Overview

Useful For
Rapid, sensitive, and specific identification of *Ureaplasma urealyticum* and *U parvum* from genitourinary, reproductive, bone and joint, and lower respiratory sources

This test is not intended for medicolegal use

Method Name
Real-Time Polymerase Chain Reaction (PCR) Using LightCycler and Fluorescent Resonance Energy Transfer (FRET)

NY State Available
Yes

Specimen

Specimen Type
Varies

Necessary Information
Specimen source is required.

Specimen Required
The high sensitivity of amplification by PCR requires the specimen to be processed in an environment in which contamination of the specimen by *Ureaplasma* DNA is not likely.

Submit only 1 of the following specimens:

Supplies:

-M4-RT (T605)

-Culturette (BBL Culture Swab) (T092)

Specimen Type: Swab

Sources: Vaginal, cervix, urethra, urogenital, chest/mediastinal; bronchus (donor swab), or upper respiratory sources (only infants <3 months: nasopharynx, nose, throat)

Container/Tube:

Preferred: Culture swab transport system (Dacron or rayon swab with aluminum or plastic shaft with either Stuart or Amies liquid medium)

Acceptable: Swab in transport media: M4, M4-RT, M5, M6, universal transport media, or ESwab

Specimen Volume: 1 swab
Collection Instructions:

Vaginal:
1. Collect specimen by swabbing back and forth over mucosa surface to maximize recovery of cells.
2. Place swab back into swab cylinder.

Urethra or Cervical:
1. Collect specimen by inserting swab 1 to 3 cm and rotating 360 degrees.
2. Place swab back into swab cylinder.

Wound:
1. Collect specimen by swabbing back and forth over wound surface to maximize recovery of cells.
2. Place swab back into swab cylinder.

Supplies: M4-RT (T605)

Specimen Type: Fluid

Sources: Pelvic, amniotic, prostatic secretions, semen, reproductive drainage or fluid, pleural/chest, chest tube, pericardial, sputum, tracheal secretions, bronchial washings, bronchoalveolar lavage, lung; or nasal washings (only infants <3 months)

Container/Tube:

Preferred: Sterile container

Acceptable: Specimen in 3 mL of transport media: M4, M4-RT, M5, M6, or universal transport media

Specimen Volume: 1-2 mL

Specimen Type: Synovial Fluid

Container/Tube:

Preferred: Lavender top (EDTA)

Acceptable: Pink top (EDTA), royal blue top (EDTA), sterile vial containing EDTA-derived aliquot, red top (no anticoagulant), or sterile container

Specimen Volume: 1 mL

Collection Instructions: Send specimen in original tube (preferred).

Specimen Type: Urine, kidney/bladder stone, or ureter

Container/Tube: Sterile container
Test Definition: URRP
Ureaplasma PCR

Specimen Volume: 10 mL or entire specimen

Specimen Type: Tissue

Sources: Placenta, products of conception, urogenital, respiratory, bronchus, chest/mediastinal, bone, or joint

Container/Tube: Sterile container

Specimen Volume: 5 mm(3)

Collection Instructions:
1. Collect fresh tissue specimen.
2. Submit fresh tissue only, do not add fluid to tissue
3. Refrigerate or freeze specimen.

Forms
If not ordering electronically, complete, print, and send a Microbiology Test Request (T244) with the specimen.

Specimen Minimum Volume
Fluid: 1 mL
Urine: 2 mL
Swab: 1 swab
Tissue: 5 mm(3)

Reject Due To
| Swab/Other | Cotton or calcium alginate-tipped swab, wooden shaft swab, transport swab containing gel or charcoal, formalin-fixed and/or paraffin-embedded tissues, Port-a-Cul tube, anaerobic fluid vials, or dry swab (no pledget or sponge); bone marrow; decalcified bone; slides |

Specimen Stability Information

<table>
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<tr>
<th>Specimen Type</th>
<th>Temperature</th>
<th>Time</th>
<th>Special Container</th>
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<tbody>
<tr>
<td>Varies</td>
<td>Refrigerated (preferred)</td>
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<tr>
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<td>Frozen</td>
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Clinical and Interpretive

Clinical Information

_Ureaplasma urealyticum_ and _U parvum_ have been associated with a number of clinically significant infections, although their clinical significance may not always be clear as they are part of the normal genital flora. _U urealyticum_ and _U parvum_ have been associated with urethritis and epididymitis. They may cause upper urinary tract infection and they have been associated with infected renal stones. _U urealyticum_ and _U parvum_ may be isolated...
Test Definition: URRP
Ureaplasma PCR

from amniotic fluid of women with preterm labor, premature rupture of membranes, spontaneous term labor, or chorioamnionitis. They may also cause neonatal infections, including meningoencephalitis and pneumonia. In addition, *U urealyticum* and *U parvum* have been reported to cause unusual infections, such as prosthetic joint infection and infections in transplant recipients.

Recently, *U urealyticum* and *U parvum* have been found to cause hyperammonemia in lung transplant recipients.(1) In lung transplant recipients with hyperammonemia, the ideal diagnostic specimen is a lower respiratory specimen (e.g., bronchoalveolar lavage fluid), although *U urealyticum* and *U parvum* may also be detected in blood. Treatment directed against these organisms has resulted in resolution of hyperammonemia.

Culture of *Ureaplasma* species is laborious, requiring a high degree of technical skill and taking several days. PCR detection is sensitive, specific, and provides same-day results. In addition, PCR allows the differentiation of *U urealyticum* and *U parvum*, which is not easily accomplished with culture. PCR assay has replaced conventional culture for *U urealyticum* and *U parvum* at Mayo Clinic Laboratories due to its speed and equivalent performance to culture.

Reference Values
Not applicable

Interpretation
A positive PCR result for the presence of a specific sequence found within the *Ureaplasma urealyticum* and *U parvum ureC* gene indicates the presence of *U urealyticum* or *U parvum* DNA in the specimen.

A negative PCR result indicates the absence of detectable *U urealyticum* and *U parvum* DNA in the specimen, but does not rule-out infection as false-negative results may occur due to inhibition of PCR, sequence variability underlying the primers and probes, or the presence of *U urealyticum* or *U parvum* in quantities less than the limit of detection of the assay.

Cautions
Interfering substances may affect the accuracy of this assay; results should always be interpreted in conjunction with clinical and epidemiological findings.

Since *Ureaplasma* species may be part of the normal flora, results should be interpreted accordingly.

Clinical Reference
Performance

Method Description
This PCR method employs a target-specific detection system including primers, as well as fluorescent resonance energy transfer (FRET) hybridization probes designed for ureC gene of *Ureaplasma urealyticum* and *U parvum*. The LightCycler instrument amplifies and monitors target nucleic acid sequences by fluorescence during PCR cycling. This is an automated PCR system that can rapidly detect amplified product development. The detection of amplified products is based on the FRET principle. For FRET product detection, a hybridization probe with a donor fluorophore, fluorescein, on the 3’ end is excited by an external light source, which emits light that is absorbed by a second hybridization probe with an acceptor fluorophore, LC-Red 640, on the 5’ end. The acceptor fluorophore then emits light of a different wavelength that is measured with a signal that is proportional to the amount of specific PCR product. The process is completed in a closed tube system and the melting temperature of the probes allows differentiation of *Ureaplasma urealyticum* from *Ureaplasma parvum*. (Cunningham SA, Mandrekar JN, Rosenblatt JE, Patel R: Rapid PCR Detection of *Mycoplasma hominis*, *Ureaplasma urealyticum*, and *Ureaplasma parvum*. Int J Bacteriol 2013 Jan 30, doi: 10.1155/2013/168742)

PDF Report
No

Day(s) and Time(s) Test Performed
Monday through Friday

Analytic Time
3 days

Maximum Laboratory Time
4 days

Specimen Retention Time
1 week

Performing Laboratory Location
Rochester

Fees and Codes

Fees
- Authorized users can sign in to Test Prices for detailed fee information.
- Clients without access to Test Prices can contact Customer Service 24 hours a day, seven days a week.
- Prospective clients should contact their Regional Manager. For assistance, contact Customer Service.

Test Classification
This test was developed and its performance characteristics determined by Mayo Clinic in a manner consistent with CLIA requirements. This test has not been cleared or approved by the U.S. Food and Drug Administration.

CPT Code Information
87798 x 2

LOINC® Information
### Test Definition: URRP
Ureaplasma PCR

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<td>Ureaplasma parvum PCR</td>
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