This “Sample Procedure” is not intended as a substitute for your facility’s Procedure Manual or reagent labeling, but rather as a model for your use in customizing for your laboratory’s needs.

Space has been provided within the document to allow you to update this template with information specific to your facility. It is suggested that a current version of the manufacturer’s directional insert be maintained as a supplement.
PROCEDURE

Title: Sekisui Diagnostics OSOM® hCG Urine Test

Prepared by: __________________________  Date: __________________________

Accepted by: __________________________  Date adopted: __________________

Reviewed by: __________________________  Date: __________________________

Discontinued by: ______________________  Date: __________________________
SAMPLE PROCEDURE MANUAL

I. TEST NAME

OSOM® hCG Urine Test

II. INTENDED USE

For the qualitative determination of human chorionic gonadotropin (hCG) in urine for early detection of pregnancy. FOR LABORATORY AND PROFESSIONAL IN VITRO DIAGNOSTIC USE ONLY.

III. SUMMARY AND EXPLANATION OF TEST

The hormone hCG is produced by the placenta. Urine hCG provides an early indication of pregnancy. However, because detectable hCG may be associated with conditions other than pregnancy, such conditions should be ruled out when diagnosing pregnancy.

IV. PRINCIPLES OF TEST

The OSOM hCG-Urine Test uses color immunochromatographic dipstick technology with mouse monoclonal and rabbit polyclonal antibodies coated on the nitrocellulose membrane. The Test Stick is dipped into the urine sample and the sample migrates along the membrane. If hCG is present in the sample, it will form a complex with the anti-hCG antibody conjugated color particles. The complex will then be bound by the anti-hCG capture antibody and a visible blue Test Line along with a red Control Line will appear to indicate a positive result.

V. KIT CONTENTS AND STORAGE

Each kit contains 50 Test Sticks in a container and one directional insert. The test sticks must be stored tightly capped at 15°–30° C (59°–86° F). Do not use Test Sticks after expiration date.

At this facility, kits are stored: ____________________________________________

VI. MATERIALS REQUIRED BUT NOT PROVIDED

Urine containers and a timer or watch.

VII. PRECAUTIONS

Follow your laboratory safety guidelines in the collection, handling, storage and disposal of patient specimens and all items exposed to patient specimens.
VIII. SPECIMEN COLLECTION & PREPARATION

This facility's procedure for patient preparation is: ________________________________

This facility’s procedure for sample labeling is: ________________________________

Urine collected anytime may be used. Urine specimens may be refrigerated (2 °C – 8 °C; 36 °F – 46 °F) and tested within 48 hours. Urine specimens and control material must be at room temperature (15 °C – 30 °C; 58 °F - 86 °F) prior to testing.

This facility’s procedure for transporting specimens is: __________________________

This facility’s procedure for rejected specimens is: ______________________________

IX. QUALITY CONTROL & ASSURANCE

If external quality control testing is desired, Control Set Catalog No. 134 may be purchased separately from Sekisui Diagnostics. Some commercial controls may contain interfering additives, therefore the use of these controls is not recommended. Test the control in the same manner as a patient sample. The red Control Line is an internal positive control; if the test has been performed correctly (e.g. the proper volume of sample was absorbed into the Test Stick) and the Test Stick is working properly (i.e. the antibodies and conjugate are active), this indicator will appear. A clear background is an internal negative control; if the test has been performed correctly and the Test Stick is working properly, the background will clear to give a discernible result. Quality Control requirements should be established in accordance with local, state and federal regulations or accreditation requirements. Minimally, Sekisui Diagnostics recommends that positive and negative external controls be run with each new lot and each untrained operator.

QC Testing Frequency and Documentation

For this facility, external QC is run: ____________________________________________

________________________________________________________________________

Results of External QC and action(s) taken when control results are unacceptable are documented: ______________________________________________________________

________________________________________________________________________

X. TEST PROCEDURE

- Remove Test Stick(s) from the container; re-cap container immediately.
- Dip the Absorbent End of the Test Stick into the sample up to the Sample Line for at least 3 seconds.
- Remove the Test Stick from the sample. Place it on a clean, flat, dry disposable surface such as a paper towel.
- Read results. Positive results can be read as soon as the red Control Line appears. Confirm negative at 5 minutes. Results read after 5 minutes are invalid.
XI. INTERPRETATION OF RESULTS

A. Positive

A blue Test Line and a red Control Line is a positive result and the sample contains hCG. Note that the blue line can be any shade of blue.

B. Negative

A red Control Line but no blue Test Line is a negative result and the sample contains no detectable hCG.

C. Invalid

If after 5 minutes, no red Control Line appears, or background color makes reading the red Control Line impossible, the result is invalid and the test should be repeated with a new Test Stick.

In the event this test becomes inoperable, this facility’s course of action for patient samples is:

________________________________________________________________________

XII. RESULT REPORTING

This facilities procedure for patient reporting is:

________________________________________________________________________

________________________________________________________________________

XIII. LIMITATIONS

1. This assay is capable of detecting only whole molecule (intact) hCG, which is the predominant form of hCG in early pregnancy. It cannot detect the presence of hCG fragments or free subunits.

2. In later term pregnancies (generally beyond the first trimester), occasional urine samples can contain very high levels of hCG fragments. Therefore the OSOM hCG Urine Test is most effective when used for the detection of pregnancy in its earlier stages.

3. For diagnostic purposes, hCG test results should always be used in conjunction with other methods and in the context of the patient’s clinical information (e.g., medical history, symptoms, results of other tests, clinical impression, etc.) Ectopic pregnancy cannot be distinguished from normal pregnancy by hCG measurements alone.4,5
4. If the hCG level is inconsistent with, or unsupported by, clinical evidence, results should also be confirmed by an alternative hCG method. Test results should be confirmed using a quantitative hCG assay prior to the performance of any critical medical procedure.

5. Interfering substances may falsely depress or falsely elevate results. These interfering substances may cause false results over the entire range of the assay, not just at low levels, and may indicate the presence of hCG when there is none. As with any immunochemical reaction, unknown interferences from medications or endogenous substances may affect results.

6. Infrequently, hCG levels may appear consistently elevated and could be due to, but not limited to, the presence of the following:6-9
   a. Trophoblastic or nontrophoblastic neoplasms: abnormal physiological states that may falsely elevate hCG levels10,11. This test should not be used in the diagnosis of these conditions.
   b. hCG like substances

7. Because of the high degree of sensitivity of the assay, specimens tested as positive during the initial days after conception may later be negative due to natural termination of the pregnancy. Overall, natural termination occurs in 22% of clinically unrecognized pregnancies and 31% of other pregnancies12. In the presence of weakly positive results, it is good laboratory practice to sample and test again after 48 hours.

8. If the test line appears very faint, it is recommended that a new sample be collected 48 hours later and tested using another OSOM hCG Urine Test Stick.

9. Dilute urine specimens may not have representative levels of hCG.

10. Detection of very low levels of hCG does not rule out pregnancy6, as low levels of hCG can occur in apparently healthy, nonpregnant subjects13,14. Additionally, post-menopausal specimens may elicit weak positive results due to low hCG levels unrelated to pregnancy. In a normal pregnancy, hCG values double approximately every 48 hours15. Patients with very low levels of hCG should be sampled and tested again after 48 hours, or tested with an alternative method.

XIV. EXPECTED RESULTS

Normally, hCG is not detected in urine of healthy men and healthy non-pregnant women. In normal pregnancy, hCG levels in urine can reach 25 mIU/mL as early as 7 to 10 days post conception, and continue rising to reach a maximum concentration in excess of 200,000 mIU/mL at the end of the first trimester.
XV. CROSS REACTIVITY

The following substances were tested individually in hCG-free urine and urine containing 25mIU/mL hCG. None of the substances, at the levels tested, affected expected results:

hLH (500 mIU/mL), hFSH (1,000 mIU/mL), hTSH (1,000 μIU/mL), Estriol (2 mg/dL), Pregnanediol (2mg/dL), Acetaminophen (20 mg/dL), Acetoacetic Acid (1,000 mg/dL), Acetone (1,000 mg/dL), Acetylsalicylic Acid (20 mg/dL), Albumin (2,000 mg/dL), Ampicillin (20 mg/dL), Ascorbic Acid (20 mg/dL), Atropine (20 mg/dL), Bilirubin (2 mg/dL), Caffeine (20 mg/dL), Gentisic Acid (20 mg/dL), Glucose (2,000 mg/dL), Hemoglobin (1,000 mg/dL), Hydroxybutyric Acid (100 mg/dL), Phenothiazine (20 mg/dL), Phenylpropanolamine (20 mg/dL), Salicylic Acid (20 mg/dL), Tetracycline (20 mg/dL).

XVI. PERFORMANCE CHARACTERISTICS & POL STUDIES

Refer to Directional Insert – OSOM® hCG-Urine Test

XVII. REFERENCES

Refer to Directional Insert – OSOM® hCG-Urine Test

XVIII. ASSISTANCE

For technical assistance, call Sekisui Diagnostics Technical Service at 800-332-1042

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