Syphilis Guide for Providers*

DEFINITION & TRANSMISSION

Syphilis is a systemic, sexually transmitted disease (STD) caused by the Treponema pallidum bacterium. If left untreated, syphilis progresses in stages.

Syphilis Transmission

Two means of syphilis transmission: sexual and vertical

- **Sexual:** Person to person via vaginal, anal, or oral sex through direct contact with syphilis sores or lesions, known as a chancre. Chancres occur at the primary stage of syphilis and can be found around the external genitals or anus, in the vagina or rectum, or in or around the mouth.
- **Sexual transmission also occurs at the secondary stage, mainly by direct contact with mucous membrane lesions such as condyloma lata and mucous patches.**
- **Vertical:** From infected mother to her unborn baby via the bloodstream.

SCREENING

Providers should routinely test for syphilis in persons who

- are pregnant (at the first prenatal visit, and at the beginning of the third trimester and delivery if risk*);
- are sexually active men who have sex with men (at least annually and more frequently if risk*);
- are living with HIV and are sexually active (annually);
- are otherwise considered to be at increased risk for syphilis*

* Risk is described in the USPSTF’s syphilis screening recommendations and the CDC STD Treatment Guidelines at www.cdc.gov/std/treatment/. Any person with signs or symptoms suggestive of syphilis should be tested for syphilis. Also, anyone with an oral, anal, or vaginal sex partner who has been recently diagnosed with syphilis should be tested for syphilis.

PRIMARY STAGE

Signs and Symptoms

- One or more chancres (usually firm, round, small, & painless but can be atypical, subtle lesions) thought to appear at site of exposure (mainly genital area) ~3 weeks after infection (range 10–90 days).
- Chancres can heal on their own in a few days to weeks, even without treatment.
- Patient is highly infectious, and in utero transmission is likely in pregnant women.

A: Courtesy of Dr. Joseph Engelmann at SJ City Clinic
B: Courtesy of National STD Curriculum
C: Courtesy of CDC, Robert E. Sumpter

FOR MORE INFORMATION

- CDC Syphils Treatment and Care: www.cdc.gov/std/syphilis/treatment.htm
- VDH Provider Resources: http://www.vdh.virginia.gov/disease-prevention/std/resources-forms/
- VDH Health Department Locator: http://www.vdh.virginia.gov/health-department-locator/
- Call 1-800-CDC-INFO
- Health care providers with STD consultation requests can also contact the STD Clinical Consultation Network. This free service is provided by the National Network of STD Clinical Prevention Training Centers & operates 5 days/week. Information is available at www.stdccn.org.
- A web-based self-study syphilis module is available for clinicians at www.std.uw.edu/go/pathogen-based/syphilis/core-concept/all.

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www.cdc.gov/std/syphilis/images.htm

**SECONDARY STAGE**

- Muco-cutaneous lesions (most commonly rashes) can occur as chancre(s) and fade ~6 weeks after infection (range 3 wks–6 mos).
- Rashes may first appear on the palms of hands or the soles of feet, but typically appear on trunk & other areas of the body.
- Lesions such as condyloma lata, a moist, wart-like lesion found in the genital area & mucous patches on the tongue occur in ~25% of patients.
- Other common findings: lymphadenopathy & constitutional symptoms. Less common: patchy alopecia (~10% of patients) & neurologic symptoms (1~2% of patients).
- Symptoms clear within 2–6 wks but may take up to 3 mos, even without treatment.
- Patient is highly infectious, especially if direct contact with a moist lesion. In utero transmission is likely in pregnant women.

**LATENT & TERTIARY STAGES**

**Early Latent:**
- Patient has reactive nontreponemal and treponemal tests within 1 year of onset of infection, but no symptoms.
- Patient is potentially infectious, as signs of primary & secondary syphilis can reoccur and go unnoticed.
- Patient is not infectious in late latent stage but may be in latent of unknown duration if onset of infection was within the past year.

**Late Latent or Latent of Unknown Duration:**
- Patient cannot be determined, but no symptoms.
- Patient is not infectious in late latent stage but may be in latent of unknown duration if onset of infection was within the past year.

**Tertiary:**
- Manifestations in skin and bones (gummas), & cardiovascular system.
- Patient is not infectious.

* In utero transmission can occur during ANY latent stage of syphilis but is more likely in early latent stage.

**NEUROSYPHILIS**

- Neurosyphilis is a site of infection and can occur at any stage of syphilis. The nervous system is infected within hours after infection but it can take weeks or years before symptoms present, if at all.
- All patients with syphilis and neurologic, ophthalmologic, or audiologic symptoms warrant a careful neurological exam and a CSF evaluation via a lumbar puncture (LP).

**NEUROSYPHILIS (cont.)**

- All patients at high risk for syphilis presenting with neurologic symptoms should be tested for syphilis and HIV.
- Early neurosyphilis usually presents a few weeks or a few years after onset of infection, and can occur at the primary and secondary stage. It typically manifests as meningitis that can affect cranial nerves (especially VI, VII and VIII), and as meningo-vascular syphilis which may present with stroke-like symptoms.
- Late neurosyphilis typically presents 10–30 years after onset of infection at the late latent stage. Manifestations include general paresis (chronic meningoencephalitis leading to dementia, muscle weakness and paralysis) or tabes dorsalis (demyelination of the posterior columns of the spinal cord).

**OCULAR SYPHILIS**

- Ocular syphilis can occur at any stage of syphilis.
- Visual complaints consist of vision loss, blurry vision, eye pain, eye redness, etc.
- Patients at high risk for STIs who present with ocular signs or symptoms should be tested for syphilis and HIV.
- Patients with syphilis & ocular symptoms should receive a careful neurological exam, an ophthalmologic assessment, and CSF evaluation via lumbar puncture.
- Treatment should not be delayed while awaiting the results of tests as severe outcomes, including permanent blindness, have been reported.

**CONGENITAL SYPHILIS**

- All pregnant women should be tested for syphilis at their first prenatal visit, as required by law in most states.
- Additional testing at the beginning of the third trimester (28 weeks) & at delivery is indicated if the woman is at increased risk or lives in a community with high syphilis prevalence rates.
- Pregnant women diagnosed with syphilis should be treated with penicillin immediately. Treatment ≥30 days prior to delivery is likely to prevent most cases of congenital syphilis. (It may not prevent stillbirth or congenital syphilis in a gravely infected fetus as evidenced by fetal syphilis on ultrasound at the time of treatment).
- All women who deliver a stillborn infant (after 20 weeks) should be tested for syphilis at time of delivery.

**CONGENITAL SYPHILIS (cont.)**

- All patients at risk for syphilis warrant a thorough physical exam & sexual history.

**DIAGNOSIS**

- Darkfield examinations & other tests (e.g., PCR) to detect T. pallidum directly from lesion exudate or tissue are definitive methods for diagnosing early syphilis and congenital syphilis, though not available in most settings.
- Presumptive diagnosis requires use of 2 serologic tests: a nontreponemal test (i.e., VDRL or RPR) & a treponemal test (i.e., FTA-ABS tests, the TP-PA assay, various EIAs, chemiluminescence immunoassays, immunoblots, or rapid treponemal assays). Persons with a reactive nontreponemal test should always receive a treponemal test to confirm the presumptive diagnosis of syphilis.
- Reverse sequence screening algorithm for syphilis testing is also used. Positive treponemal screening tests are confirmed with a standard nontreponemal test with titer. More information at www.cdc.gov/mmwr/ preview/mmwrhtml/mm6005a1.htm.
- All positive nontreponemal tests must be quantified at time of treatment, as titers are used to monitor treatment success.
- Note: In primary syphilis, nontreponemal serologic tests are only ~75% sensitive.
- Note: In primary & secondary syphilis, “prozone effect” is possible (false negative RPR occurring when high antibody titers prevent antibody/antigen lattice formation).
- All patients with positive syphilis serologic tests & a presumptive diagnosis of syphilis must be staged to determine the recommended treatment regimen. Information needed to determine stage: complete sexual & medical history including history of syphilis testing/treatment; thorough physical exam; epidemiologic information such as risk factors & sexual partner(s) history of syphilis.
- All patients at risk for syphilis warrant a thorough physical exam & sexual history.