

Rapid Amphetamine Urine Test

Ref No. 0118

INTENDED USE

Acro Rapid Amphetamine Urine Test is a lateral flow, rapid immunoassay for the qualitative detection of Amphetamine in human urine at a cutoff of 1000 ng/mL. The test is used to obtain a visual qualitative result and is intended for laboratory use only.

This assay provides only a preliminary result. Clinical consideration and professional judgment must be applied to a drug test result, particularly in evaluating a preliminary positive result. In order to obtain a confirmed analytical result, a more specific alternate chemical method is needed. Gas Chromatography/ Mass Spectroscopy (GC/MS) analysis is preferred.

SUMMARY AND EXPLANATION

Amphetamine is a potent central nervous system stimulant. Acute higher doses induce euphoria, alertness, and sense of increased energy and power. More acute responses produce anxiety, paranoia, psychotic behavior, and cardiac dysrhythmias.

The length of time following drug use for which a positive result may occur is dependent upon several factors, including the frequency and amount of drug, metabolic rate, excretion rate, drug half-life, the drug user's age, weight, activity and diet.

TEST PRINCIPLE

The Acro Rapid Amphetamine Urine Test is based on the principle of competitive immunochemical reaction between an immobilized drug-protein conjugate and the drug or drug metabolites, which may be present in the urine sample for limited binding sites for the drug/drug metabolites of a labeled drug antibody. The test device contains a nitrocellulose membrane strip pre-coated with drug-protein conjugate in the test region and a wicking pad containing colored antibody-colloidal gold conjugate. During the test, the urine sample is allowed to migrate upward and hydrate the antibody-colloidal gold conjugate. The mixture then migrates along the membrane chromatographically by capillary action to the immobilized drug-protein band in the test region. When drug is absent in the urine, the colored antibody-colloidal gold conjugate and immobilized drug-protein bind specifically to form a visible line in the test region (test line). When amphetamine is present in the urine sample, it will compete with the drug-protein conjugate for the limited antibody binding sites. When the drug is present in sufficient concentration in the urine sample, it will fill the limited antibody binding sites, which will inhibit attachment of the colored antibody-colloidal gold conjugate to the drug-protein conjugate in the test region. Therefore the presence of the test line indicates a **negative** result for amphetamine and the absence of the test line indicates a preliminary **positive** result for amphetamine.

Another visible line, control line, generated by a different antigen/antibody reaction is also present at a control region (C) next to the test region of the test strip. The control line should always appear, regardless of the presence of the drug or drug metabolites in the urine sample. This means that a **negative** urine sample will produce **two** lines (test line and control line), and a **positive** urine sample will generate only **one** line (control line). The presence of control line serves as a built-in control, which demonstrates that the test is performed properly.

REAGENTS & MATERIALS SUPPLIED

1. Test Cassette contains membrane-embedded reagents in a protein matrix containing sodium azide as a preservative.
2. Anti-amphetamine monoclonal antibody is from murine ascites.
3. Dropper. A transfer pipette is included with each test device inside the foil pouch.
4. Test Instructions

MATERIALS REQUIRED, BUT NOT PROVIDED

1. Timer
2. Sample container

WARNINGS AND PRECAUTIONS

1. For laboratory *in vitro* diagnostic use only
2. Urine specimens may be potentially infectious. Proper handling and disposal procedures should be followed.
3. Avoid cross-contamination of urine samples. Use a new dropper or transfer pipette for adding each test sample.
4. Test device should remain sealed until ready for use.
5. Do not use the test kit after the expiration date.

STORAGE AND STABILITY

Store at 2-30°C (36-86°F) in the original sealed pouch. Do not freeze.

SPECIMEN COLLECTION AND HANDLING

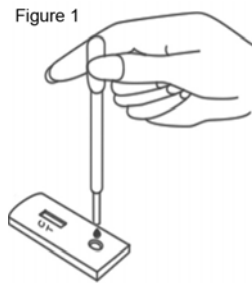
Fresh urine does not require any special handling or pretreatment. A clean dry plastic or glass container may be used for specimen collection. If the specimen is not immediately tested after the specimen collection, the specimen may be refrigerated at 2-8°C for up to 3 days or frozen at -20°C for longer period of time. Specimens that have been refrigerated

must be equilibrated to room temperature prior to testing. Frozen specimens must be thawed and mixed thoroughly prior to testing.

Note: Urine specimens and all materials coming in contact with the specimens should be handled and disposed as if capable of transmitting infection. Avoid contact with skin by wearing gloves and proper laboratory attire.

ASSAY PROCEDURE

1. Bring the test components and urine sample to room temperature (15° - 28° C) before testing. Do not open the foil pouch until ready to begin testing.
2. Open the foil pouch at the notch and remove the test device and dropper prior to testing. Place the device on a clean, level surface.
3. Hold the dropper vertically and dispense 2 full drops (80 µl) of urine sample without air bubbles into the sample well "S" of the test device (figure 1).
4. Read the result between 5-10 minutes (figure

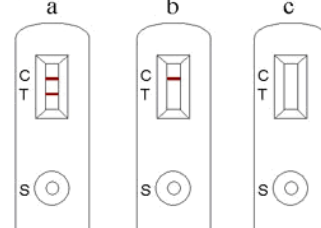


IMPORTANT: The result must be read at five - eight minutes. Waiting more than eight minutes may cause the test result to be inaccurate.

INTERPRETATION OF RESULTS

1. **Preliminary Positive:** a *rose-pink* color band appears in the Control Zone "C" but not in the Test Zone "T". A preliminary positive result indicates amphetamine level in the urine sample is at or above the detection sensitivity of 1000 ng/mL. The sample should be confirmed.
2. **Negative:** two horizontal *rose-pink* color bands appear, one in the Control zone "C" and one in the Test Zone "T". A negative result indicates amphetamine level in the urine sample is below the detection sensitivity of 1000 ng/mL.
3. **Invalid:** no *rose-pink* bands appear, or a band appears in the Test Zone "T", but not in the Control Zone "C". An invalid result may be due to improper testing procedures or deterioration of the kit components.

Figure 2



Note: There is no meaning attributed to line color intensity or width.

QUALITY CONTROL

An internal procedure control is included in the test device. A control line must form regardless of the presence or absence of drugs or metabolites. The presence of the line in the Control region indicates that a proper sample volume has been used. If the line in the Control region does not form, the test is considered invalid.

To ensure proper kit performance, it is recommended that the test devices be tested once a week or prior to use with external controls. External controls are available from commercial sources. It is important to make sure that the control values are within established limits. If the values of external controls do not fall within established limits, the test results are invalid. Additional controls may be tested according to guidelines or requirement of local state, and/or federal regulations or accrediting organizations.

LIMITATIONS OF PROCEDURE

1. The assay is designed for use with human urine only.
2. A preliminary positive result indicates only the presence of amphetamine and does not indicate or measure intoxication
3. There is a possibility that technical or procedural error as well as other substances as factors not listed may interfere with the test and cause false results. See SPECIFICITY section for substances that will produce positive results, or that do not interfere with the test performance.
4. If adulteration is suspected, the test should be repeated with a new sample.
5. Certain over the counter or prescription medications (or certain foods) may cause false results.

PERFORMANCE CHARACTERISTICS

1. **Sensitivity.** The Acro Rapid Amphetamine Urine Test detects amphetamine and its metabolites in urine at concentrations equal to or greater than 1000 ng/mL.
2. **Specificity.** Interference of substances that may be present in urine specimens, as well as sample effect of sample pH and specificity were studied.
 - a. Cross-reactivity of non-amphetamine related compounds at concentrations much higher than normally found in the urine of people using or abusing them were tested using the assay devices. No cross-reactivity was detected with the substances listed in Table I.
 - b. Table II lists Amphetamine related substances and concentrations that produced results approximately equivalent to the cutoff level for amphetamine.

- c. Varying sample pH within the range of 4 and 9 has no significant effect on the assay results.
- d. Varying sample specific gravity within the range of 1.003 and 1.040 has no significant effect on the assay results.

Table-I: Compounds tested and found not to cross-react with the test at the concentrations of 10 µg/mL and 100 µg/mL in urine.

<i>Acetaminophen</i>	<i>Ibuprofen</i>
<i>Acetone</i>	<i>(+/-)-Isoproterenol</i>
<i>Albumin</i>	<i>Ketamine</i>
<i>Ampicillin</i>	<i>Levorphanol</i>
<i>Ascorbic Acid</i>	<i>Lidocaine</i>
<i>Aspartame</i>	<i>(+)-Naproxen</i>
<i>Aspirin</i>	<i>Niacinamide</i>
<i>Atropine</i>	<i>Nicotine</i>
<i>Benzocaine</i>	<i>(+/-)-Norephedrine</i>
<i>Bilirubin</i>	<i>Oxalic Acid</i>
<i>Caffeine</i>	<i>Penicillin-G</i>
<i>Chloroquine</i>	<i>Pheniramine</i>
<i>(+)-Chlorpheniramine</i>	<i>Phenothiazine</i>
<i>(+/-)-Chlorpheniramine</i>	<i>1-Phenylephrine</i>
<i>Creatine</i>	<i>β-Phenylethylamine</i>
<i>Dexbrompheniramine</i>	<i>Procaine</i>
<i>Dextromethorphan</i>	<i>Quinidine</i>
<i>Diphenhydramine</i>	<i>Ranitidine</i>
<i>Dopamine</i>	<i>Riboflavin</i>
<i>(+/-)-Epinephrine</i>	<i>Sodium Chloride</i>
<i>Erythromycin</i>	<i>Sulindac</i>
<i>Ethanol</i>	<i>Theophylline</i>
<i>Furosemide</i>	<i>Tyramine</i>
<i>Glucose</i>	<i>4-</i>
<i>Guaiacol Glyceryl Ether</i>	<i>Dimethylaminoantipyrine</i>
<i>Hemoglobin</i>	<i>(1R, 2S)-(-)-N-Methyl-Ephedrine</i>

Table-II: Concentration of amphetamine-related compounds showing a positive response approximately equivalent to the amphetamine cut-off set for the test.

Compound /	Concentration in ng/ml
<i>d-Amphetamine</i>	1,000
<i>dl-Amphetamine</i>	2,500
<i>(+/-) 3,4-MDA</i>	1,250
<i>d-Methamphetamine</i>	50,000
<i>(+/-)3,4-MDMA</i>	50,000

3. Accuracy

Comparison with GC/MS and predicate device: The accuracy study was performed by testing urine samples with GC/MS analysis results and predicate device results of the same samples.

Acro AMP	GC/MS, Cutoff 1000ng/ml			
	Near Cutoff Negative (-50% to cutoff)	Near Cutoff Positive (cutoff to +50%)	Positive (>+50%)	% Agreement with GC/MS
Positive	3	13	45	58/59 = 98%
Negative	10	0	1	10/13 = 77%

% Total Agreement 68/72 =94%

Acro AMP	Predicate Device, Cutoff 500ng/ml	
	Negative	Positive
Positive	0	59
Negative	62	1
% Agreement with Predicate Device	100%	98%

4. Cut-off Study. The cut-off of the test was determined by the repetitive assaying of six levels of amphetamine controls. The resultant data are summarized as follows:

Amphetamine conc.	# Tested	# Positive (+)	# Negative (-)	% Correct Results
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0 ng/mL	68	0	68	100%
500 ng/mL	68	0	68	100%
750 ng/mL	68	17	51	75%
1250 ng/mL	68	51	17	75%
1500 ng/mL	68	68	0	100%
2000 ng/mL	68	68	0	100%

5. Reproducibility. The reproducibility was evaluated at four different sites. The Acro Rapid Amphetamine Urine Test was tested against blind-labeled urine controls containing 0, 500, 750, 1250, 1500 and 2000 ng/mL amphetamine at each site. The results are summarized as follows:

Test Sites	0 ng/mL		500ng/mL		750ng/mL		1250ng/mL		1500ng/mL		2000ng/mL	
	#	Result	#	Result	#	Result	#	Result	#	Result	#	Result
1	20	20-	20	20-	20	4+, 16-	20	14+, 6-	20	20+	20	20+
2	16	16-	16	16-	16	4+, 12-	16	13+, 3-	16	16+	16	16+
3	16	16-	16	16-	16	4+, 12-	16	12+, 4-	16	16+	16	16+
4	16	16-	16	16-	16	5+, 11-	16	12+, 4-	16	16+	16	16+
Total	68	68-	68	68-	68	17+, 51-	68	51+, 17-	68	68+	68	68+

6. Stability Study. As determined by temperature accelerated stability study method, the shelf life of the product under the specified storage condition is 24 months from the date of production.

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