INTRODUCTION
The “See Now” Methamphetamine (MET) Test is intended for the qualitative detection of the presence of MET and its metabolites in urine at or above the cutoff level of 1000 ng/ml. The device is designed for professional use. This assay provides only a preliminary result.

SUMMARY OF THE TEST
Methamphetamine is among the five most common illicit drugs. As a central nervous system stimulant, Methamphetamine produces potent dopaminergic and sympathomimetic effects, including euphoria, improved cognitive and sensory performance, generalized improvement in mood, increased physical endurance, and appetite suppression1–4. Both d and L forms of the isomers are controlled substances, and the mandated allowable level for methamphetamine is set at 1000 ng/ml in urine. Methamphetamine is excreted primarily in urine, with little biliary excretion of the parent drug or metabolites5. The urinary pH plays an important role in the excretion of methamphetamine6–12. The percentage of the dose excreted as parent drug can range from as low as 2% in alkaline (pH ≥8.0) to 76% in acidic urine (pH ≤5.0). In normal urine (pH 6–8), 37–54% of a dose is excreted as parent drug and 4–7% as amphetamine. The principal of the “See Now” Methamphetamine Test is a solid phase, competitive inhibition immuno-chromatographic assay, in which a chemically labeled drug (drug conjugate) competes with the drug that may be present in urine, for limited antibody binding sites. When the absorbent pad is soaked with urine, the urine will migrate via capillary action toward the test window where the test reaction occurs. A negative specimen produces two distinct color bands, one in the test zone and one in the control zone. A positive specimen produces only one color band in the control zone.

SPECIMEN COLLECTION AND STORAGE
- Urine specimen may be collected at any time in a clean, dry container without preservatives.
- If specimen cannot be assayed immediately, they can be stored at 2–8°C for up to 72 hours prior to testing or frozen at –20°C for longer period of time.
- Specimens should be equilibrated to room temperature before testing if they were refrigerated or frozen.
- Urine specimens exhibiting visible precipitates should be filtered, centrifuged, or allowed to settle so that clear aliquots can be obtained for testing.

TEST PROCEDURE
- Remove the test device from pouch when ready to perform the test. Label the test device with patient or control identification.
- Remove the test device from the sealed pouch by tearing at the notch. Then place the testing device on a level surface.
- Holding the sample dropper vertically, add 5 drops (0.2 ml) of specimen without air bubbles into the sample well.
- For strip test, immerse the strip into the urine cup and take out the strip after 10 sec. Lay the strip on a flat, clean, dry, non-absorbent surface.
- Read the results at 10 minutes. Ensure that the background of the test area is white before interpreting the result.

INTERPRETATION OF RESULTS

Positive
- Only one color band appears at the control region. No apparent band at the test region. This indicates that drug presence is above the cutoff concentration.

Negative
- Two distinct color bands appear at the control and test regions. This indicates that there is no drug in the sample or drug presence is below the cutoff concentration.

Invalid
- No visible band at the control region. Repeat with a new test kit. If test still fails, please contact the distributor with the lot number.
- Note: A faint line at the test region indicates the drug in sample is near the cut-off level for the test. These samples should be re-tested or confirmed with a more specific method before a clinical determination is made.

STORAGE AND STABILITY
The test kit can be stored at temperature (2 to 30°C) in the sealed pouch to the date of expiration. The test kit should be kept away from direct sunlight, moisture and heat.

PRECAUTION
- FOR IN VITRO DIAGNOSTIC USE ONLY.
- Don’t use it after the expiration date.
- The test device should not be re-used.

PERFORMANCE CHARACTERISTICS
- Sensitivity
The “See Now” Methamphetamine test detects Methamphetamine and its metabolites in urine at concentrations equal to or greater than 1000 ng/ml.
- Specificity
The “See Now” Methamphetamine test relative specificity is 99%.
- Accuracy
Accuracy of the “See Now” Methamphetamine test has been evaluated. A total of 80 clinic samples tested (40 negative and 40 positive). The two assays gave an overall of 98.5%.
- Reproducibility
The precision was determined by replicate assays of both positive and negative urine samples with devices from three different production lots. The resultant data indicated no appreciable inter lot variation when testing both positive and negative samples across three different lots of devices.
“See Now” Methamphetamine (MET) Cassette Test
In human oral fluid specimens
For in vitro Diagnosis Use
Product Code: SN 7.6.S

INTRODUCTION
The “See Now” Methamphetamine (MET) cassette test is an immunochromatography based one step in vitro test. It is designed for qualitative determination of methamphetamine in human oral fluid specimens (saliva). This assay provides only a preliminary analytical test result.

SUMMARY OF THE TEST
Methamphetamine (MET) and its metabolites are potent sympathomimetic agents. Acute higher doses lead to enhanced stimulation of the central nervous system and symptoms include euphoria, alertness, and a sense of increased energy and power. More acute responses produce anxiety, paranoia, psychotic behavior, and cardiac dysrhythmias. The pattern of psychosis which may appear at high doses may be indistinguishable from schizophrenia.

The “See Now” MET cassette test is an immunoassay based on the principle of competitive binding. Drugs that may be present in the oral fluid specimen compete against their respective drug conjugate for binding sites on their specific antibody. During testing, a portion of the oral fluid specimen migrates upward by capillary action. A drug, if present in the oral fluid specimen below its cut-off concentration, will not saturate the binding sites of its specific antibody. The antibody will then react with the drug-protein conjugate and a visible colored line will show up in the test line region of the specific drug strip. The presence of drug above the cut-off concentration in the oral fluid specimen will saturate all the binding sites of the antibody. Therefore, the colored line will not form in the test line region. A drug-positive oral fluid specimen will not generate a colored line in the specific test line region of the strip because of drug competition, while a drug-negative oral fluid specimen will generate a line in the test line region because of the absence of drug competition.

To serve as a procedural control, a colored line will always appear at the control line region, indicating that proper volume of specimen.

MATERIALS PROVIDED
1. The “See Now” MET cassette test.
2. Instruction for use.

MATERIALS REQUIRED BUT NOT PROVIDED
1. Timer or clock.
2. Oral fluid collection swabs

SPECIMEN COLLECTION AND STORAGE
The test device should be stored at 4 to 30°C and will be effective until the expiration date stated on the package. The product is humidity-sensitive and should be used immediately after being opened. Any improperly sealed product should be discarded.

Donors should avoid placing anything (including food, drink, gum and tobacco products) in their mouth for at least 10 minutes prior to specimen collection. Donors should avoid placing anything (including food, drink, gum and tobacco products) in their mouth for at least 10 minutes prior to specimen collection.

1. Remove the oral fluid collection swab from the sealed pouch and insert the sponge end of the oral fluid collection swab into the mouth. Actively swab the inside of the mouth and tongue to collect oral fluid for a total of 3 minutes until the sponge becomes fully saturated. Gently press the sponge between the tongue and teeth will assist saturation. No hard spots should be felt on the sponge when saturated.
2. Remove the oral fluid collection swab from the mouth.
3. Place the saturated swab into the collection chamber and press the sponge firmly down on the plastic strainer to release as much liquid as possible.
4. Tightly shut the outer cap of the collection tube.

TEST PROCEDURE
Bring tests, specimens, and/or controls to room temperature (15-30°C) before use.
1. Remove the test card from sealed pouch and use it within one hour.
2. Flip open the dropper tip of the collection tube.
3. Invert the collection tube and transfer 2 drops of oral fluid (approximately 80 uL) into the specimen well (S) of the test card. Avoid ping air bubbles in the specimen well.
4. Read results at 10 minutes.

Note: Do not interpret the results after 20 minutes.

INTERPRETATION OF RESULTS
The “See Now” MET cassette test is a qualitative assay. It identifies the drug in human saliva at its cut-off concentration or higher. The concentration of the drug can not be determined by this assay. The test is intended to distinguish negative result from presumptive positive result. All positive results must be confirmed using an alternate method, preferably GC/MS.

Positive:
Two colored bands form. The appearance of two colored bands, one in test line zone and the other in control line zone, indicates negative result for that particular test(s). The negative result does not indicate the absence of drug and their metabolites in the specimen, it only indicates the level of tested drug and their metabolites in the specimen is less than cut-off level.

Negative:
One colored band forms. One colored band appears in control line zone. No colored band is found in test line zone. This is an indication the level of tested drug and their metabolites in the specimen is above the cut-off level.

Invalid:
If there is no colored band in control line zone of any strip, the test result is invalid. Retest the sample with a new device.

Note: A borderline(±) in test line zone should be considered negative result.

LIMITATION OF PROCEDURE
1. The “See Now” MET cassette test provides only a qualitative, preliminary analytical result. A secondary analytical method must be used to obtain a confirmed result. Gas chromatography/mass spectrometry (GC/MS) or gas chromatography/tandem mass spectrometry (GC/MS/MS) is preferred in confirmatory methods.
2. A positive test result does not indicate the concentration of drug in the specimen or the route of administration.
3. A negative result may not necessarily indicate a drug-free specimen. Drug may be present in the specimen below the cutoff level of the assay.

PERFORMANCE CHARACTERISTICS
A. Sensitivity
The cut-off concentration (sensitivity level) of the “See Now” MET cassette test is determined to be: 50 ng/mL.
A phosphate-buffered saline (PBS) pool was spiked with drugs to
B. Precision
The results of 30 samples each of 50% above and 50% below cut-off specimens are 100% agreed by three observers. The test results were found to have no significant differences between these three observers.
C. Specificity
The “See Now” MET saliva cassette test relative specificity it is 99%.