

\*Early primary syphilis and late untreated syphilis possible if RPR/VDRL are nonreactive; see below for recommended actions

## Table 1: Interpretation of Syphilis Serologies, Traditional Algorithm Non Non

	Treponemal (RPR/VDRL)	Treponemal (TPPA)	Possible Interpretations		Recommended Actions
		Nonreactive or not done	<ol> <li>No syphilis</li> <li>Early/incubating syphilis (too early to be detected by serology)</li> </ol>	•	If syphilis unlikely, no further action needed. If early syphilis suspected, consider ordering a treponemal test (if not done initially) and repeating an RPR/VDRL in 1-2 weeks; if either test is reactive, treat for syphilis. If concerned for early syphilis (e.g., chancre present or known exposure) treat presumptively. If treating presumptively, repeat RPR/VDRL on day of treatment and, if nonreactive, again in 2-4 weeks to assess for seroconversion.
	Nonreactive	Reactive	<ol> <li>Prior treated syphilis</li> <li>Untreated syphilis</li> </ol>	•	Treponemal tests (e.g., TPPA) often stay reactive for life; if patient has a history of adequate treatment for syphilis & no new exposures/ symptoms, no further action needed. If early syphilis suspected (e.g., chancre present or known exposure), treat presumptively according to stage. If treating presumptively, repeat RPR/VDRL on day of treatment and, if nonreactive, again in 2-4 weeks to assess for seroconversion. If no signs or symptoms, order a second treponemal test (e.g., EIA or CIA); see table 2 for recommendations based on results.
		Nonreactive	1. False positive RPR or VDRL	8	Likely false positive (not syphilis). <sup>b</sup> In pregnancy or in patients at high risk for syphilis, consider rescreening with serologic testing in 2-4 weeks – if unchanged, no action needed. <sup>c</sup>
	Reactive	Reactive	<ol> <li>Current syphilis</li> <li>Treated syphilis with residual/ persistent RPR/VDRL titer</li> </ol>	•	If RPR/VDRL is newly reactive, stage and treat. If previously treated and sustained (≥2 weeks) 4- fold rise in RPR/VDRL titer, manage as treatment failure versus re-infection. <sup>d</sup> Note that RPR/VDRL may still be reactive after treatment; if there is a fourfold decline within 12- 24 months, treatment is considered to have been adequate even if RPR/VDRL remains reactive. Some treated patients may have a persistent low level RPR/VDRL titer for a prolonged period; re- treatment is not necessary in the absence of new exposures or symptoms.

<sup>a</sup> The traditional algorithm starts with a non-treponemal test (RPR or VDRL) which, if reactive, is followed by a confirmatory treponemal test (TPPA). In interpreting

serologies, it is helpful to know which testing algorithm (traditional vs reverse) is being used in your lab.

<sup>b</sup> False positives can be seen in pregnancy and/ in patients with autoimmune diseases, Lyme disease, certain viral infections (including HIV), injection drug use, and other conditions.

<sup>c</sup> In California, <u>all pregnant people should be screened for syphilis three times during pregnancy</u>: (1) at confirmation of pregnancy or first prenatal encounter, (2) early in the third trimester (at approximately 28 weeks gestation or as soon as possible thereafter), and (3) at delivery. The American College of Obstetrics and Gynecology also recommends <u>screening all pregnant patients universally for syphilis three times</u>: once at the first prenatal care visit, again during the third trimester, and again at birth.

<sup>d</sup> For patients determined to have new syphilis or treatment failure, refer to the Centers for Disease Control STD treatment guidelines at <a href="https://www.cdc.gov/std/treatment-guidelines/syphilis.htm">https://www.cdc.gov/std/treatment-guidelines/syphilis.htm</a> for treatment and follow up recommendations.

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## Table 2: Interpretation of Syphilis Serologies, Reverse Screening Algorithm

Immuno- assay (CIA or EIA)	RPR/ VDRL	ТРРА	Possible Interpretations	Recommended Actions
Non- reactive	Non- reactive or not done	Non- reactive or not done	<ol> <li>Syphilis unlikely</li> <li>Early/incubating syphilis (too early to be detected by serology)</li> </ol>	<ul> <li>If syphilis unlikely, no further action needed.</li> <li>If immunoassay nonreactive but high clinical suspicion (such as a chancre or known exposure), treat presumptively for early syphilis. If treating presumptively, obtain RPR/VDRL on day of treatment and, if nonreactive, again in 2-4 weeks to assess for seroconversion.</li> </ul>
Reactive	Non- reactive	Non- reactive or not done	<ol> <li>False positive immunoassay</li> <li>Early/incubating syphilis</li> <li>Latent or prior syphilis (treated or untreated)</li> <li>Latent or prior syphilis (treated or untreated)</li> <li>Early syphilis (prior to RPR/VDRL seroconversion)</li> </ol>	<ul> <li>If no signs/symptoms and low risk for syphilis, most likely a false positive immunoassay.<sup>b</sup> No further action needed.</li> <li>If concerned for early infection or in pregnant patients, re-screen in 2-4 weeks.<sup>c</sup></li> <li>If signs/symptoms or contact to syphilis, treat presumptively. Repeat RPR/VDRL on day of treatment and, if nonreactive, again in 2-4 weeks to assess for seroconversion.</li> <li>No further action needed if patient treated appropriately for syphilis in past, assuming no new exposures/symptoms and a negative clinical exam.</li> <li>If no symptoms and no known prior adequate treatment, treat presumptively for latent syphilis.</li> <li>If early syphilis suspected (symptoms or known exposure), treat presumptively. Obtain DRP (VDR) and a solution of the sector of the secto</li></ul>
	Reactive	Not done or Reactive	<ol> <li>Current syphilis</li> <li>Prior syphilis (treated or untreated)</li> </ol>	<ul> <li>repeat in 2-4 weeks to assess for seroconversion.</li> <li>If RPR/VDRL is newly reactive, stage and treat.</li> <li>If previously treated and sustained (≥2 weeks) 4-fold rise in RPR/VDRL titer, manage as treatment failure versus re-infection.<sup>d</sup></li> <li>If known prior adequate treatment for stage of infection and RPR/VDRL declining appropriately (i.e., a fourfold decline within 12-24 months), no further action needed.</li> <li>Some treated patients may have a persistent low level RPR/VDRL titer for a prolonged period; re- treatment is not necessary in the absence of new exposures or symptoms.</li> </ul>

<sup>a</sup> The reverse algorithm starts with an immunoassay detecting syphilis antibodies which, if reactive, is followed by an RPR/VDRL. If there is a discrepancy between the immunoassay and RPR (one reactive, one nonreactive), a treponemal test (TPPA) serves as the tie-breaker. In interpreting serologies, it is helpful to know which testing algorithm (traditional vs reverse) is being used in your lab. <sup>b</sup> False positive immunoassays can occur with Lyme disease or non-syphilis treponemal infections.

<sup>c</sup> In California, all pregnant people should be screened for syphilis three times during pregnancy: (1) at confirmation of pregnancy or first prenatal encounter, (2) early in the third trimester (at approximately 28 weeks gestation or as soon as possible thereafter), and (3) at delivery. The American College of Obstetrics and Gynecology also rec nends screening all pregnant patients universally for syphilis three times: once at the first prenatal care visit, again during the third trimester, and again at birth.

<sup>d</sup> For patients determined to have new syphilis or treatment failure, refer to the Centers for Disease Control STD treatment guidelines at uidelines/syphilis.htm for treatment and follow up recommendations. http:

