	0.2 ng/mL				
	Adrenalin	Plasma	8 pg/mL	1			
Analytical Sensitivity	N	Urine	0.6 ng/mL	Manager al (Zama Chandand) + 200			
(Limit of Detection)	Noradrenalin	Plasma	20 pg/mL	Mean signal (Z	Mean signal (Zero-Standard) + 2SE		
		Urine	5 ng/mL				
	Dopamine	Plasma	4 pg/mL				
Precision			Range (ng/mL)	CV (%)			
	A day I'	Urine	5.0 - 64.3	8.7			
	Adrenalin	Plasma	0.046 - 1.060	6.8			
Intra-Assay	<b>N</b> I I II	Urine	16.0 – 256	7.3			
	Noradrenalin	Plasma	0.560 - 12.38	7.4			
	<b>.</b>	Urine	37 – 1549	7.0			
	Dopamine	Plasma	0.046 - 1.402	10.9			
		Urine	5.2 - 74.5	12.1			
	Adrenalin	Plasma	0.051 - 0.979	15.2			
		Urine	15.4 - 391.5	12.1			
Inter-Assay	Noradrenalin	Plasma	0.569 - 1.945	12.5			
		Urine	47 – 990	10.8			
	Dopamine	Plasma	0.213 - 1.055	16.3			
			Range (ng/mL)	Serial dilution u	p to	Range (%)	
	Adrenalin	Urine	2.7 – 114.6	1:32		85 - 105	
		Plasma	0.002 - 0.837	1:32		76 - 120	
Linearity		Urine	5.1 - 423	1:32		85 - 115	
Linearity	Noradrenalin	Plasma	0.02 - 8.2	1:32		89 - 111	
	D	Urine	141 - 5732	1:32		87 - 114	
	Dopamine	Plasma	0.04 – 1.3	1:16		100 - 135	
		No Hig	gh dose hook effe	ect detected.			
			Mean (%)	Range (%)			
Recovery	A due := = liss	Urine	95	85 - 106			
	Adrenalin	Plasma	100	85 - 120		a. <b>-</b>	
	Noradrenalin	Urine	100.9	81 - 116		% Recovery	
		Plasma	97.5	83 - 111		after spiking	
		Urine	101.5	89 - 113			
	Dopamine	Plasma	91	70 - 113			
	Adrenalin		DRG= 0.91 x HPL	C + 14.0; r = 0.96	9; n = <sup>-</sup>	120	
Method Comparison versus HPLC	Noradrenalin	DRG = 0.75 x HPLC + 4.8; r = 0.945; n = 134					
VEISUS IIF LU	Dopamine	DRG = 1.06 x HPLC - 0.154; r = 0.985; n = 90					
	•						

# 17 SHORT PROTOCOL TRICAT (ADRENALINE, NORADRENALINE AND DOPAMINE) ELISA

Total Assay Time	appr. 7 hours
Specimen	EDTA Plasma, Urine
Extraction and Acylation	
Standards, Controls and Urine Samples	20 μL (Automated version 30 μL)
Plasma Samples	500 μL (Automated version 750 μL)
Addition of bidest. water to Standards, Controls and Urine samples	500 μL (Automated version 750 μL)
Extraction buffer	1 mL
Incubation	30 min; RT; on orbital shaker (600-900 rpm)
Washing with bidest. water	2 mL
Incubation	5 min; RT; on orbital shaker (600-900 rpm)
Extraction buffer	150 μL
Acylation Reagent	50 μL
Incubation	20 min; RT; on orbital shaker (400-600 rpm)
Washing with bidest. water	2 mL
Incubation	5 min; RT; on orbital shaker (600-900 rpm)
Release buffer	300 μL (Automated version: 450 μL)
Incubation	30 min; RT; on orbital shaker (400-600 rpm)

Extraction plate can be stored overnight covered at 2 °C - 8 °C

<b>For Dopamine only</b> : Predilution of extracted standards, controls and urine samples with Release Buffer in separated tubes	10 μL + 500 μL			
Samples are now ready for the ELISA				
Microtiter plate procedure		1		
	Adrenalin	Noradrenalin	Dopamine	
COMT enzymatic solution	75 μL	25 μL	75 µL	
Prediluted extracted Dopamine Standards, Controls and Urine samples	_	-	100 µL	
Extracted Dopamine Plasma Samples (without predilution)	-	-	100 µL	
Extracted Standards, Controls and Samples (Urine and Plasma)	100 µL	25 µL	-	
Antiserum	50 μL (green)	50 µL (blue)	50 µL (violet)	
Incubation	120 min; RT; on orbital shaker (400-600 rpm)			
Washing	6 x 300 μL			
Enzyme conjugate	100 µL			
Incubation	60 min; RT; on orbital shaker (400-600 rpm)			
Washing	6 x 300 μL			
Substrate	200 µL			
Incubation	40 min (30 min Automated Version) ; RT; on orbital shaker (400-600 rpm)			
Stop solution		50 µL		
Measurement Optical density	within 60 min. at 405 nm / 620-650 nm			

#### **18 LITERATURE REFERENCES**

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- 2. Creces J., Appleton Ch.: Catecholamines and their Metabolites: Evaluation of a commercial ELISA. Clin. Biochem., QML Pathology, Brisbane QLD (2004)
- 3. Adams, J. M. et al. Effects of 17β-Estradiol on hypoglycemia-induced increases in plasma catecholamines in the rat. Poster Society for Neuroscience, Annual Meeting, New Orleans (2003)
- 4. Westermann J, Hubl W, Kaiser N, Salewski L, Simple, rapid and sensitive determination of epinephrine and norepinephrine in urine and plasma by non-competitive enzyme immunoassay, compared with HPLC method. Clin. Lab., 48: 61-71 (2002)

# SYMBOLS USED

Symbol	English	Deutsch	Italiano	Español	Français
(6	European Conformity	CE-Konformitäts- kennzeichnung	Conformità europea	Conformidad europea	Conformité normes européennes
Í	Consult instructions for use *	Gebrauchsanweisung beachten *	Consultare le istruzioni per l'uso	Consulte las instrucciones de uso	Consulter les instructions d'utilisation
IVD	<i>In vitro</i> diagnostic medical device *	<i>In-vitro</i> -Diagnostikum *	Dispositivo medico- diagnostico in vitro	Producto sanitario para diagnóstico In vitro	Dispositif médical de diagnostic in vitro
REF	Catalogue number *	Artikelnummer *	Numero di Catalogo	Nûmero de catálogo	Référence de catalogue
LOT	Batch code *	Fertigungslosnummer, Charge *	Codice del lotto	Codigo de lote	Numéro de lot
Σ	Contains sufficient for <n> tests *</n>	Ausreichend für <n> Prüfungen *</n>	Contenuto sufficiente per "n" saggi	Contenido suficiente para <n> ensayos</n>	Contenu suffisant pour "n" tests
	Temperature limit *	Temperaturbegrenzung *	Temperatura di conservazione	Temperatura de conservacion	Température de conservation
	Use-by date *	Verwendbar bis *	Utilizzare prima del	Establa hasta	Utiliser jusque
AAA	Manufacturer *	Hersteller *	Fabbricante	Fabricante	Fabricant
8	Biological risks*	Biologische Risiken*	Rischi biologici	Riesgos biológicos	Risques biologiques
$\triangle$	Caution *	Achtung *	Attenzione	Precaución	Attention
RUO	For research use only	Nur für Forschungszwecke	Solo a scopo di ricerca	Sólo para uso en investigación	Seulement dans le cadre de recherches
Distributed by	Distributed by	Vertreiber	Distributore	Distribuidor	Distributeur
Content	Content	Inhalt	Contenuto	Contenido	Contenu
Volume/No.	Volume / No.	Volumen/Anzahl	Volume/Quantità	Volumen/Número	Volume/Quantité