OneStep
Scrub Typhus IgM Serum
/WB/Plasma
RapiDip™ InstaTest

INTENDED USE
The Cortez Diagnostics Scrub Typhus IgM RapiDip™ InstaTest is a
rapid immunochromatographic immunoassay for the qualitative
detection of IgM antibodies to members of O. tsutsugamushi (OT)
species in human serum, plasma or blood. It is intended for research
use only. Not for use in diagnostic procedures. It is not intended for
use by blood donor centers or blood component manufacturers.

SUMMARY AND EXPLANATION
Scrub Typhus (ST) is an infectious disease that is caused by Orientia
tsutsugamushi (OT; formerly Rickettsia), a tiny parasite about the size
of bacteria that belongs to the family Rickettsiaceae (1). A bite from
the larval trombiculid mite, a parasite of rodents, will transmit the
disease. An ulcer of the skin is characteristic of a bite from a trombiculid mite, followed by symptoms including fever, a spotted rash on the torso, and swelling of the lymph glands. After the onset of symptoms, the IgM antibody titers increased gradually over 2–3
weeks, peaked at about 4 weeks, and started to decrease rapidly
between 4 and 5 weeks. Over the first 2 weeks, IgG antibody titers
increase sharply, peaked at about 4 weeks and decreased rather
gradually thereafter (1). Scrub typhus generally occurs after exposure
to areas with secondary (scrub) vegetation, which is where its name is
derived from. However, the disease can also be prevalent in sandy,
mountainous, and tropical areas. Scrub Typhus is a worldwide illness,
but particular to South East Asia and the Western Pacific. It accounts
for approximately 20% of fever in some regions in South East Asia,
where it is endemic. Illness lasts for a period of 10 to 12 days after the
initial bite. With therapy, the fever will break within 36 hours, but if
left untreated, complications or death may occur.

TEST PRINCIPLE
The Cortez Diagnostics Scrub Typhus IgM RapiDip™ InstaTest is a
qualitative, membrane-based immunoassay for the detection of IgM.
The membrane is pre-coated with a mixture of novel recombinants
(representing several geographical isolates; 2, 3) on the test line region
and appropriate control antigen on the control line region. During
testing, the sample reacts with the dye conjugate (anti-human IgM
colloidal gold conjugate) which has been pre-coated in the test device.
The mixture then migrates upward on the membrane chromatographically by capillary action to react with ST derived recombinant antigens on the membrane and generates a red line.
Presence of this red line indicates a positive result, while its absence
indicates a negative result. Independent of the presence or absence
of antibody to scrub typhus antigens, a red line at the control line region
will always appear when the sera-gold migrates to the immobilized
control region. The presence of this red line verifies for sufficient
sample volume and proper flow of reagents.

SPECIMEN COLLECTION AND PREPARATION
- The Scrub Typhus IgM Test system can be performed using
serum, plasma or whole blood. Note: Do not use heat
inactivated samples.
- Testing should be performed immediately after the specimens
have been collected. Do not leave the specimens at room
temperature for prolonged periods. The plasma or serum may be
stored at 2-8°C for up to 3 days. For long term storage,
specimens should be kept below -20°C.
- Bring specimens to room temperature prior to testing. Frozen
specimens must be completely thawed and mixed well prior to
testing. Specimens should not be frozen and thawed repeatedly.
If specimens are to be shipped, they should be packed in
compliance with federal regulations for transportation of
etiologic agents.

MATERIALS AND COMPONENTS
Materials provided with the test kits
ScrubTyphus IgM rapid test strip’s membrane is pre-coated with a
mixture of ST derived recombinant r-56 antigens on the test line
region and Protein A binding, or anti-human IgM (for IgM detection)
reagents on the control line region.
The Kit contains the following:
1. Twenty-five (25) individually pouched Test Strips or twenty-five
(25) test strips in a vial with desiccant on the vial cap.
2. One (1) vial of Chase Buffer solution.

PRECAUTIONS
- For research use only. Not for use in diagnostic procedures.
- All human source materials used in the preparation of controls
have tested negative for antibodies to HIV 1&2, Hepatitis C and
Hepatitis B surface antigen. However, no test method can ensure
100% efficiency. Therefore, all human controls and antigen
should be handled as potentially infectious material. The Center
for Disease Control and the National Institute of Health
recommend that potentially infectious agents be handled at the
Biosafety Level 2.
- Avoid drinking and/or eating in testing areas.
- A thorough understanding of this package insert is necessary for
successful use of the product. Reliable results will only be
obtained by using precise laboratory techniques and accurately
following the package insert.
- Do not mix various lots of any kit component within an
individual assay.
- Do not use any component beyond the expiration date shown on
its label.
- Avoid exposure of the reagents to excessive heat or direct sunlight
during storage and incubation.
- Wear protective clothing, eye protection and disposable gloves
while performing the assay. Wash hands thoroughly afterwards.
- Use a clean disposable pipette tip for each reagent, Standard,
Control or specimen.
- Cover working area with disposable absorbent paper.
- Avoid all possible contact with skin and mucous membranes.
ASSAY PROCEDURE
1. Allow the samples to reach room temperature prior to testing.
2. Remove the Scrub Typhus IgM test from the foil pouch or vial.
3. Add 10 μl of sera or 1 μl whole blood to the test strip in the area beneath the arrow (sample pad area). For addition of the sera, the strip may be placed horizontally on a flat surface or vertically in a microtiter well or test tube.
4. If the test strip is placed horizontally on a flat surface, add 3 drops (or 90-120 μl) of the Chase Buffer solution into a test tube or plastic well (with a minimum capacity of 200 μl); then place the strip, facing downward (as indicated by arrows on the strip), into the well.
5. If the test strip is placed vertically in the well, then add 3 drops (90-120 μl) of the Chase Buffer solution directly into the well.
6. Read the results in 15 minutes.
7. To ensure the detection of all samples, including those yielding a weak signal in the test region, it is imperative that the testing period be complete and that the background on the strip cleared prior to reading the result. Results interpreted after 15 minutes can be inaccurate.

Note: Do not test this product with the Chase Buffer solution alone. 10 μl of human serum or 10 μl of whole blood must be added first.

RESULTS
A Positive Result
The test is positive when only the control line (C) and test line (T) appears in the test area (Figure 1A). A positive result indicates that the Cortez Scrub Typhus IgM rapid dipstick detected antibodies to mixture of OT derived recombinant antigens. A faint line is considered a positive result. As a guide for interpretation, the red color in the test region (T) will vary depending on the concentration/affinity of antibodies present.

Figure 1
Schematic representation of Cortez Scrub Typhus reactivity

A Negative Result
The test is negative when only the control line appears (Figure 1B). A negative result indicates that the Cortez Scrub Typhus IgM rapid dipstick did not detect antibodies to members of OT derived recombinant antigens.

An Invalid Result
1. No lines appear at either the control or test line areas.
2. No control line appears, but a test line is seen (Figure 1C).

LIMITATIONS OF PROCEDURE
- The Cortez Scrub Typhus IgM rapid Test Device is for Research Use Only. Not for use in diagnostic procedures. The test should be used for the detection of antibodies to OT derived antigens in specimen.
- The Cortez Scrub Typhus IgM rapid Test Device will only indicate the presence of antibodies to OT antigens in the specimen and may not imply active infection.
- Do not use serum samples containing any glycerol or other viscous materials. This will decrease the sensitivity of the assay.
- The Cortez Scrub Typhus assays have not been tested in persons with advanced HIV infection or other immunocompromised diseases that are known to have low or undetectable specific antibodies.
- Rheumatoid Factor containing (RF) sera may produce false positive results.

STORAGE
The sealed pouch or vial containing the test strip and the bottle containing the Chase Buffer is designed to be stored at room temperature (20-28°C) for the duration of its shelf life. Exposure to temperatures over 30°C can impact the performance of the test and should be minimized. The strips should not be frozen. The test should be used within 1 hour after removal from the pouch or vial to prevent exposure to humidity.

REFERENCE