



April 4, 2025

Hangzhou Alltest Biotech Co.,Ltd
% Jenny Xia
Director
LSI International Inc
504E Diamond Ave., Suite H
Gaithersburg, Maryland 20877

Re: K250727

Trade/Device Name: AllTest Multi-Drug Rapid Urine Test Cup; AllTest Multi-Drug Urine Test Cup
Regulation Number: 21 CFR 862.3100
Regulation Name: Amphetamine Test System
Regulatory Class: Class II
Product Code: NFT, PTH, NGL, NFV, NFY, PTG, NGG, QBF, NGM, QAW, NFW
Dated: March 9, 2025
Received: March 11, 2025

Dear Jenny Xia:

We have reviewed your section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (the Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database available at <https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm> identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Additional information about changes that may require a new premarket notification are provided in the FDA guidance documents entitled "Deciding When to Submit a 510(k) for a Change to an Existing Device" (<https://www.fda.gov/media/99812/download>) and "Deciding When to Submit a 510(k) for a Software Change to an Existing Device" (<https://www.fda.gov/media/99785/download>).

Your device is also subject to, among other requirements, the Quality System (QS) regulation (21 CFR Part 820), which includes, but is not limited to, 21 CFR 820.30, Design controls; 21 CFR 820.90, Nonconforming product; and 21 CFR 820.100, Corrective and preventive action. Please note that regardless of whether a change requires premarket review, the QS regulation requires device manufacturers to review and approve changes to device design and production (21 CFR 820.30 and 21 CFR 820.70) and document changes and approvals in the device master record (21 CFR 820.181).

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801 and Part 809); medical device reporting (reporting of medical device-related adverse events) (21 CFR Part 803) for devices or postmarketing safety reporting (21 CFR Part 4, Subpart B) for combination products (see <https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products>); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR Part 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR Parts 1000-1050.

All medical devices, including Class I and unclassified devices and combination product device constituent parts are required to be in compliance with the final Unique Device Identification System rule ("UDI Rule"). The UDI Rule requires, among other things, that a device bear a unique device identifier (UDI) on its label and package (21 CFR 801.20(a)) unless an exception or alternative applies (21 CFR 801.20(b)) and that the dates on the device label be formatted in accordance with 21 CFR 801.18. The UDI Rule (21 CFR 830.300(a) and 830.320(b)) also requires that certain information be submitted to the Global Unique Device Identification Database (GUDID) (21 CFR Part 830 Subpart E). For additional information on these requirements, please see the UDI System webpage at <https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/unique-device-identification-system-udi-system>.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance>) and CDRH Learn (<https://www.fda.gov/training-and-continuing-education/cdrh-learn>). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory->

[assistance/contact-us-division-industry-and-consumer-education-dice](#)) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

**JOSEPH A.
KOTAREK-S**

Digitally signed by JOSEPH A.
KOTAREK-S
Date: 2025.04.04 07:32:33
-04'00'

Joseph Kotarek, Ph.D.
Branch Chief
Division of Chemistry and Toxicology Devices
OHT7: Office of In Vitro Diagnostics
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number (if known)
K250727

Device Name

AllTest Multi-Drug Urine Test Cup;
AllTest Multi-Drug Rapid Urine Test Cup

Indications for Use (Describe)

AllTest Multi-Drug Urine Test Cup tests are competitive binding, lateral flow immunochromatographic assays for qualitative and simultaneous detection of Amphetamine, Buprenorphine, Secobarbital, Benzodiazepine, Cocaine, Methamphetamine, Methylenedioxymethamphetamine, Morphine, Methadone, Oxycodone, Phencyclidine, Nortriptyline, Marijuana, Fentanyl, 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine, Tramadol, Propoxyphene and 6-monoacetylmorphine in human urine at the cutoff concentrations of:

Drug (Identifier)	Cut-off level
Amphetamine (AMP)	500 ng/mL or 1000 ng/mL
Buprenorphine (BUP)	10 ng/mL
Secobarbital (BAR)	300 ng/mL
Benzodiazepine (BZO)	300 ng/mL
Cocaine (COC)	150 ng/mL or 300 ng/mL
Methamphetamine (MET)	500 ng/mL or 1000 ng/mL
Methylenedioxymethamphetamine (MDMA)	500 ng/mL
Morphine (MOP/OPI)	300 ng/mL or 2000 ng/mL
Methadone (MTD)	300 ng/mL
Oxycodone (OXY)	100 ng/mL
Phencyclidine (PCP)	25 ng/mL
Nortriptyline (TCA)	1000 ng/mL
Marijuana (THC)	50 ng/mL
Fentanyl(FYL)	1 ng/mL
2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP)	300 ng/mL
Tramadol (TRA)	100 ng/mL
Propoxyphene (PPX)	300 ng/mL
6-monoacetylmorphine (6-MAM)	10 ng/mL

AllTest Multi-Drug Urine Test Cup can be a single drug test cup or used for any combinations of the above listed analytes. It is for in vitro diagnostic use only. It is intended for OTC use.

The tests may yield positive results for the prescription drugs when taken at or above prescribed doses. It is not intended to distinguish between prescription use or abuse of these drugs. Clinical consideration and professional judgment should be applied to any drug of abuse test result, particularly in evaluating a preliminary positive result.

The tests provide only preliminary results. To obtain a confirmed analytical result, a more specific alternate chemical method must be used. GC/MS or LC/MS is the recommended confirmatory method.

AllTest Multi-Drug Rapid Urine Test Cup tests are competitive binding, lateral flow immunochromatographic assays for qualitative and simultaneous detection of Amphetamine, Buprenorphine, Secobarbital, Benzodiazepine, Cocaine, Methamphetamine, Methylenedioxymethamphetamine, Morphine, Methadone, Oxycodone, Phencyclidine, Nortriptyline, Marijuana, Fentanyl, 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine, Tramadol, Propoxyphene and 6-monoacetylmorphine in human urine at the cutoff concentrations of:

Drug (Identifier)	Calibrator	Cut-off (ng/mL)
Amphetamine (AMP)	d-Amphetamine	500 or 1000

Buprenorphine (BUP)	Buprenorphine	10
Secobarbital (BAR)	Secobarbital	300
Benzodiazepine (BZO)	Oxazepam	300
Cocaine (COC)	Benzoyllecgonine	150 or 300
Methamphetamine (MET)	d-Methamphetamine	500 or 1000
Methylenedioxymethamphetamine (MDMA)	d,l-Methylenedioxymethamphetamine	500
Morphine (MOP/OPI)	Morphine	300 or 2000
Methadone (MTD)	Methadone	300
Oxycodone (OXY)	Oxycodone	100
Phencyclidine (PCP)	Phencyclidine	25
Nortriptyline (TCA)	Nortriptyline	1000
Marijuana (THC)	11-nor- Δ^9 -THC-9 COOH	50
Fentanyl (FYL)	Fentanyl	1
2-ethylidene-1,5-dimethyl- 3,3-diphenylpyrrolidine (EDDP)	2-ethylidene-1,5-dimethyl- 3,3-diphenylpyrrolidine	300
Tramadol (TRA)	Tramadol	100
Propoxyphene (PPX)	Propoxyphene	300
6-monoacetylmorphine (6-MAM)	6-monoacetylmorphine	10

AllTest Multi-Drug Rapid Urine Test Cup can be a single drug test cup or used for any combination of the above listed analytes. It is for in vitro diagnostic use only.

The tests may yield positive results for the prescription drugs when taken at or above prescribed doses. It is not intended to distinguish between prescription use or abuse of these drugs. Clinical consideration and professional judgment should be applied to any drug of abuse test result, particularly in evaluating a preliminary positive result.

The tests provide only preliminary results. To obtain a confirmed analytical result, a more specific alternate chemical method must be used. GC/MS or LC/MS is the recommended confirmatory method.

Type of Use (Select one or both, as applicable)

☐ Prescription Use (Part 21 CFR 801 Subpart D)

☒ Over-The-Counter Use (21 CFR 801 Subpart C)

CONTINUE ON A SEPARATE PAGE IF NEEDED.

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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510(k) SUMMARY
K250727

- 1. Date:** March 26, 2025
- 2. Submitter:** Hangzhou AllTest Biotech Co., Ltd.
Plant Bldg. 3, 4, 5, No. 550 Yinhai Street, Baiyang Street, Hangzhou ETDZ, Jianggan District
- 3. Contact person:** Jenny Xia
LSI International Inc.
504 East Diamond Ave., Suite H
Gaithersburg, MD 20877
Telephone: 301-525-6856
Email: jxia@lsi-consulting.org
- 4. Device Name:** AllTest Multi-Drug Urine Test Cup
AllTest Multi-Drug Rapid Urine Test Cup
- 5. Classification:** Class II

Product Code Target Drug	Regulation Section	Panel
NFT Amphetamine (AMP)	862.3100, Amphetamine Test System	Toxicology
PTH Secobarbital (BAR)	862.3150, Barbiturate Test System	Toxicology
NGL Buprenorphine (BUP) Fentanyl (FYL) Morphine (MOP/OPI) Oxycodone (OXY) 6-Monoacetylmorphine(6-MAM) Tramadol (TML)	862.3650, Opiate Test System	Toxicology
NFV Oxazepam (BZO)	862.3170, Benzodiazepine Test System	Toxicology
NFY Cocaine (COC)	862.3250, Cocaine and cocaine metabolite test system	Toxicology
PTG 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP) Methadone (MTD)	862.3620, Methadone Test System	Toxicology
NGG Methylenedioxymethamphetamine (MDMA) Methamphetamine (MET)	862.3610, Methamphetamine Test System	Toxicology
QBF Propoxyphene(PPX)	862.3700 Propoxyphene test system.	Toxicology
NGM	Unclassified	Toxicology

Phencyclidine (PCP)		
QAW Nortriptyline (TCA)	862.3910 Tricyclic antidepressant drugs test system	Toxicology
NFW Cannabinoids (THC)	862.3870, Cannabinoids Test System	Toxicology

6. Predicate Devices:

AllTest Multi-Drug Rapid Test Cup (K244043)

7. Intended Use

AllTest Multi-Drug Urine Test Cup tests are competitive binding, lateral flow immunochromatographic assays for qualitative and simultaneous detection of Amphetamine, Buprenorphine, Secobarbital, Benzodiazepine, Cocaine, Methamphetamine, Methylenedioxymethamphetamine, Morphine, Methadone, Oxycodone, Phencyclidine, Nortriptyline, Marijuana, Fentanyl, 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine, Tramadol, Propoxyphene and 6-monoacetylmorphine in human urine at the cutoff concentrations of:

Drug (Identifier)	Cut-off
Amphetamine (AMP)	500 ng/mL or 1000 ng/mL
Buprenorphine (BUP)	10 ng/mL
Secobarbital (BAR)	300 ng/mL
Benzodiazepine (BZO)	300 ng/mL
Cocaine (COC)	150 ng/mL or 300 ng/mL
Methamphetamine (MET)	500 ng/mL or 1000 ng/mL
Methylenedioxymethamphetamine (MDMA)	500 ng/mL
Morphine (MOP/OPI)	300 ng/mL or 2000 ng/mL
Methadone (MTD)	300 ng/mL
Oxycodone (OXY)	100 ng/mL
Phencyclidine (PCP)	25 ng/mL
Nortriptyline (TCA)	1000 ng/mL
Marijuana (THC)	50 ng/mL
Fentanyl (FYL)	1 ng/mL
2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP)	300 ng/mL
Tramadol (TRA)	100 ng/mL
Propoxyphene (PPX)	300 ng/mL
6-monoacetylmorphine (6-MAM)	10 ng/mL

AllTest Multi-Drug Urine Test Cup can be a single drug test cup or used for any combinations of the above listed analytes. It is for in vitro diagnostic use only. It is intended for OTC use.

The tests may yield positive results for the prescription drugs when taken at or above prescribed doses. It is not intended to distinguish between prescription use or abuse of these drugs. Clinical consideration and professional judgment should be applied to any drug of abuse test result, particularly in evaluating a preliminary positive result.

The tests provide only preliminary results. To obtain a confirmed analytical result, a more specific alternate chemical method must be used. GC/MS or LC/MS is the recommended confirmatory method.

AllTest Multi-Drug Rapid Urine Test Cup tests are competitive binding, lateral flow immunochromatographic assays for qualitative and simultaneous detection of Amphetamine, Buprenorphine, Secobarbital, Benzodiazepine, Cocaine, Methamphetamine, Methylenedioxymethamphetamine, Morphine, Methadone, Oxycodone, Phencyclidine, Nortriptyline, Marijuana, Fentanyl, 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine, Tramadol, Propoxyphene and 6-monoacetylmorphine in human urine at the cutoff concentrations of:

Drug (Identifier)	Calibrator	Cut-off (ng/mL)
Amphetamine (AMP)	d-Amphetamine	500 or 1000
Buprenorphine (BUP)	Buprenorphine	10
Secobarbital (BAR)	Secobarbital	300
Benzodiazepine (BZO)	Oxazepam	300
Cocaine (COC)	Benzoylecgonine	150 or 300
Methamphetamine (MET)	d-Methamphetamine	500 or 1000
Methylenedioxymethamphetamine (MDMA)	d,l-Methylenedioxymethamphetamine	500
Morphine (MOP/OPI)	Morphine	300 or 2000
Methadone (MTD)	Methadone	300
Oxycodone (OXY)	Oxycodone	100
Phencyclidine (PCP)	Phencyclidine	25
Nortriptyline (TCA)	Nortriptyline	1000
Marijuana (THC)	11-nor- Δ^9 -THC-9 COOH	50
Fentanyl (FYL)	Fentanyl	1
2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP)	2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine	300
Tramadol (TRA)	Tramadol	100
Propoxyphene (PPX)	Propoxyphene	300
6-monoacetylmorphine (6-MAM)	6-monoacetylmorphine	10

AllTest Multi-Drug Rapid Urine Test Cup can be a single drug test cup or used for any combination of the above listed analytes. It is for in vitro diagnostic use only.

The tests may yield positive results for the prescription drugs when taken at or above prescribed doses. It is not intended to distinguish between prescription use or abuse of these drugs. Clinical consideration and professional judgment should be applied to any drug of abuse test result, particularly in evaluating a preliminary positive result.

The tests provide only preliminary results. To obtain a confirmed analytical result, a more specific alternate chemical method must be used. GC/MS or LC/MS is the recommended confirmatory method.

8. Device Description

AllTest Multi-Drug Urine Test Cup and AllTest Multi-Drug Rapid Urine Test Cup are immunochromatographic assays that use a lateral flow system for the qualitative detection of single or multiple drugs in human urine.

The devices are a cup format. Each test device is sealed with sachets of desiccant in an aluminum pouch. The device is in a ready-to-use format.

9. Substantial Equivalence Information

Similarities		
Item	Device	Predicate (K244043)
Intended use	Qualitative detection of drugs of abuse in urine. For prescription use or over-the-counter use	Same
Methodology	Competitive binding, lateral flow immunochromatographic assay based on antigen-antibody reaction	Same
Type of Test	Qualitative	Same
Specimen Type	Human urine	Same
Target Drug and Cut Off Values	Target Drugs	Cutoff (ng/mL)
	Amphetamine(AMP)	1000 or 500
	Secobarbital (BAR)	300
	Buprenorphine (BUP)	10
	Oxazepam (BZO)	300
	Cocaine (COC)	150 or 300
	2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP)	300
	Methylenedioxymethamphetamine (MDMA)	500
	Methamphetamine (MET)	1000 or 500
	Morphine (MOP300/OPI2000)	2000 or 300
	Methadone (MTD)	300
	Oxycodone (OXY)	100
	Phencyclidine (PCP)	25
	Propoxyphene(PPX)	300
	Nortriptyline (TCA)	1000
	Cannabinoids (THC)	50
	6-Monoacetylmorphine(6-MAM)	10
	Fentanyl (FYL)	1
	Tramadol (TRA)	100
Configurations	Test cup	Cup
Test Strips	MET, AMP, MOP/OPI, and TCA strips differ	MET, AMP, MOP/OPI, and TCA strips

10. Standard/Guidance Document Reference (if applicable)

None referenced.

11. Test Principle

AllTest Multi-Drug Rapid Urine Test Cup or AllTest Multi-Drug Urine Test Cup is a competitive

immunoassay that is used to screen for the presence of various drugs and drug metabolites in urine. It is chromatographic absorbent device in which, drugs within a urine sample, competitively combined to a limited number of drug monoclonal antibody (mouse) conjugate binding sites.

When the test is activated, the urine is absorbed into each test strip by capillary action, mixes with the respective drug monoclonal antibody conjugate, and flows across a pre-coated membrane. When drug within the urine sample is below the detection level of the test, respective drug monoclonal antibody conjugate binds to the respective drug-protein conjugate immobilized in the Test Region (T) of the test strip. This produces a colored Test line in the Test Region (T) of the strip, which, regardless of its intensity, indicates a negative test result.

When sample drug levels are at or above the detection level of the test, the free drug in the sample binds to the respective drug monoclonal antibody conjugate, preventing the respective drug monoclonal antibody conjugate from binding to the respective drug-protein conjugate immobilized in the Test Region (T) of the device. This prevents the development of a distinct colored band in the test region, indicating a preliminary positive result.

To serve as a procedure control, a colored line will appear at the Control Region (C) of each strip, if the test has been performed properly.

12. Performance Characteristics

A. Analytical performance

a. Precision/Reproducibility:

Precision studies were carried out for samples with concentrations of +100% cutoff, +75% cutoff, +50% cutoff, +25% cutoff, cutoff, -25% cutoff, -50% cutoff, -75% cut off and -100% cutoff. Other samples were prepared by spiked target drug in drug-free urine samples. Each drug concentration was confirmed by LC-MS/MS. For each concentration, tests were performed two runs per day for 25 days using three lots of test cups. The results obtained are summarized in the following table for Amphetamine (AMP 1000), Benzoylcegonine (COC 300), Methamphetamine (MET 1000), Morphine (OPI/MOP 2000), 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP), Tramadol (TRA), Propoxyphene (PPX) and 6-monoacetylmorphine (6-MAM). The precision for the remaining analytes is supported by data in the previously cleared submission K241428.

MOP 2000

Concentration (ng/mL)	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
Lot Number	4211.662	3462.759	3030.531	2431.941	2132.487	1511.933	1015.242	515.108	0.000
Lot 1	0-/50+	0-/50+	0-/50+	0-/50+	22-/28+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	1-/49+	23-/27+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	0-/50+	22-/28+	50-/0+	50-/0+	50-/0+	50-/0+

EDDP

Concentration (ng/mL)	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
Lot Number	587.020	517.618	487.965	355.764	294.641	223.848	157.757	81.993	0.000
Lot 1	0-/50+	0-/50+	0-/50+	0-/50+	22-/28+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	0-/50+	21-/29+	49-/1+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	0-/50+	23-/27+	50-/0+	50-/0+	50-/0+	50-/0+

COC 300

Concentration (ng/mL)	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
Lot Number	609.678	515.747	451.622	376.434	299.386	232.496	149.316	81.012	0.000
Lot 1	0-/50+	0-/50+	0-/50+	0-/50+	21-/29+	49-/1+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	0-/50+	22-/28+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	0-/50+	23-/27+	50-/0+	50-/0+	50-/0+	50-/0+

TRA

Concentration (ng/mL)	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
Lot Number	204.718	169.671	149.969	129.661	97.564	74.650	47.592	26.174	0.000
Lot 1	0-/50+	0-/50+	0-/50+	0-/50+	21-/29+	49-/1+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	1-/49+	22-/28+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	0-/50+	23-/27+	50-/0+	50-/0+	50-/0+	50-/0+

PPX

Concentration (ng/mL)	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
Lot Number	558.684	559.396	472.685	352.338	311.349	206.969	158.805	69.773	0.000
Lot 1	0-/50+	0-/50+	0-/50+	0-/50+	23-/27+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	0-/50+	21-/29+	49-/1+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	0-/50+	21-/29+	50-/0+	50-/0+	50-/0+	50-/0+

AMP 1000

Concentration (ng/mL)	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
Lot Number	2025.713	1760.840	1497.784	1226.200	1063.069	739.120	487.230	250.343	0.000
Lot 1	0-/50+	0-/50+	0-/50+	0-/50+	23-/27+	49-/1+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	0-/50+	22-/28+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	1-/49+	23-/27+	50-/0+	50-/0+	50-/0+	50-/0+

MET 1000

Concentration (ng/mL) Lot Number	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
	1976.054	1710.253	1592.753	1244.352	1072.277	702.076	486.231	242.057	0.000
Lot 1	0-/50+	0-/50+	0-/50+	0-/50+	21-/29+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	0-/50+	20-/30+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	1-/49+	24-/26+	50-/0+	50-/0+	50-/0+	50-/0+

6-MAM

Concentration (ng/mL) Lot Number	+100% cutoff	+75% cutoff	+50% cutoff	+25% cutoff	Cutoff	-25% cutoff	-50% cutoff	-75% cutoff	-100% cut-off
	21.827	16.871	15.734	12.237	10.188	7.566	4.659	2.503	0.000
Lot 1	0-/50+	0-/50+	0-/50+	1-/49+	23-/27+	50-/0+	50-/0+	50-/0+	50-/0+
Lot 2	0-/50+	0-/50+	0-/50+	0-/50+	24-/26+	48-/2+	50-/0+	50-/0+	50-/0+
Lot 3	0-/50+	0-/50+	0-/50+	0-/50+	25-/25+	49-/1+	50-/0+	50-/0+	50-/0+

b. Linearity/assay reportable range:

Not applicable. This device is intended for qualitative use only.

c. Stability:

The device is stable at 2-30°C for 24 months based on real time stability study.

d. Analytical specificity/Interference:

To test the specificity, drug metabolites and other components that are likely to cross-react in urine samples were spiked into drug-free urine. These urine samples were tested using three lots of the device. The results obtained are summarized in the following table for Amphetamine (AMP 1000), Benzoylcegonine (COC 300), Methamphetamine (MET 1000), Morphine (OPI/MOP 2000), 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP), Tramadol (TRA), Propoxyphene (PPX) and 6-monoacetylmorphine (6-MAM). The specificity for the remaining analytes is supported by data in the previously cleared submission K241428.

Percent cross-reactivity, provided in the below table, was calculated as the cutoff concentration divided by the concentration of analyte tested that yielded a positive result, multiplied by 100.

Drug/Cutoff	Compound	Minimum concentration required to obtain a positive result (ng/mL)	% Cross- Reactivity
MOP 2000	Morphine	2000	100%
	Codeine	5000	40%
	Codeine-6-β-D-glucuronide	>100000	<2%

	Hydrocodone	10000	20%
	Hydromorphone	5000	40%
	Oxycodone	>100000	<2%
	Procaine hydrochloride	>100000	<2%
	Oxymorphone	>100000	<2%
	Normorphine	>100000	<2%
	Ethylmorphine	500	400%
	Norcodeine	50000	4%
	Thebaine	>100000	<2%
	Acetylcodeine	>100000	<2%
	Dihydrocodeine	30000	6.7%
	Nalorphine	2000	100%
	Levorphanol tartrate	10000	20%
	Morphine-3-β-D-glucuronide	5000	40%
	s-Monoacetylmorphine	10000	20%
	6-Monoacetylmorphine (6-MAM)	10000	20%
	Norpropoxyphene	>100000	<2%
	Diacetylmorphine (Heroin)	4000	50%
	6-Acetylmorphine	10000	20%
	Morphine-6β-D-glucuronide	5000	40%
EDDP	EDDP	300	100%
	Doxylamine	>100000	<0.3%
	Methadone	>100000	<0.3%
	Methadol	>100000	<0.3%
COC 300	Benzoylecgonine	300	100%
	Cocaethylene	500	60%
	Cocaine hydrochloride	250	120%
	Ecgonine	>100000	<0.3%
	Norcocaine	>100000	<0.3%
	Ecgonine methyl ester	>100000	<0.3%
AMP 1000	Hydroxyamphetamine	10000	10%
	(+/-)- Methylenedioxyamphetamine(MDA)	50	2000%
	D,L-Amphetamine	1000	100%
	D-Amphetamine	1000	100%

	Diethylstilbestrol	>100000	<1%
	L-Amphetamine	1000	100%
	Phentermine	1000	100%
	β-Phenylethylamine	>100000	<1%
	Tyramine	>100000	<1%
	p-Hydroxynorephedrine	>100000	<1%
	D,L-Norephedrine	>100000	<1%
	p-Hydroxyamphetamine	10000	10%
	D-Methamphetamine	>100000	<1%
	L-Methamphetamine	>100000	<1%
	Ephedrine hydrochloride	>100000	<1%
	(+/-)3,4-Methylenedioxyamphetamine (MDMA)	>100000	<1%
	Phenylpropanolamine	>100000	<1%
	Benzphetamine	>100000	<1%
	L-Ephedrine	>100000	<1%
	L-Epinephrine	>100000	<1%
	D,L-Epinephrine	>100000	<1%
	(+/-)3,4-Methylenedioxyethylamphetamine (MDEA)	>100000	<1%
	D,L-Methamphetamine	>100000	<1%
	Pseudoephedrine	>100000	<1%
MET 1000	(+/-)3,4-Methylenedioxy-n-ethylamphetamine(MDEA)	10000	10%
	(±)-MDMA	2000	50%
	D-Methamphetamine	1000	100%
	L-Methamphetamine	20000	5%
	Fenfluramine	100000	1%
	p-Hydroxymethamphetamine	500	200%
	D,L-Methamphetamine	500	200%
	β-Phenylethylamine	>100000	<1%
	Mephetermine	50000	2%
	L-Amphetamine	>100000	<1%
	Methoxyphenamine hydrochloride	>100000	<1%

	D-Amphetamine	>100000	<1%
	D,L-Amphetamine	500	200%
	Chloroquine	50000	2%
	Ephedrine hydrochloride	>100000	<1%
	(+/-)3,4-Methylenedioxyamphetamine (MDA)	>100000	<1%
	Trimethobenzamide	>100000	<1%
	L-phenylephrine	>100000	<1%
	(1R,2S)-(-)-Ephedrine	>100000	<1%
	Procaine hydrochloride	>100000	<1%
	Phentermine	>100000	<1%
	Pseudoephedrine	>100000	<1%
TRA 100	Tramadol	100	100%
	N-Desmethyl-cis-tramadol	200	50%
	O-Desmethyl-cis-tramadol	1000	10%
	Venlafaxine	100000	0.1%
	(±)-O-Desmethylvenlafaxine	100000	0.1%
PPX 300	(+)-Propoxyphene	300	100%
	(+)-Norpropoxyphene	500	60%
6-MAM 10	6-Monoacetylmorphine	10	100%
	Hydrocodone	>100000	<0.01%
	Hydromorphone	>100000	<0.01%
	Morphine	>100000	<0.01%
	Oxymorphone	>100000	<0.01%
	Procaine	>100000	<0.01%
	Thebaine	>100000	<0.01%
	Diacetylmorphine (heroin)	300	3.3%
	Acetylcodeine	>100000	<0.01%
	Buprenorphine	>100000	<0.01%
	Dihydrocodeine	>100000	<0.01%
	Nalorphine	>100000	<0.01%
	Mitragynine (kratom)	>100000	<0.01%
	Norbuprenorphine	>100000	<0.01%
	s-Monoacetylmorphine	10	100%
	Codeine	>100000	<0.01%

Ethylmorphine	>100000	<0.01%
Levorphanol tartrate	>100000	<0.01%
Morphine-3-β-D-glucuronide	>100000	<0.01%
Norcodeine	>100000	<0.01%
Normorphine	>100000	<0.01%
Oxycodone	>100000	<0.01%
Dextromethorphan	>100000	<0.01%
Imipramine hydrochloride	>100000	<0.01%
Levacetylmethadol (LAAM)	>100000	<0.01%
Meperidine	>100000	<0.01%
(±)-Methadone	>100000	<0.01%
Naloxone hydrochloride	>100000	<0.01%
Naltrexone hydrochloride	>100000	<0.01%
Naproxen	>100000	<0.01%
Noroxycodone HCL	>100000	<0.01%
Noroxymorphone HCL	>100000	<0.01%
(+)-Norpropoxyphene maleate	>100000	<0.01%
Oxymorphone-3β-D-glucuronide	>100000	<0.01%
Tapentadol HCl	>100000	<0.01%
Tramadol hydrochloride	>100000	<0.01%
Chlordiazepoxide	>100000	<0.01%
Clobazam	>100000	<0.01%
D-Amphetamine	>100000	<0.01%
(±)-Amphetamine	>100000	<0.01%
Morphine-6β-D-glucuronide	>100000	<0.01%
Norbuprenorphine glucuronide	>100000	<0.01%
Norhydrocodone HCl	>100000	<0.01%

To evaluate potential interference, non-structurally related compounds were added to drug-free urine and to urine samples containing the target drugs at 50% below and 50% above each corresponding cutoff.

Compounds that show no interference at a concentration of 100µg/mL or specified concentrations are summarized in the following table.

Acetaminophen	D-Pseudoephedrine	Norfentanyl
Acetone (1000mg/dL)	Duloxetine	Noscapine
Acetophenetidin	4-Dimethyl-aminoantipyrine	(+)-Naproxen
Acetylsalicylic Acid	5, 5-Diphenylhydantoin	19-Norethindrone

Acyclovir	Ecgonine methyl ester	Octopamine
Albumin (100mg/dL)	EMDP	O-Hydroxyhippuric acid
Albuterol sulfate (Proair HFA)	Ephedrine	Olanzapine
Alpha Methadol	Erythromycin	Omeprazole
Aminophylline	Esomeprazole Magnesium (except MTD test)	Oxalic acid (100 mg/dL)
Aminopyrine	Estradiol	Oxolinic acid
Amoxicillin	Estrone	Oxymetazoline
Ampicillin	Ethanol (1%)	Paliperidone
Aripiprazole	Fenofibrate	Papaverine
Aspartame	Fenoprofen	Penicillin-G
Aspirin	Fentanyl(except FYL test)	Penicillin V Potassium
Atomoxetine	Fluoxetine Hydrochloride	Perphenazine
Atorvastatin Calcium	Fluphenazine	Phenacetin
Atropine	Fotemustine	Phenelzine
Azithromycin	Furosemide	Phenethylamine
Baclofen	Gabapentin	Phenylethylamine
Benzilic acid	Galactose(10mg/dL)	Phenylpropanolamine
Benzocaine	Gamma Globulin(500mg/dL)	Prednisone
Benzoic Acid	Gatifloxacin	Pregablin
Benzphetamine	Gemfibrozil	Procaine
Bilirubin	Gentisic acid	Promazine
Boric Acid (1%)	Glucose(3000mg/dL)	Promethazine
Bupropion	Guaiacol glyceryl ether	Propoxyphene (except PPX test)
Caffeine	Hemoglobin	Propranolol
Cannabidiol	Hydralazine	Pseudoephedrine
Captopril	Hydrochlorothiazide	Pyridoxine
Carbamazepine	Hydrocortisone	Pyrilamine
Carfentanil (except FYL test)	3-Hydroxytyramine	Pyrogallol
Carisoprodol	Ibuprofen	Quetiapine
Cefradine	Isoxsuprine	Quinine
Cephalexin	(+/-)-Isoproterenol	Quinolinic Acid
Chloralhydrate	Ketamine	R(-)-Apomorphine
Chloramphenicol	Ketoprofen	Ranitidine
Chlordiazepoxide (except BZO test)	LAAM HCl	Riboflavin (10mg/dL)
Chloroquine (except MET test)	Labetalol	Rifampicin
Chlorothiazide	L-Ascorbic Acid	Risperidone
Chlorpromazine	L-Ephedrine	Salicylic Acid
Cholesterol	L-Epinephrine	Serotonin(5- Hydroxytyramine)
Ciprofloxacin Hydrochloride	Levofloxacin Hydrochloride	Sertraline
Citalopram	Levonorgestrel	Sildenafil Citrate
Clarithromycin	Levothyroxine Sodium	Simvastatin
Clonidine	Lidocaine Hydrochloride	Sulfamethazine
Clozapine	Lisinopril	Sulindac
Conjugated Estrogens	Loperamide	Telmisartan
Cortisone	Loratadine	Tetracycline
Creatine Hydrate	L-phenylephrine	Tetrahydrocortisone 3-(β-Dglucuronide)
Creatinine	Magnesium	Tetrahydrocortisone, 3-acetate
Cyclobenzaprine	Maprotiline	Tetrahydrozoline
Cyclodextrin	Meperidine	Theophylline
(-)-Cotinine	Meprobamate	Thiamine
(+)-Chlorpheniramine	Methapyrilene	Thioridazine
D,L-Epinephrine	Methaqualone	Tramadol (except TRA test)

D,L-Octopamine	Methoxyphenamine (except AMP/MET test)	Triamterene
d,l-Propranolol	Methylphenidate	Trifluoperazine
D,L-Tryptophan	Metoprolol Tartrate	Trimethobenzamide
D,L-Tyrosine	Metronidazole (300ug/ml)	Trimethoprim
Delorazepam (except BZO test)	Mifepristone	Tryptamine
Deoxycorticosterone	N-Acetylprocainamide	Tyramine
Desloratadine	NaCl(4000mg/dL)	Urea (2000mg/dL)
Dextromethorphan	Nalidixic Acid	Uric Acid
Diclofenac	Naloxone hydrochloride (except OXY test)	Valproic acid (250ug/ml)
Diflunisal	Naltrexone	Venlafaxine HCl (except TRA test)
Digoxin	Niacinamide	Verapamil
Diphenhydramine HCl	Nicotine	Vitamin B2
Disopyramide (except MTD test)	Nicotinic Acid	Vitamin C
Dopamine HCl	Nifedipine	Zaleplon
Doxepin	Nitroglycerin	Zomepirac sodium salt
Doxylamine	Nordoxepin	

Interference by pH and specific gravity were also evaluated using pooled urine specimens with concentrations of 0 (drug-free), at 50% below and 50% above each corresponding cutoff. The results demonstrated that pH levels of 4 to 9 and specific gravity levels of 1.000 to 1.035 do not affect the results of the assays.

B. Method comparison study

The method comparison studies for the device were performed in-house with three operators. Operators ran 80 (40 negative and 40 positive) unaltered urine clinical samples for each drug. The samples were blind labeled and compared to LC-MS/MS results. The results are presented in the table below for Amphetamine (AMP 1000), Benzoylcegonine (COC 300), Methamphetamine (MET 1000), Morphine (OPI/MOP 2000), 2-ethylidene-1,5-dimethyl-3,3-diphenylpyrrolidine (EDDP), Tramadol (TRA), Propoxyphene (PPX) and 6-monoacetylmorphine (6-MAM). The method comparison for the remaining analytes is supported by data in the previously cleared submission K241428.

Drug test	Test Cup Result		Drug-Free	Low Negative by LC-MS/MS (less than - 50%)	Near Cutoff Negative by LC-MS/MS (Between - 50% and the Cutoff)	Near Cutoff Positive by LC-MS/MS (Between the cutoff and +50%)	High Positive by LC-MS/MS (greater than +50%)
MOP 2000	Operator A	+	0	0	0	13	27
		-	12	15	13	0	0
	Operator B	+	0	0	0	12	27
		-	12	15	13	1	0
	Operator C	+	0	0	1	13	27
		-	12	15	12	0	0
	Operator	+	0	0	0	13	26

AMP 1000	A	-	12	12	16	1	0
		+	0	0	1	13	26
	Operator B	-	12	12	15	1	0
		+	0	0	1	14	26
	Operator C	-	12	12	15	0	0
		+	0	0	0	10	29
MET 1000	Operator A	-	12	11	17	1	0
		+	0	0	1	11	29
	Operator B	-	12	11	16	0	0
		+	0	0	1	10	29
	Operator C	-	12	11	16	1	0
		+	0	0	1	10	29
COC 300	Operator A	-	12	10	17	0	0
		+	0	0	1	10	30
	Operator B	-	12	10	18	1	0
		+	0	0	1	9	30
	Operator C	-	12	10	17	1	0
		+	0	0	1	9	30
EDDP	Operator A	-	12	15	12	1	0
		+	0	0	1	12	27
	Operator B	-	12	15	12	0	0
		+	0	0	1	13	27
	Operator C	-	12	15	13	0	0
		+	0	0	0	13	27
TRA	Operator A	-	12	14	13	1	0
		+	0	0	1	10	29
	Operator B	-	12	14	13	0	0
		+	0	0	0	11	29
	Operator C	-	12	14	14	1	0
		+	0	0	0	10	29
PPX	Operator A	-	12	13	14	0	0
		+	0	0	1	10	30
	Operator B	-	12	13	15	0	0
		+	0	0	0	10	30
	Operator C	-	12	13	14	1	0
		+	0	0	1	9	30
6-MAM	Operator A	-	12	11	16	1	0
		+	0	0	1	13	26
	Operator B	-	12	11	16	0	0
		+	0	0	1	14	26
	Operator C	-	12	11	16	1	0
		+	0	0	1	13	26

Discordant Results are summarized below.

Drug	Operator	Sample Number	LC/MS/MS	Discordant
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			Result (ng/mL)	Device Result
MOP 2000	B	S1854	2061.2	-
	C	S1836	1853.5	+
AMP 1000	A	S1667	1070.8	-
	B	S1636	1058.2	-
	B	S1610	923.0	+
	C	S1642	987.8	+
MET 1000	B, C	S1732	974.6	+
	A, C	S1744	1102.9	-
COC 300	A	S1535	297.0	+
	B, C	S1547	313.0	-
	C	S1562	294.6	+
EDDP	A, B	S1927	291.3	+
	A	S1918	300.6	-
TRA	A, B	S2148	91.6	+
	A, C	S2157	103.0	-
PPX	A	S2052	286.3	+
	C	S2017	280.6	+
	C	S2024	311.3	-
6-MAM	C	S2249	9.5	+
	A, B	S2236	9.6	+
	A, C	S2228	10.2	-

C. Lay person study

A lay user study was performed at three intended user sites with 280 lay persons. 117 male and 163 female tested AllTest Multi-Drug Urine Test Cup. They had diverse educational and professional backgrounds and their age range from 20 to > 50. Urine samples were prepared at the following concentrations; -100%, +/-75%, +/-50%, +/-25% of the cutoff by spiking drug(s) into drug free-pooled urine specimens. The concentrations of the samples were confirmed by LC-MS/MS. Each sample was aliquoted into individual containers and blind-labeled. Each participant was provided with the package insert, 1 blind labeled sample and a device. The results are summarized below.

Result of AllTest Multi-Drug Urine Test Cup: Configuration 1

Drug	Cutoff (ng/mL)	Results	Concentration						
			-100% cutoff	-75% cutoff	-50% cutoff	-25% cutoff	+25% cutoff	+50% cutoff	+75% cutoff
AMP	1000	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
BAR	300	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%

BZO	300	Negative	20	20	20	19	2	0	0
		Positive	0	0	0	1	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	90%	100%	100%
BUP	10	Negative	20	20	20	19	2	0	0
		Positive	0	0	0	1	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	90%	100%	100%
COC	300	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
MDMA	500	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
MET	1000	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
MOP	2000	Negative	20	20	20	19	2	0	0
		Positive	0	0	0	1	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	90%	100%	100%
MTD	300	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%
OXY	100	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
PCP	25	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%
TCA	1000	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
THC	50	Negative	20	20	20	19	2	0	0
		Positive	0	0	0	1	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	90%	100%	100%
FYL	1	Negative	20	20	20	18	0	0	0

		Positive	0	0	0	2	20	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	100%	100%	100%
6-MAM	10	Negative	20	20	20	19	2	0	0
		Positive	0	0	0	1	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	90%	100%	100%
TRA	100	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
PPX	300	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
EDDP	300	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%

Result of AllTest Multi-Drug Urine Test Cup: Configuration 2

Drug	Cutoff (ng/mL)	Results	Concentration						
			-100% cutoff	-75% cutoff	-50% cutoff	-25% cutoff	+25% cutoff	+50% cutoff	+75% cutoff
AMP	500	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
BAR	300	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%
BZO	300	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
BUP	10	Negative	20	20	20	19	2	0	0
		Positive	0	0	0	1	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	90%	100%	100%
COC	150	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%
MDMA	500	Negative	20	20	20	19	1	0	0

		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
MET	500	Negative	20	20	20	19	2	0	0
		Positive	0	0	0	1	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	90%	100%	100%
MOP	300	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
MTD	300	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
OXY	100	Negative	20	20	20	19	0	0	0
		Positive	0	0	0	1	20	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	100%	100%	100%
PCP	25	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
TCA	1000	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%
THC	50	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%
FYL	1	Negative	20	20	20	18	0	0	0
		Positive	0	0	0	2	20	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	100%	100%	100%
6-MAM	10	Negative	20	20	20	18	2	0	0
		Positive	0	0	0	2	18	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	90%	100%	100%
TRA	100	Negative	20	20	20	19	1	0	0
		Positive	0	0	0	1	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	95%	95%	100%	100%
PPX	300	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20

		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%
EDDP	300	Negative	20	20	20	18	1	0	0
		Positive	0	0	0	2	19	20	20
		Total	20	20	20	20	20	20	20
		Agreement (%)	100%	100%	100%	90%	95%	100%	100%

Participants were given surveys on the ease of understanding the instruction for use. All participants indicated that the device instruction is easy to understand and follow. A Flesch-Kincaid reading analysis was performed on each package insert and the scores revealed a reading Grade Level of 7.

Clinical Studies:

Not applicable.

13. Conclusion

Based on the test principle and performance characteristics of the device including precision, cut-off, interference, specificity, method comparison and lay-user studies of the devices, it's concluded that AllTest Multi-Drug Rapid Urine Test Cup and AllTest Multi-Drug Urine Test Cup are substantially equivalent to the predicate device.