Send reports to:

Arizona Department of Health Services Office of Infectious Disease Services 150 North 18th Avenue, Suite 140 Phoenix, AZ 85007 602-364-3676 or 602-364-3199 (fax) See page 2 for electronic submission

ARIZONA LABORATORY REPORTING REQUIREMENTS

Send isolates or specimens to:

Arizona State Laboratory 250 North 17th Avenue Phoenix, AZ 85007



=	Anaplasma spp.	4O&	Francisella tularensis	=	Plasmodium spp.
)* 4	Arboviruses	3 * ^{4,5}	Haemophilus influenzae, from a normally sterile site)*	Rabies virus from a human
=	Babesia spp.	①	Hantavirus)* 4	Rabies virus from an animal
≙≊ *	Bacillus anthracis	D ¹	Hepatitis A virus (anti-HAV-IgM serologies, detection of viral nucleic acid, or genetic sequencing)	=	Respiratory syncytial virus
)* 4	Bordetella pertussis	<u> </u>	Hepatitis B virus (anti-Hepatitis B core-IgM serologies, Hepatitis B surface or envelope antigen serologies, detection of viral nucleic acid, or genetic sequencing)	҈* ⁴	Rickettsia spp. – any test result
)*	Brucella spp.	<u>≡</u> 1	Hepatitis C virus	3 1*	Rubella virus and anti-rubella-IgM serologies
)*	Burkholderia mallei and B. pseudomallei	<u>≡</u> 1	Hepatitis D virus)*	Salmonella spp.
* 4	Campylobacter spp.	 ■¹*⁴	Hepatitis E virus)* 4	Shigella spp.
≡* ⁴	Carbapenem-resistant Enterobacteriaceae (CRE)	=	HIV—any test result (by culture, antigen, antibodies to the virus, detection of viral nucleic acid, or genetic sequencing), except from a negative screening test		Streptococcus group A, from a normally sterile site
=	CD ₄ -T-lymphocyte count	=	HIV—any test result for an infant (by culture, antigen, antibodies to the virus, detection of viral nucleic acid, or genetic sequencing)	=	Streptococcus group B, from a normally sterile site in an infant younger than 90 days of age
)* ⁴	Chikungunya virus	*	Influenza virus	≅ *⁴	Streptococcus pneumoniae and its drug sensitivity pattern, from a normally sterile site
=	Chlamydia trachomatis)+	Legionella spp. (excluding single serological results)	1	Treponema pallidum (syphilis) or rapid plasma reagin
=	Chlamydia psittaci /Chlamydophila psittaci	①	Leptospira spp.	=	Trypanosoma cruzi (Chagas disease)
<u> </u>	Clostridium botulinum toxin (botulism)	①	Lymphocytic choriomeningitis virus	① *	Vancomycin-resistant or Vancomycin-intermediate Staphylococcus aureus
≡ * ⁴	Coccidioides spp.)*	Listeria spp., from a normally sterile site	₽2	Variola virus (smallpox)
①	Coxiella burnetti	≊ ¹*	Measles virus and anti-measles-IgM serologies)*	Vibrio spp.
①	Cryptosporidium spp.	<u>≡</u> 2	Methicillin-resistant Staphylococcus aureus, from a normally sterile site	₽2	Viral hemorrhagic fever agent
①	Cyclospora spp.	1*	Mumps virus and anti-mumps-IgM serologies	=	West Nile virus
)* 4	Dengue virus)* 3	Mycobacterium tuberculosis complex and its drug sensitivity pattern	☎ *	Yellow fever virus
=	Ehrlichia spp.	≡ **	Neisseria gonorrhoeae and, if performed, the drug sensitivity pattern	≙≊ *	Yersinia pestis (plague)
₽	Emerging or exotic disease agent	*	Neisseria meningitidis, from a normally sterile site)*	Yersinia spp. (other than Y. pestis)
=	Entamoeba histolytica	①	Norovirus)*	Zika virus
) *	Escherichia coli, Shiga toxin-producing	*	Novel coronavirus infection (e.g., SARS or MERS)		

Key:

- Submit a report immediately after receiving one specimen for detection of the agent. Report the receipt of subsequent specimens within five working days after receipt.
- Submit a report within 24 hours after obtaining a positive test result.
- D Submit a report within one working day after obtaining a positive test result.
- Submit a report within five working days after obtaining a positive test result or a test result specified in the above table.
- Submit an isolate of the organism for each positive culture, if available, or a specimen for each positive test result to the Arizona State Laboratory within one working day.
- + Submit an isolate of the organism for each positive culture to the Arizona State Laboratory within one working day.

When appearing after one of the symbols to the left, the following modify the requirement:

- 1 When reporting a positive result for any of the specified tests, report the results of all other tests performed for the subject as part of the disease panel or as a reflex test.
- 2 Submit a report only when an initial positive result is obtained for an individual.
- 3 Submit an isolate or specimen of the organism, as applicable, only when an initial positive result is obtained for an individual, when a change in resistance pattern is detected, or when a positive result is obtained ≥ 12 months after the initial positive result is obtained for an individual.
- 4 Submit an isolate or specimen, as applicable, only by request.
- Submit an isolate of the organism, if available, or a specimen when a positive result is obtained for an individual < 5 years of age.

Arizona Administrative Code R9-6-204 Clinical Laboratory* Director Reporting Requirements

A director of a clinical laboratory that obtains a test result described in the reporting table or that receives a specimen for detection of an infectious agent or toxin listed shall, either personally or through a representative, submit a report, in a Department-provided format, and, if applicable, an isolate or a specimen to the Department within the time limitation and as specified.

For each **specimen** for which an immediate report ((a)) is required, a clinical laboratory director shall ensure the report includes:

- 1. The **name** and **address** of the laboratory;
- 2. The name and telephone number of the director of the clinical laboratory;
- 3. The name and, as available, the address, telephone number, and email address of the subject;
- 4. The date of birth of the subject;
- The gender of the subject;
- 6. The laboratory identification number;
- 7. The **specimen type**;
- 8. The date of collection of the specimen;
- 9. The type of test ordered on the specimen; and
- The ordering health care provider's name, address, telephone number, and, if available, email address.

For each **test result** for a subject for which a report is required, a clinical laboratory director shall ensure the report includes:

- 1. The **name** and **address** of the laboratory;
- The name and telephone number of the director of the clinical laboratory;
- 3. The **name** and, as available, the **address**, **telephone number**, and **email address** of the subject;
- 4. The date of birth of the subject;
- 5. The **gender** of the subject;
- 6. The laboratory identification number;
- 7. The **specimen type**;
- 8. The date of collection of the specimen;
- 9. The date of the result of the test;
- 10. The **type of test completed** on the specimen;
- 11. The **test result**, including quantitative values and reference ranges, if applicable; and
- 12. The ordering health care provider's name, address, telephone number, and, if available, email address

Reports may be submitted via electronic laboratory reporting (ELR), by entry in MEDSIS (https://my.health.azdhs.gov/), or by phone, fax or mail.

Visit http://azdhs.gov/labreporting for information about initiating the ELR engagement process for your laboratory or obtaining MEDSIS access.

Additional reporting resources, including the laboratory reporting form and the list of reportable tests and results, are available at http://azdhs.gov/labreporting

*Clinical laboratory is defined in A.R.S. § 36-451: http://www.azleg.gov/ars/36/00451.htm.