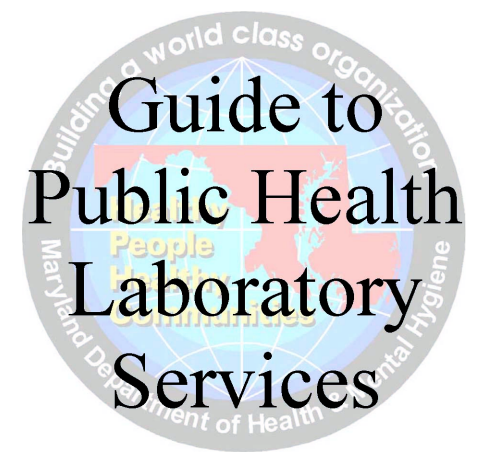
# State of Maryland Department of Health and Mental Hygiene Laboratories Administration



The J. Mehsen Joseph Public Health Laboratory 1770 Ashland Avenue, Baltimore MD 21205 Telephone: 443-681-3800 Fax: 443-681-4501 http://dhmh.maryland.gov/laboratories/

December 2016

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Maryland Department of Health and Mental Hygiene

Larry Hogan, Governor - Boyd Rutherford, Lt. Governor - Van Mitchell, Secretary

Laboratories Administration

Robert A. Myers, Ph.D., Director

May 14, 2015

Dear Health Professionals:

This reference guide lists public health laboratory services available to health officers, physicians, and other health professionals to assist in the prevention, diagnosis, and control of human diseases. The listing of laboratory services is arranged alphabetically by test and includes contact information for the laboratory that performs the test.

Specimens and samples submitted to the central and regional laboratories should be collected and submitted in special kits provided by the Laboratories Administration. These kits may also be obtained from the regional laboratories or county health departments. Use of these kits assures collection of the proper type of specimen, preservation of specimen integrity, proper demographic/epidemiological information, and prompt distribution for examination when received in the laboratory.

Records of patient information and test results are treated as confidential information and will be released only to the submitting physician or other legally authorized individual.

Public Health professionals and physicians using the Administration's services are invited to visit the central laboratory in Baltimore or their regional laboratory. A few minutes spent in the laboratory can often result in clarification of points regarding types of tests performed, specimen kits available, and many other points important to effective use of laboratory services. This personal contact not only improves services but also can be informative to the physician and stimulating to the laboratorian in supporting the practice of modern scientific medicine.

The most up-to-date version of this guide is available for downloading and printing off the internet at: http://dhmh.maryland.gov/laboratories/docs/guide.pdf

(Please note the attached listing of new contact numbers, all extensions will be updated in the future)

Polant A. Maye

Robert A. Myers, Ph.D.

Director

P.O. BOX 2355 • Baltimore, Maryland 21203-2355 410-767-6100 • TTY for Disabled - Maryland Relay Service 1-800-735-2258 Toll Free 1-877-4MD-DHMH • Web Site: http://dhmh.maryland.gov/laboratories/

# GENERAL ORGANIZATION OF THE LABORATORIES ADMINISTRATION

EGISTRATION & LABORATORY REPORTS	
PECIMEN ACCESSIONING LABORATORY	443-681-3793/443-681-3842
PECIMEN KIT PREPARATION UNIT	443-681-3777
OFFICE OF FISCAL ADMINISTRATION: Fax# 443-681-4503	
BILLING OFFICE	443-681-3812
PROCUREMENT OFFICE	443-681-3813
FFICES OF LABORATORY QUALITY ASSURANCE, SAFETY, and TRAINING: Fax# 443-68	31-4503
QUALITY ASSURANCE OFFICER	
TRAINING COORDINATOR	443-681-3792
OFFICE OF SAFETY AND SECURITY	
IVISION OF PUBLIC HEALTH MICROBIOLOGY: Fax# 443-681-4506	
DIVISION CHIEF	443-681-3941
DIVISION MANAGER	
BIOTERRORISM LABORATORY	
	-
DAIRY BACTERIOLOGY/CHEMISTRY	-
FOOD/SHELLFISH	
GC	•
GLASSWARE PREPARATION	•
MEDIA PREPARATION	
MYCOBACTERIOLOGY (TB)	443-681-4569/443-681-3950
PARASITOLOGY	443-681-3952/443-681-3953
WATER MICROBIOLOGY	443-681-3959/443-681-3960
DIVISION CHIEF	
CORE SEQUENCING LABORATORY	443-681-3874
MOLECULAR DIAGNOSTICS LABORATORY	443-681-3924
MOLECULAR EPIDEMIOLOGY LABORATORY	443-681-3879
RETROVIROLOGY LABORATORY	443-681-3877
VIRAL DISEASE ASSESSMENT LABORATORY	443-681-3878
IVISION OF NEWBORN AND CHILDHOOD LABORATORY SCREENING: Fax# 443-681-4	505
DIVISION CHIEF	443-681-3900
NEWBORN SCREENING:	
BIOCHEMICALS	443-681-3913
ENDOCRINOLOGY	443-681-3913/443-681-3912
HEMOGLOBINOPATHIES	
	-
SEVERE COMBINED IMMUNODEFICIENCY DISEASE (SCID)	443-681-3913
SEVERE COMBINED IMMUNODEFICIENCY DISEASE (SCID) TANDEM MASS SPECTROMETRY	443-681-3913 443-681-3915
TANDEM MASS SPECTROMETRY	443-681-3913 443-681-3915
TANDEM MASS SPECTROMETRY	443-681-3913 443-681-3915 443-681-4590/443-681-3910
TANDEM MASS SPECTROMETRY IVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF	443-681-3913 443-681-3915 443-681-4590/443-681-3910 443-681-3930
TANDEM MASS SPECTROMETRY IVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF ARBOVIRUS SEROLOGY	443-681-3913 443-681-3915 443-681-4590/443-681-3910 443-681-3930 443-681-3937
TANDEM MASS SPECTROMETRY IVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF ARBOVIRUS SEROLOGY CHLAMYDIA	443-681-3913 443-681-3915 443-681-4590/443-681-3910 443-681-3930 443-681-3937 443-681-3937
TANDEM MASS SPECTROMETRY VIVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF ARBOVIRUS SEROLOGY CHLAMYDIA HEPATITIS.	443-681-3913 443-681-3915 443-681-4590/443-681-3910 443-681-3930 443-681-3937 443-681-3937 443-681-3889
TANDEM MASS SPECTROMETRY VIVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF ARBOVIRUS SEROLOGY CHLAMYDIA HEPATITIS MICROBIAL SEROLOGY	443-681-3913 443-681-3915 443-681-4590/443-681-3910 443-681-3930 443-681-3937 443-681-3937 443-681-3889 443-681-3938
TANDEM MASS SPECTROMETRY VIVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF ARBOVIRUS SEROLOGY CHLAMYDIA HEPATITIS MICROBIAL SEROLOGY RABIES & ZOONOTIC DISEASES	443-681-3913 443-681-3915 443-681-4590/443-681-3910 443-681-3930 443-681-3937 443-681-3889 443-681-3938 443-681-3772
TANDEM MASS SPECTROMETRY DIVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF ARBOVIRUS SEROLOGY CHLAMYDIA HEPATITIS MICROBIAL SEROLOGY RABIES & ZOONOTIC DISEASES SYPHILLIS & TREPONEMAL SEROLOGY	443-681-3915 443-681-3915 443-681-4590/443-681-3910 443-681-3930 443-681-3937 443-681-3937 443-681-3938 443-681-3938 443-681-3938
TANDEM MASS SPECTROMETRY DIVISION OF VIROLOGY and IMMUNOLOGY: Fax # 443-681-3844 DIVISION CHIEF ARBOVIRUS SEROLOGY CHLAMYDIA HEPATITIS MICROBIAL SEROLOGY RABIES & ZOONOTIC DISEASES SYPHILLIS & TREPONEMAL SEROLOGY VACCINE PREVENTABLE DISEASES	443-681-3913 443-681-3915 443-681-4590/443-681-3910 443-681-3930 443-681-3937 443-681-3937 443-681-3889 443-681-3772 443-681-3938 443-681-3938 443-681-3889
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#### OFFICE OF LABORATORY EMERGENCY PREPAREDNESS and RESPONSE: Fax# 443-681-4509

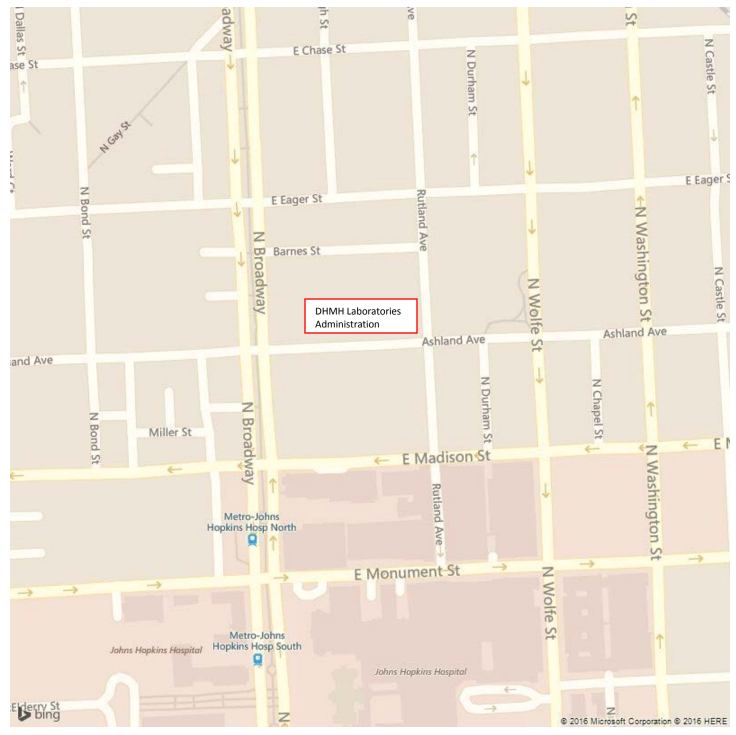
BIOTERRORISM CHIEF	443-681-3787
BIOTERRORISM COORDINATOR	443-681-3788
BIOLOGICAL AGENTS REGISTRY (BAR) PROGRAM	443-681-3789

#### DIVISION OF ENVIRONMENTAL CHEMISTRY: Fax# 443-681-4507

[Refer to "Guide to Environmental Chemistry Laboratory Services" for inf	ormation on testing in this division]
DIVISION CHIEF	443-681-3851
AIR QUALITY SECTION	443-681-3855
CHEMICAL EMERGENCY PREPAREDNESS AND RESPONSE	443-681-3857
ENVIRONMENTAL METALS SECTION	443-681-4596
GENERAL CHEMISTRY SECTION	443-681-3855
NUTRIENTS SECTION	443-681-3855
QUALITY ASSURANCE OFFICE	443-681-3856
RADIATION SECTION	443-681-4596
SEMI-VOLATILES SECTION	443-681-3857
VOLATILES ORGANICS SECTION	443-681-3857

DHMH-Laboratories Administration

The J. Mehsen Joseph Public Health Laboratory



#### A. GENERAL INFORMATION

#### A.1. CENTRAL LABORATORY

Hours: Monday thru Friday 8:00 a.m. – 4:30 p.m. Saturday 7:30 a.m. – 10:30 a.m. Sunday Closed Location: 1770 Ashland Avenue Baltimore, MD 21205 Mailing Address: Laboratories Administration P.O. Box 2355 Baltimore, MD 21203-2355

#### NON-EMERGENCY NUMBERS:

DIRECTOR'S OFFICE	443-681-3800
CENTRAL LABORATORY FAX	443-681-4501
REGISTRATION and LABORATORY REPORTS	443-681-3820
SPECIMEN ACCESSIONING LABORATORY	443-681-3793/443-681-3842

#### 24-HOUR EMERGENCY NUMBERS:

#### **ANIMAL RABIES EMERGENCY EXAMINATION REQUESTS (See page 16)**

# NON-RABIES CASES

NON-RADIES CASES	
LABORATORY EMERGENCY PREPAREDNESS	
AND RESPONSE CELL PHONE: 410	0-925-3121
DIRECTOR'S EMERGENCY CELL PHONE:	
DR. ROBERT MYERS 44	3-928-0925
A.2. REGIONAL PUBLIC HEALTH LABORATORIES HOURS AND LOCATIONS A.2.a. EASTERN SHORE REGIONAL LABORATORY (ESRL-Salisbury):	
Hours: Monday thru Friday 8:00 a.m. – 4:30 p.m.	
Saturday/Sunday Closed	
Location: 926 Snow Hill Road-Cottage 500	
Salisbury, MD 21804-1939	
Director, Robert A. Myers, Ph.D 44	3-928-0925

Director, Robert A. Myers, Ph.D	. 443-928-0925
ESRL Office	410-219-9005
ESRL FAX	410-749-1173

# 24-HOUR EMERGENCY NUMBER: 443-523-5056 (cell-Primary) 443-928-0925 (cell-Backup)

# A.2.b. WESTERN MARYLAND REGIONAL LABORATORY (WMRL – Cumberland):

Hours: Monday thru Friday 8:00 a.m. – 4:30 p.m.	
Saturday/Sunday Closed	
Location: 12503 Willowbrook Road	
The Brook Building, Entrance #6	
Cumberland, MD 21502	
Director, Robert A. Myers, Ph.D	. 443-928-0925
Chief, Jo Ann Flinn	301-759-5115
WMRL FAX	301-777-2021

#### 24-HOUR EMERGENCY NUMBER: 301-268-4468 (cell)

#### A.3. COURIER SERVICE

The Laboratories Administration contracts to provide specimen courier service for many local health departments. Problems concerning the courier service should be reported immediately by calling 443-681-3820.

### A.4. SPECIMEN REJECTION POLICY

The Laboratories Administration's "Specimen/Sample Acceptance and Rejection Criteria" policy helps to assure the accuracy, reliability, and timeliness of laboratory test results by eliminating the testing of unacceptable specimens. When the laboratory determines that a specimen is unacceptable for testing, the laboratory, whenever feasible, notifies the submitter immediately by telephone, confirms the notification in writing, and temporarily retains the specimen for possible future testing (e.g., in cases where additional information provided by the submitter would make the specimen acceptable for testing).

# A.5. BILLING

Questions concerning client billing, laboratory billing, and laboratory reimbursement by the Maryland Medical Assistance Program or other third party payer should be directed to the Head of the Laboratory Administration's Billing Unit by telephoning 443-681-3810.

# B. SPECIMEN SUPPLIES, PACKAGING, TRANSPORT, AND DELIVERY

#### **B.1. PACKAGING FOR TRANSPORT:**

Care must be taken to ensure a proper transport environment for specimens. Collect recommended quantities of test specimen and follow all directions for recording date and, where appropriate, time of specimen collection. Also make every effort to see that specimens are transported at required temperatures and in appropriate collection containers. Collection containers and other specimen supplies are available from the Laboratory's Supply Unit (443-681-3777). In addition, always separate glass tubes by using either protective material or separate biohazard bags to prevent breakage and cross-contamination during transport (see Basic Triple Packaging on page 10). A submitter using a courier service should take similar precautions by submitting individual tubes and requisition slips in separate, sealable plastic biohazard bags protected in an appropriate shipping container.

# TEST COLLECTION COMPONENTS AND OTHER LABORATORY SUPPLIES:

The Laboratories Administration provides test request forms and specimen collection components (e.g., tubes, bags, etc.). Requests and questions about supplies should be directed to the nearest Regional Laboratory or the Central Laboratory Supplies Unit at 443-681-3777 or fax the "Requisition for Specimen-Mailing Assemblies" form to 443-681-3850. To obtain the "Requisition for Specimen-Mailing Assemblies" visit our website at <a href="http://dhmh.maryland.gov/laboratories/docs/Requisition%20for%20Specimen-Mailing%20Assemblies.pdf">http://dhmh.maryland.gov/laboratories/docs/Requisition%20for%20Specimen-Mailing%20Assemblies.pdf</a>.

Note that various tests and specimens require different types of collection devices, transport media, and transport containers. Using the incorrect kit, collection component, or container will often render a test specimen unacceptable for analysis. If you have a question regarding the acceptable collection container contact the testing laboratory.

# **B.1.a. VIA STATE CONTRACTED COURIER**

Counties using the state contracted courier service must pack specimens and/or samples according to the temperature storage requirements. Specimens requiring freezing should be frozen and packed with adequate cooling (dry ice) material to maintain their proper temperature for up to 36 hours. Coolers are required to transport all specimens and/or samples through the state contracted courier. Therefore, it is essential that all coolers be properly labeled. Each cooler should specify the conditions for storage on all visible outer surfaces – "ROOM TEMPERATURE", "REFRIGERATE", or "FREEZE". Each cooler for specific laboratories should be labeled on all visible outer surfaces for "ENVIRONMENTAL" or "RABIES". Both Environmental and Rabies coolers must only be used as labeled. DO NOT use or re-use Environmental or Rabies cooler for any other types of specimens/samples, or add any other types of specimens to these coolers. A "RABIES" cooler must only be used for rabies samples, and an "ENVIRONMENTAL" cooler must only be used for environmental or Rabies cooler that are received in an Environmental or Rabies cooler that are not intended for Environmental or Rabies testing will be rejected and discarded for safety reasons. (Please see Rabies Section on page 15 for detailed information on animal rabies submissions).

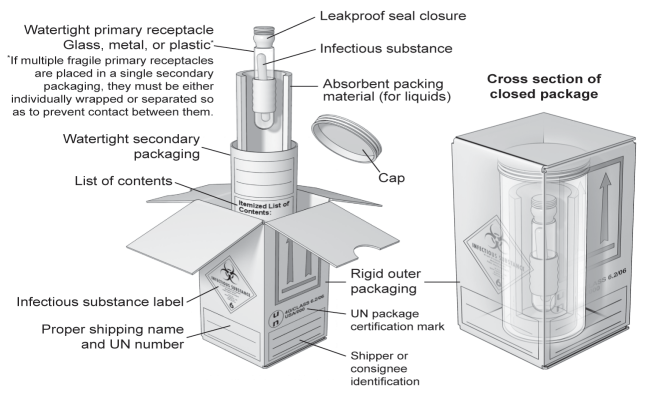
# B. 1. b. VIA U.S. MAIL OR OTHER CARRIER:

Due to regulations published by IATA (International Air Transportation Association), US DOT (United States Department of Transportation), and the USPS (United States Postal Service), the Laboratories Administration's specimen collection components may be used only when sending specimens via private or state-contracted courier. These containers are not approved or certified for use in the USPS system or other common carriers (e.g., FedEx, UPS, etc.). Infectious substances sent through the mail or by other common carriers must be packaged by individuals trained and certified in Infectious Substances shipping. **Certified packaging systems are not supplied by the Laboratories Administration.** 

Before using the USPS or other carrier, the shipper must refer to the current IATA, USPS and DOT regulations. IATA has divided infectious substances into two categories. IATA "Category A Infectious Substance" includes substances that are "transported in a form that, when exposure to it occurs, is capable of causing permanent disability, life-threatening or fatal disease in otherwise healthy humans or animals." "Category A Infectious Substances" are subject to the more stringent packing requirements described in IATA Packing Instruction 602. An IATA "Category B Infectious Substance" is defined as "an infectious substance that does not meet the criteria for inclusion in Category A." The proper shipping name of UN 3373 is Biological substance Category B. This includes human or animal material transported for research, diagnosis, disease treatment, etc., and diagnostic or clinical cultures. These specimens must be mailed and transported in packaging that meets IATA Packing Instruction 650.

# BASIC TRIPLE PACKAGING (Refer to tests for specific details)

Basic triple packaging systems include a primary receptacle such as a tube with adhesive tape around the screw cap or a plate with parafilm around the edges. The primary (1°) receptacle, along with required absorbent and cushioning material, is placed inside a secondary (2°) container. The 2° container for diagnostic specimens should be a sealed biohazard or Ziploc bag. The 2° container is then securely placed within an outer shipping container (tertiary (3°) container), generally a corrugated cardboard box with cushioning material inside to surround the 2° container. This outermost container bears the name, address, and telephone number of shipper, name of person responsible with 24/7 telephone number, and the complete name, shipping address, and telephone number of the recipient, plus all the required markings. Include an itemized list of contents in a sealed plastic bag, placed between the 3° and 2° containers. Specific instructions for various tests can be found in the test list section of this guide.



Example of a correctly prepared and labeled triple package for Biological specimen, Category B (UN 3373) (previously known as Clinical specimen and Diagnostic Specimen. A Category B infectious substance is one that does not meet the criteria for inclusion in Category A. A Category B infectious substance does not cause permanent disability or life-threatening or fatal disease to humans or animals when exposure to it occurs. The proper shipping name for a Category B infectious substance, "Biological specimen, Category B," is assigned to identification number "UN 3373." The proper shipping names "Diagnostic specimen" and "Clinical specimen" may no longer be used (as of January 1, 2007). (Modified from Biosafety in Microbiological and Biomedical Laboratories [BMBL], 5th edition) **BASIC TRIPLE PACKAGING:** 

1) A watertight primary receptacle.

2) A watertight secondary receptacle.

3) An outer packaging of adequate strength for its capacity, mass and intended use.

Note: For a liquid specimen, absorbent material must be placed between the primary and secondary containers and be capable of absorbing the entire contents of the primary receptacle(s).

Certified packaging systems are designed to withstand specific pressure changes and drop tests. Packaging systems that meet the packing instruction standards are currently available from vendors specializing in products certified to meet the IATA, USPS, and other carriers' requirements. Packaging systems using fiberboard or aluminum canisters, zip-lock bags, or other uncertified components may not be in compliance.

# IT IS THE RESPONSIBILITY OF THE SHIPPER TO COMPLY WITH ALL LAWS AND REGULATIONS REGARDING THE SHIPPING OF INFECTIOUS SUBSTANCES.

Questions may be referred to the DHMH Laboratories Administration's Quality Assurance Officer, Heather Peters, by calling 443-681-3791 or by email <u>heather.peters@maryland.gov</u>.

Resources: http://www.cdc.gov/biosafety/publications/bmbl5/ http://www.usps.com/

# **B.2. DELIVERY/DROP-OFF TO CENTRAL LABORATORY**

Specimens intended for the Central Laboratory should be directed to 1770 Ashland Avenue Baltimore, MD 21205. The Laboratory facility is located at the corner of Ashland and Rutland Avenues. All specimen and sample deliveries to the laboratory must be delivered to the loading dock located on Rutland Avenue. Temporary parking is available at the loading dock. Couriers delivering specimens are required to sign a loading dock security log sheet upon arrival.

# **B.2.a. Specimen/Sample Deliveries Accepted**

#### Clinical

Monday-Friday 8:00am-6:00pm Saturday: 7:30am-10:30am

# Newborn Screening

Monday-Friday 8:00am-6:00pm Saturday: 7:30am-2:00pm

Rabies specimens and testing: Contact Rabies on-call staff (see page 15).

# B.2.b. HOLIDAYS

A detailed holiday schedule can be found on the Laboratories Administration website at <a href="http://dhmh.maryland.gov/laboratories/">http://dhmh.maryland.gov/laboratories/</a>

# B.2.c. OTHER EMERGENCY REQUESTS INVOLVING DROP OFF OR LABORATORY SERVICES

Emergency on-call numbers:

(1.) Biological, chemical or radiological terrorism:	
Jim Svrjcek	410-925-3121 (cell)
Robert Myers	443-928-0925 (cell)
(2.) Microbiology emergency:	
Prince Kassim	202-271-4447 (cell)
Robert Myers	443-928-0925 (cell)
(3.) Environmental Chemistry emergency:	
Prince Kassim	202-271-4447 (cell)
Robert Myers	443-928-0925 (cell)

For unknown powders and environmental samples for bioterrorism/chemical terrorism see the Laboratories Administration website at http://dhmh.maryland.gov/laboratories/ or call a phone number under B.2.c.(1.), above.

#### C. SPECIMEN COLLECTION, PREPARATION, AND HANDLING

# C.1. GENERAL

Specimen quality is a product of the nature of the specimen itself, how well it was collected, and the manner in which it is or was transported to the laboratory. A laboratory can provide accurate and clinically relevant test results only if it receives good test specimens. Before attempting to collect a specimen, look up the desired test(s) in this reference guide. Check to see if there are specific requirements for:

- 1. Specimen type or volume;
- 2. Collecting procedures;
- 3. Collecting devices or containers.

Use the correct test request form and properly and legibly complete this form to ensure accurate and efficient laboratory service. Use a soft pencil or black ballpoint to print the information. Be sure to include proper identifying information on the test request form and the specimen itself.

Please note the clinic's full mailing address, test request authorized by personnel, and telephone number to assure proper return of test results. Then see that the test request form accompanies the specimen. The following sections provide practical guidelines to physicians, nurses, and other non-laboratory health personnel who must routinely collect and submit clinical specimens to one of the State's public health laboratories (i.e., Department of Health and Mental Hygiene's Laboratories Administration).

#### **C.1.a. PATIENT PREPARATION**

Prior to the time scheduled to collect a patient's specimen the patient should receive appropriate instructions concerning fasting, diet, and medication restriction. For example, a patient about to submit a specimen for a microbiology culture should have specimen(s) collected before starting antimicrobial therapy.

# C.1.b. SPECIMEN HANDLING BY SUBMITTER

The most common specimen handling errors include failing to:

- 1. Tighten specimen container lids or caps;
- 2. Label a specimen correctly; and
- 3. Provide all pertinent clinical information.

Properly identifying specimens is extremely important. Legibly label each specimen container or tube with the patient's full name, and date of specimen collection, just as they appear on the test request form. Information on specimens should be checked against information on the test request form for agreement before the specimen is sent to the laboratory.

#### C.2. PROCUREMENT AND SUBMISSION REQUIREMENTS, PRECAUTIONS, AND PROBLEMS BY SPECIMEN TYPE

#### C.2.a. BLOOD/SERUM

#### C.2.a.(1.) HEMOLYSIS

In general, grossly or even moderately hemolyzed blood specimens may not be acceptable for testing. Hemolyzed serum is pink or red, rather than the normal clear straw color. Most cases of hemolysis can be avoided by observing the steps below.

1. Use a needle no smaller than 20- or 21- gauge. (On occasion, however, it may be necessary to use a 22- or 23- gauge needle for patients from elderly and pediatric populations with small or difficult veins.) Hemolysis can be avoided by not placing small gauge Butterfly needles into Vacutainer tubes. Carefully and safely remove Butterfly and replace with a 16-gauge needle before penetrating Vacutainer tube.

2. If there is air leakage around the needle or loss of vacuum in the tube, replace the vacuum tube.

3. Collect blood in room temperature containers unless the specimen requirement specifies otherwise.

4. When a vacuum tube fills too slowly due to an incomplete venipuncture, damage to the red blood cells may result. Correct by deeper vein entry or select another puncture site and collect a second specimen.

5. Do not remove the needle from the vein until the vacuum tube is completely filled or the tube is pulled back from holder to release pressure. Premature removal causes a rush of air to enter the tube, with resultant damage to the red cells.

#### C.2.a.(2.) PAIRED SERA/PARALLEL TESTING

Both acute and convalescent sera are required to determine recent infection. Acute sera may be tested immediately and then stored until the convalescent sera are submitted. When both sera are available parallel testing under identical testing conditions will be performed to ensure an accurate comparison of acute and convalescent antibody titers. See Submission of Specimen for requested serological test.

#### C.2.a.(3.) VACUUM TUBES CONTAINING ANTICOAGULANTS

When using vacuum tubes containing anticoagulants and preservatives:

1. Tap the tube gently at a point just below the stopper to release any additive adhering to the tube or stopper.

2. Permit the tube to fill completely to ensure the proper ratio of blood to additive.

3. To ensure adequate mixing of blood with the anticoagulant or preservative, use a slow rolling wrist motion to invert the tube gently five or six times. Rapid wrist motion or vigorous shaking contributes either to small clot formation or hemolysis and fails to initiate proper mixing action.

4. Check to see that all the preservative or anticoagulant is dissolved. If any preservative powder is visible, continue inverting the tube slowly until the powder is dissolved.

#### C.2.a.(4.) VACUUM TUBES WITHOUT ANTICOAGULANTS

When using vacuum tubes containing no anticoagulants or preservatives, or SST serum Separator Tubes:

1. Permit the tube to fill completely.

2. Let the specimen stand for a minimum of 30 minutes and not longer than 45 minutes prior to centrifugation. This allows time for the clot to form. If the specimen is allowed to stand longer than 45 minutes, chemical activity and degeneration of the cells within the tube will take place, and test results will be altered as a consequence.

3. Centrifuge the specimen at the end of the 30 to 45 minute period in strict accordance with manufacturer's instructions for speed and duration of centrifugation.

#### C.2.a.(5.) QUANTITY NOT SUFFICIENT (QNS)

One of the most common errors in specimen collection is the submission of an insufficient quantity of specimen for testing. To ensure an adequate amount of specimen:

1. Always draw whole blood in an amount 2 ½ times the required volume of serum needed for a particular test. For example, if 4mL serum are required, draw at least 10mL whole blood.

2. For most profile testing submit one full tube of serum (8-10mL).

#### C.2.b. ENTOMOLOGICAL SPECIMENS

Identification of insects of medical importance (e.g., ticks) can be provided as a referral service. Please call the Microbiology Division (443-681-3943/443-681-3952) prior to submitting insect specimens.

C.2.c. RABIES SPECIMENS

# C.2.c.(1.) HOURS OF OPERATION

The DHMH Laboratories Administration Rabies Laboratory operates from 8:00 AM to 4:30 PM weekdays (Monday through Friday except on holidays. On-call laboratory scientists are available for requests that require test results as soon as possible so that a medical determination on rabies post-exposure prophylaxis (PEP) can be made.

Specimens must be received at the DHMH Laboratories Administration by 12:00 PM on Fridays to have the test results reported by Friday 4:30 PM. Specimens received on Fridays after 12:00 PM will have the results ready the next regular workday.

Specimens received on evenings from Monday through Friday, Fridays from 12:00 PM to 4:30 PM, on a weekend, or on a State holiday will be processed on the next regular workday, except for situations that require test results as soon as possible so that a medical determination about rabies PEP can be made (emergency examination). In these situations, prior approval by epidemiology staff in the DHMH Office of Infectious Disease Epidemiology and Outbreak Response (IDEOR) is necessary before testing will be initiated by on-call laboratory scientists. (For details, please see the Emergency Examination Requests section below).

# C.2.c.(2.) DELIVERY PROCEDURES

Delivery of specimens must be from Monday through Friday 7:30AM to 6:00PM (regular workdays) to the DHMH Laboratories Administration Loading Dock at 1770 Ashland Ave Baltimore, Maryland 21205. All animal submission of specimen must be routed through the local health department and sent via courier service. <u>Do not use</u> the U.S. Postal Service or other public transportation service to send specimens. (For emergency examination situations, please see the Emergency Examination Requests section below).

# C.2.c.(3.) ORDERING TESTS

For routine testing Monday through Friday, all local health departments must use the DHMH Laboratories Administration's MyLIMS (Laboratory Information Management Systems)

(http://starlims.dhmh.md.gov/starlims10.dhmhlabs.prod/) for submission of specimens. The updated rabies submission form (DHMH 1188 11/10) will print out automatically when the animal rabies test is ordered through the MyLIMS system, and must be included in the cooler and attached to the specimen being submitted. One Rabies Submission Form should accompany each animal submitted. Specimens approved for emergency testing must be accompanied by a fully completed handwritten Rabies Submission Form if access to MyLIMS is not possible. An emergency contact name and phone number must be listed on the Rabies Submission Form. The updated rabies submission form (DHMH 1188 11/10) can be downloaded from our website at http://dhmh.maryland.gov/laboratories/Pages/Rabies.aspx.

# C.2.c.(4.) CRITERIA FOR ANIMAL SUBMISSION

Live animals will **NOT** be accepted in the laboratory. Terrestrial animals acceptable for submission to DHMH are rabies vector species (e.g., raccoons, foxes, skunks, etc.) that expose humans, livestock, or pets. Exposure is defined as a bite that breaks the skin or contact of mucous membranes or broken skin with either animal saliva or nervous tissue. Birds, fish, reptiles and amphibians will not be accepted for rabies testing under any circumstances. Small rodents, including squirrels, chipmunks, gerbils, guinea pigs, hamsters, rabbits, mice, rats, voles, shrews and moles, will not be accepted for testing unless (1) the animal has bitten a human and (2) prior approval for testing has been authorized by the DHMH IDEOR epidemiology staff. Most recent human cases of rabies in the U.S. have been associated with bats, and bat bites may be difficult to recognize.Bats should be submitted for testing in all cases of direct human contact with a bat or when bite or mucous membrane contact cannot be ruled out. Live animals will **NOT** be accepted in the laboratory.

**Please Note:** Large animal heads (e.g. horse and cow) should be submitted to the Maryland Department of Agriculture for brain tissue extraction.

# C.2.c.(5.) EMERGENCY EXAMINATION REQUESTS

Some situations that occur after regular business hours may require rabies test results as soon as possible so that a medical determination about rabies PEP can be made. In these Situations, on-call laboratory scientists are available; and specimens may be examined Fridays from 12:00 PM to 4:30 PM, on a weekend, or on a State holiday, with prior approval of DHMH PHPA (Prevention and Health Promotion Administration) epidemiology staff. To reach the epidemiology staff during regular business hours, contact the DHMH PHPA for Zoonotic and Vector-borne Diseases (CZVBD) at 410-767-5649 (main); 410-767-6703 (DHMH State Public Health Veterinarian ); or 410-767-6618 (CZVBD) Rabies Chief). After hours, use the DHMH IDEORB (Infectious Disease Epidemiology and Outbreak Response Bureau) Epidemiologist-On-Call pager at 410-716-8194 or call the SYSCOM operator at 410-795-7365 and ask to be directed to the Epidemiologist-on-Call for all rabies consultations.

After receiving approval for an emergency examination request, contact one of the following DHMH Laboratories Administration staff **(in the order listed below)** to arrange for testing and appropriate submission. (NOTE In addition to the rabies submission form, the specimen should be accompanied by the submitter's after-hours contact information to receive results).

- 1) Rabies Lab On-Call No: 443-735-1291
- 2) Rabies Lab Supervisor (Kenneth Okogi): 443-799-9490
- 3) OLEPR (Jim Svrjcek or BT Coordinator): 410-925-3121
- 5) Laboratory Director, Dr. Robert Myers: 443-928-0925

# C.2.c.(6.) SPECIMEN COLLECTION

Live animals will **NOT** be accepted in the laboratory. Animals should be euthanized in a manner that will not destroy the brain tissues to be examined in the diagnosis of rabies. When possible, only the animal's head should be submitted for diagnostic purposes. For animals weighing more than 20 pounds, particularly large dogs, only the head may be submitted for testing. If an animal is being submitted to DHMH Labs from an animal pathology or diagnostic laboratory, and the animal has already been prepared for necropsy, the submitter should submit all or a cross section of the brainstem and half of the cerebrum.

**Please Note:** Large animal heads (e.g. horse and cow) should be submitted to the Maryland Department of Agriculture for brain tissue extraction.

# C.2.c.(7.) PACKAGING AND SHIPPING

•All rabies specimens must be placed into coolers that are clearly marked as rabies coolers. No other non-rabies clinical samples may be placed into rabies coolers or these samples will be rejected.

- •Rabies coolers must fully close and must be waterproof.
- •Each specimen must be individually packaged in a leak-proof bag and clearly labeled.
- •Each specimen must be accompanied by a Rabies Submission Form for proper identification.
- •All Rabies Submission Forms must be filled out correctly and legibly including exposure type.
- •Coolers may be shipped with ice or ice packs but the ice should not occupy more than 1/3 of the cooler.
- Live animals will **NOT** be accepted in the laboratory.
- •Submitters should avoid freezing specimens. If frozen specimens are received, testing will be delayed.
- Trash **MUST** not be sent in rabies coolers.

•Animal rabies packaging and training video available at <a href="http://dhmh.maryland.gov/laboratories/Pages/Rabies-Animal-DFA.aspx">http://dhmh.maryland.gov/laboratories/Pages/Rabies-Animal-DFA.aspx</a>

#### D. GUIDE TO PUBLIC HEALTH LABORATORY TESTS:

TEST:	<b>ABCs (previously BIDS)</b> includes <i>Neisseria meningitidis, Haemophilus influenzae</i> , Group A streptococcus, Group B Streptococcus, and <i>Streptococcus pneumoniae</i> . <i>Listeria</i>
	<i>monocytogenes</i> is handled as an ABCs isolate and evaluated by the National Antimicrobial Resistance Monitoring Systems (NARMS) Program.
Synonym:	Active Bacterial Core Surveillance (Bacterial Invasive Disease Surveillance)
Laboratory/Phone:	Microbiology / 443-681-3952
Turnaround Time:	N/A
Specimen Required:	Pure culture on agar slant in screw cap tube.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Bacterial isolate
Specimen Volume (Minimum):	N/A
Collect:	N/A
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate ABCs # and organism identification on test request form. Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	AT ROOM TEMPERATURE - DO NOT REFRIGERATE ISOLATE - DO NOT FREEZE.
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> <li>Specimen frozen</li> </ul>
Results and Interpretation:	N/A
Reference Range:	
Additional Information:	SUBCULTURE TO AGAR SLANT BEFORE TRANSPORTING. DO NOT SEND CULTURE PLATES.
Purpose of Test:	Active Bacterial Core Surveillance (ABCs) is a core component of the CDC's Emerging Infections Programs Network (EIP).
Method:	Isolate is subcultured and identified prior to submission to CDC.
Interfering Substances/Limitations:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
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Comment:	Active Bacterial Core Surveillance (ABCs) is a core component of the CDC's Emerging
	Infections Programs Network (EIP), collaboration between CDC, state health departments,
	and universities. ABCs is an active laboratory and population-based surveillance system
	for invasive bacterial pathogens of public health importance. For each case of invasive
	disease in the surveillance population, a case report with basic demographic information
	is completed and bacterial isolates are sent to CDC and other reference laboratories for
	additional laboratory evaluation.
	ABCs was initially established in four (4) states in 1995. It currently operates among ten
	(10) EIP sites across the United States, representing a population of over 38 million
	persons. At this time, ABCs conducts surveillance for six (6) pathogens: Group A and
	Group B streptococcus (GAS, GBS), Haemophilus influenzae, Neisseria meningitidis,
	Streptococcus pneumoniae, and Listeria monocytogenes. The Maryland Department of
	Health and Mental Hygiene is an EIP site with partner Johns Hopkins Bloomberg School of
	Public Health.

TEST:	Adenovirus, Viral Culture
Synonym:	Adenovirus: Virus Culture, Virus isolation: Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934
Turnaround Time:	3-28 days

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TEST:	AFB/Acid-fast Bacilli culture (Mycobacterium tuberculosis identification)
Synonym:	AFB/Acid Fast Bacteria Identification (Acid Fast Bacilli); M. Tuberculosis culture: Refer to
	instructions for Mycobacterium tuberculosis culture.
Laboratory/Phone:	Mycobacteriology / 443-681-3942

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TEST:	Amoebiasis (Ova and Parasites Microscopic Examination)
Synonym:	Amoebiasis; Amebiasis: Refer to instructions for Ova and Parasites Microscopic
	Examination.
Laboratory/Phone:	Microbiology / 443-681-3952

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TEST:	Anthrax, Cutaneous
Synonym:	Bacillus anthracis, Woolsorters' disease
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952
Turnaround Time:	2-7 days [from specimen receipt in the Laboratory]
Specimen Required:	1. Vesicular Stage: Vesicular fluid
	2. Eschar Stage: Eschar material
	3. Isolate
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	1. Vesicular Stage: Collect vesicular fluid on sterile swab from previously unopened
	vesicles.
	2. Eschar Stage: Collect eschar material by carefully lifting the eschar's outer edge,
	insert sterile swab, then slowly rotate for 2-3 seconds beneath the edge of the
	eschar without removing it.
	3. Isolate: Pure culture, 24 hours old, growing on sheep blood agar plate.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on the form.
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Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	<ol> <li>Swabs: Transport directly to laboratory at room temperature. For transport time &gt; 1 hour, transport at 2-8°C.</li> <li>Isolate: Transport the specimen at room temperature on a sealed sheep blood agar</li> </ol>
Specimen Rejection Criteria:	plate.The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.Unlabeled or improperly labeled specimenNon-sterile or leaking containerInappropriate specimen transport conditionsIllegible, or no submitter information on the request formBroken specimen/sample containerThe wrong specimen for test requestInappropriate outfit for requested testIllegible or no patient information on the specimen
	Expired transport media
Availability: Results and Interpretation:	24 hours/day, 7 days/week Bacillus anthracis isolated/detected. Bacillus anthracis not found.
Additional Information:	Call 410-925-3121 before sending specimen to the Laboratory.
Purpose of Test:	To confirm diagnosis of cutaneous anthrax.
Method:	LRN Methods
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland
Comment:	Call 410-925-3121 before sending to the Laboratory.

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TEST:	Anthrax, Gastrointestinal
Synonym:	Bacillus anthracis, Woolsorters' disease
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952
Turnaround Time:	2-7 days [from specimen receipt in the Laboratory]
Specimen Required:	1. Blood Cultures
	2. Stool
	3. Rectal swab (for patients unable to pass a specimen)
	4. Isolate
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must be
	properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	<ol> <li>Blood Cultures: Collect appropriate blood volume and number of sets per routine laboratory protocol.</li> </ol>
	<ol> <li>Stool: Transfer ≥ 5g of stool directly into a clean, dry, sterile, wide-mouth, leak-proof container.</li> </ol>
	3. Rectal swab: Insert a sterile swab one (1) inch beyond the anal sphincter.
	4. Isolate: Pure culture, 24 hours old, growing on a sheep blood agar plate.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on the form.
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	The 5. Wenser Joseph Tuble Tealth Laboratory
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	1. Blood Cultures: Transport directly to the laboratory at room temperature.
	2. Stool: Transport unpreserved stool to laboratory within one (1) hour. For transport
	time > 1 hour, transport at 2-8°C. Cary-Blair or equivalent transport media is
	acceptable.
	3. Rectal Swab: Transport swab(s) directly to laboratory at room temperature. For
	transport time $> 1$ hour, transport at 2-8°C.
	4. Isolate: Transport the specimen at room temperature on a sealed sheep blood plate.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results
	and to avoid misleading information that might lead to misdiagnosis and inappropriate
	therapy. A request for a new specimen will provide appropriate materials and clinically
	relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	Expired transport media
Availability:	24 hours/day, 7 days/week
Results and Interpretation:	Bacillus anthracis is isolated/detected.
	Bacillus anthracis not found.
Additional Information:	Call 410-925-3121 before sending specimen to the Laboratory.
Purpose of Test:	To confirm diagnosis of gastrointestinal anthrax.
Method:	LRN Methods
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Call 410-925-3121 before sending specimen to the Laboratory.

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TEST:	Anthrax, Inhalational
Synonym:	Bacillus anthracis, Woolsorters' disease
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response: <b>410-925-3121 (24/7 emergency contact number)</b> Select Agents Microbiology Laboratory: 443-681-3954 Division of Microbiology Laboratory: 443-681-3952
Turnaround Time:	2-7 days [from specimen receipt in the Laboratory]
Specimen Required:	Blood Cultures     Sputum     Isolate
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	<ol> <li>Blood Cultures: Collect appropriate blood volume and number of sets per routine laboratory protocol.</li> <li>Sputum: Collect &gt;1 ml of a lower respiratory specimen into a sterile container.</li> <li>Isolate: Pure culture, 24 hours old, growing on a sheep blood agar plate.</li> </ol>
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on the form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
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Transport Conditions:	<ol> <li>Blood Cultures: Transport directly to the laboratory at room temperature.</li> <li>Sputum: Transport in sterile, screw-capped container at room temperature when transport time is &lt;1 hour. For transport time &gt; 1 hour, transport at 2-8°C.</li> <li>Isolates: Transport at room temperature on a sealed sheep blood agar plate.</li> <li>The following rejection criteria are designed to prevent the reporting of inaccurate</li> </ol>
Specimen Rejection Criteria:	<ul> <li>results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> </ul>
Availability:	24 hours/day, 7 days/week
Results and Interpretation:	Bacillus anthracis isolated/detected; Bacillus anthracis not found.
Additional Information:	Call 410-925-3121 before sending specimen to the Laboratory.
Purpose of Test:	To confirm diagnosis of Inhalational Anthrax.
Method:	LRN Methods
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Call 410-925-3121 before sending to the Laboratory.

TEST:	Antimicrobial Susceptibility Test
Synonym:	Disk Diffusion Susceptibility Testing, E-test, Susceptibility Testing or Microbroth Dilution
	Susceptibiliy Testing
Laboratory/Phone:	Microbiology / 443-681-3952
Turnaround Time:	48-72 hrs [from specimen receipt in the Laboratory]
Specimen Required:	Original specimen or pure isolate of rapidly growing non-fastidious aerobic bacteria.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Viable pure isolate on an appropriate slant.
Specimen Volume (Minimum):	N/A
Collect:	N/A
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate
	results and to avoid misleading information that might lead to misdiagnosis and
	inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
	Non-viable organism
Availability:	Monday through Friday

Results are reported as S-I-R, following Clinical Laboratory Standards Institute (CLSI) criteria for organism/source combination.
CSLI guidelines
If original specimen is submitted, pathogenic bacteria should be isolated from it.
To assist the physician in choosing an appropriate antimicrobial agent(s) for therapy.
Disk Diffusion
Administration of antimicrobial agents before specimen collection.
DHMH Laboratories Administration, Central Laboratory
1770 Ashland Avenue, Baltimore, Maryland 21205
The antibiotics tested and reported will follow the latest CLSI recommendations appropriate for the bacterial species submitted for testing; the methodology used will also follow CLSI recommendations.

TEST:	Antimicrobial Susceptibility Test, Minimum Inhibitory Concentration (MIC),
	Aerobic Bacteria
Synonym:	N/A
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	48-72 hrs [from specimen receipt in the Laboratory]
Specimen Required:	Original specimen or a pure isolate of aerobic bacteria.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
specifici lucitimation.	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Viable pure isolate on an appropriate slant.
Specimen Volume (Minimum):	N/A
Collect:	N/A
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
Tomi.	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
r dekaging and Snipping .	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate
	results and to avoid misleading information that might lead to misdiagnosis and
	inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Results are reported as S-I-R following Clinical Laboratory Standard Institute (CLSI)
Results and interpretation.	criteria for organism/source combination.
Reference Range:	CSLI guidelines
Additional Information:	Test is performed on aerobic possible pathogens.
Purpose of Test:	To assist the physician in choosing an appropriate drug therapy, monitoring emerging
Purpose of Test:	resistance, monitoring percentage susceptibility trend.
Method:	E-Test, Microbroth Dilution, or Vitek
Interfering Substances:	Administration of antimicrobial before specimen collection.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
Commont	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Arbovirus Culture
Synonym:	
Laboratory/Phone:	Virology: 443-681-3937
Turnaround Time:	3-6 weeks for both negatives and positives
Specimen Required:	CSF, throat washing, brain and spinal cord tissue
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time
	of collection. The specimen/sample must be properly labeled and match the test
	requisition or electronic test order.
Specimen Volume (Optimum):	≥ 2ml or 4 grams of tissue
Specimen Volume (Minimum):	2ml or 4 grams of tissue
Collect:	Sterile container with leak-proof lid.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Deliver on dry ice.
Specimen Rejection Criteria:	Unlabeled specimen, mismatch between labeling of specimen and test request form.
Availability:	Monday-Friday
Results and Interpretation:	Isolated or No viruses isolated
Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name given to
	viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).
	Arboviruses that cause human encephalitis are members of three virus families: The
	Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae.
	For more information, see the CDC link at:
	https://www.cdc.gov/ncezid/dvbd/
Purpose of Test:	Virus isolation to determine probable cause of infection and aid in the diagnosis of viral
	disease or to further characterization for epidemiological purposes.
Method:	Viral culture
Interfering Substances:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	

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TEST:	Arbovirus Endemic Panel
	Panel includes WNV and SLE IgM MIA and EEEV IgM EIA. LaCrosse (LAC) IgM
	testing available based on patient's travel history.
Synonym:	Arthropod-borne virus: WNV (West Nile Virus), EEEV (Eastern Equine Encephalitis Virus),
	SLEV (St. Louis Encephalitis Virus)
Laboratory/Phone:	Virology: 443-681-3937
Turnaround Time:	5-10 working days during Arbovirus Season (excluding PRNT Testing)
Specimen Required:	Serum (blood); CSF
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time
	of collection. The specimen/sample must be properly labeled and match the test
	requisition or electronic test order.
Specimen Volume (Optimum):	2 ml serum; 2ml CSF
Specimen Volume (Minimum):	1 ml serum; 0.5 ml CSF
Collect:	Red top vacuum tube, transfer serum to sterile tube: CSF in sterile container with leak- proof cap.
Request Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	For testing to be initiated, the following information MUST be provided: date of onset,
	and date specimen collected. Also please provide: patient's date of birth, diagnosis,
	symptoms, fatality, travel history, immunizations, and immunocompromised.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
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Transport Conditions:	Store refrigerated and chin on cold packs in a cooler. If chinning is delayed hereard 49
Transport Conditions:	Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48 hours CSE must be frequent at 20°C and chipped on dry ice
Constitution Delivation Criteria	hours, CSF must be frozen at -20°C and shipped on dry ice.
Specimen Rejection Criteria:	Grossly hemolyzed specimens, unlabeled specimen, leaking container, duplicate
	specimen type (ex: two CSF specimens collected the same day- one tube will not be
	tested), and mismatch between labeling of specimen and test request form.
Availability:	Monday through Friday.
Results and Interpretation:	(EIA) IgM: Negative, High Background, Equivocal, Positive
	(MIA) IgM: Positive, Negative, Nonspecific
	Serum and CSF that tests positive for IgM is consistent with acute infection. A positive
	IgG antibody and a negative IgM antibody are consistent with infection in the distant past
	and are not consistent with acute infection.
Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name give to
	viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).
	Arboviruses that cause human encephalitis are members of three virus families: The
	Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae. For more information,
	see the CDC link at: http://www.cdc.gov/ncidod/dvbid/arbor/arbdet.htm
	The MD State DHMH laboratory routinely tests for IgM ab to WNV, SLEV and EEEV, which
	are endemic to this area. Confirmatory testing by PRNT (plaque reduction neutralization
	test) may be necessary on positive samples. A convalescent serum sample (collected > 10
	days after onset date) is needed for PRNT testing. Please contact the Arbovirus
	laboratory with any questions regarding PRNT. Patients with travel history supporting
	suspicion of other arboviruses will be forwarded to the CDC for testing. LAC IgM serology
	testing is available based on patient's travel history.
Purpose of Test:	For the presumptive detection of IgM antibody to WNV, SLEV, EEEV, and LAC.
	Confirmatory testing by PRNT may be required.
Method:	EIA, MIA (Microimmunoassay), PCR, PRNT
Interfering Substances:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	If eligible for PCR (all CSF specimens and acute serum samples (<10 days between onset
comment.	date and collection date.) If the laboratory receives both serum and CSF, only the CSF will
	be tested by PCR.) PCR testing is performed first. If the volume is low, the PCR assay
	cannot be performed.
	Paired specimens are NOT required:
	IgM antibody for WNV & SLEV is performed on all specimens. If the sample volume
	permits, EEEV IgM testing will also be performed.

TEST:	Arthropod Identification
Synonym:	Tick identification/Ectoparasite
Laboratory/Phone:	Microbiology/ 443-681-3952
Turnaround Time:	48-72 hrs [from specimen receipt in the Laboratory]
Specimen Required:	Whole parasite
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Whole parasite
Specimen Volume (Minimum):	N/A
Collect:	Collect the whole parasite; put it in a clean container with a tight fitting lid with alcohol.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature
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Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> <li>Received only partial parasite</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Genus/species
Reference Range:	N/A
Additional Information:	N/A
Purpose of Test:	Identify disease carrying arthropods
Method:	Macroscopic examination
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Aspergillus serology
Synonym:	Aspergillosis antibody test
Laboratory/Phone:	Virology: 443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include the patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer tube
Form:	DHMH 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to Page 9 & 10). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis
Availability:	Monday through Friday
Results and Interpretation:	POSITIVE- Antibodies against (A. fumigatus, A. flavus, A. niger) detected. NEGATIVE- Antibodies against (A. fumigatus, A. flavus, A. niger) not detected.
Additional Information:	
Purpose of Test:	For the detection of antibody to A. fumigatus, A. flavus, A. niger
Method:	Micro-immunodiffusion
Interfering Substances:	Hemolysis
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

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TEST:	Babesia serology
Synonym:	Babesia microti, Babesiosis
Laboratory/Phone:	Virology: 443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen Identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer tube
Form:	DHMH 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	<ul> <li>≥1:64: Reflect infection at an undetermined time by <i>Babesia microti</i></li> <li>&lt;1:64: <i>Babesia</i> antibody not detected. Another specimen should be drawn if the original was taken soon after onset</li> </ul>
Additional Information:	http://www.cdc.gov/parasites/babesiosis/
Purpose of Test:	Detect IgG antibodies which may be due to a Babesia microti infection
Method:	IFA
Interfering Substances:	Cross reaction with <i>Plasmodium spp.</i> has been documented. Cross-reactivity with <i>Babesia divergens,</i> which causes a more severe infection in European patients is possible.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

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TEST:	Bacillus anthracis Culture
Synonym:	For Bacillus anthracis culturing: Refer to Anthrax, Cutaneous, Anthrax, Gastrointestinal,
	or Anthrax, Inhalational, for specific instructions as required.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Bacillus cereus Culture
Synonym:	Bacillus cereus Culture: For specific instructions refer to Foodborne Pathogens (Bacillus
	cereus, Clostridium perfringens, Staph aureus).
Laboratory/Phone:	Microbiology / 443-681-3952

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TEST:	Bacterial Culture, Routine
Synonym:	Aerobic culture, routine culture, eye culture, ear culture, genital culture, nose culture, respiratory culture, throat culture, urine culture, wound culture, sterile fluid culture.
Laboratory/Phone:	Microbiology / 443-681-3952
Turnaround Time:	Varies depending on culture site and organisms isolated, usually 2-4 days (or longer if fastidious organism isolate) [from specimen receipt in the Laboratory].
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HMH-Laboratories Administration	The J. Mehsen Joseph Public Health Laboratory
Specimen Required:	Swab from site in transport media (Amies, Stuarts, culturette) Aseptically aspirated pus or tissue Clean-catch urine
	Fluid in sterile container with leak-proof lid
	Do not send a syringe with needle attached. (Specimen will be rejected)
Specimen identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Swab or 0.5 ml fluid
Specimen Volume (Minimum):	N/A
Collect:	Most sites: Use swab to collect and place in transport media (Amies or Stuarts).
	Urine: fresh, clean-catch urine in screw cap jar, refrigerate, must reach lab within 24 hours, ship promptly on cold packs.
	Wound: Disinfect contiguous areas of skin or mucous membrane containing resident normal flora prior to culture collection. Collect exudates from the interior of productive lesions. Keep tissue samples moist.
	A thin, air-dried smear for Gram stain obtained from the same site as the culture is recommended.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature: abscesses, burn swabs, dental cultures, ear (inner ear), eye specimens, sterile body fluids, genital, Intra Uterine Device (IUD), spore testing, tissues, wound swabs, nasopharynx, upper respiratory cultures.
	At refrigerator (4°C) if kept > 2 hours: catheters, ear (external ear), feces for C. difficile Toxin A&B ( <b>frozen if test not done within three (3) days</b> ), sputum, urine – all types, autopsy tissue.
	At 37°C (or room temperature, if unavailable): blood culture bottles, bone marrow, cerebrospinal fluid (CSF)
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> <li>Specimen received after prolonged delay (usually more than 72 hours)</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Identification of potentially pathogenic organisms and antimicrobial susceptibilities, if clinically appropriate.
Reference Range:	No growth, routine/normal skin flora, routine/normal "body site" flora.
Additional Information: Purpose of Test:	N/A Isolation, identification and if clinically appropriate, antimicrobial susceptibilities of
Method:	potentially pathogenic organisms. Culture, staining, biochemical testing, antimicrobial susceptibility testing.
Interfering Substances/Limitations:	Only rapid-growing, no fastidious aerobic organisms can be recovered and identified by routine culture methods. "Bacterial culture, routine" will not detect anaerobic bacteria, chlamydia, viruses, fungi, or mycobacteria.

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Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Bacterial Referred Culture for ID
Synonym:	Isolate for Identification; referred culture
Laboratory/Phone:	Microbiology / 443-681-3952
Turnaround Time:	Varies depending on organisms submitted.
Specimen Required:	Isolate subcultured on agar slant with a leak-proof screw top lid.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	N/A
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store and ship at the proper temperature
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate
	results and to avoid misleading information that might lead to misdiagnosis and
	inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	Expired transport media
	Specimen received after prolonged delay (usually more than 72 hours)
Availability:	Monday through Friday
Results and Interpretation:	Identification of submitted isolate.
Reference Range:	N/A
Additional Information:	N/A
Purpose of Test:	Identification and if clinically appropriate, antimicrobial susceptibilities of potentially
	pathogenic organisms.
Method:	Culture, staining, biochemical testing, and MALDI-TOF.
Interfering Substances/Limitations:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Bang's Disease (Brucella serology and Brucella species culture)
Synonym:	Bang's Disease, Undulant fever, Malta Fever, and Rock of Gibraltar Fever: Refer to
	instructions for Brucella serology or Brucella species, culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Blood Culture (limited to Medical Examiner and special requests only)
Synonym:	N/A
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	Seven (7) days [from specimen receipt in the Laboratory]
Specimen Required:	Blood collected in B-D blood culture bottle
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	10 ml of right-heart blood
Specimen Volume (Minimum):	N/A
Collect:	Best collected before body is handled too much or opened. Decontaminate skin or seal
	surface of heart or other organ before inserting needle.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and
	inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	The wrong specimen for test request
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	If $\leq$ 3 organisms then Genus/species.
	If $\geq$ 3 organisms – no identification (hold organism for 10 days).
Reference Range:	No growth after seven (7) days incubation.
Additional Information:	N/A
Purpose of Test:	Assist Medical Examiner to establish the cause of death.
Method:	Culture, biochemical, and MALDI-TOF.
Interfering Substances:	Antibiotic therapy
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

TEST:	Bordetella Pertussis Culture
Synonym:	Pertussis, Whooping cough; B. pertussis culture, PCR
Laboratory/Phone:	Microbiology: 443-681-3952
Turnaround Time:	7-10 days [from receipt in the Laboratory], preliminary as soon as positive is detected.
Specimen Required:	Nasopharyngeal aspirates or nasopharyngeal swabs are both acceptable. Throat swabs are less suitable since <i>B. pertussis</i> exhibits tropism for ciliated respiratory epithelium, which is not found in the pharynx. However, throat swabs may be suitable for PCR diagnosis. Dacron <sup>TM</sup> swabs are to be used for both culture and PCR. Cotton-tipped swabs are to be avoided since they contain fatty acids that are toxic and may inhibit the growth of <i>B. pertussis</i> .
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
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Specimen Volume (Optimum):	Culture: Nasopharyngeal specimen on Dacron™ swab inserted in Regan-Lowe transport media.
	PCR: Nasopharyngeal specimen on Dacron <sup>™</sup> swab, submitted in Regan-Lowe transport media.
Specimen Volume (Minimum):	N/A
Collect:	Collect according to kit instructions. To order Pertussis culture kit, call 443-681-3777.
	Use Dacron™-tipped swabs only.
	1. Remove swabs from sterile package.
	2. Infants and young children should be supine. The infant/child's head must be held immobile by an assistant.
	3. Pass two (2) swabs simultaneously through one nostril and gently along the floor of
	the nasopharyngeal cavity until it reaches the posterior nares. <b>NOTE: Do not force</b> <b><u>swabs</u>.</b> Obstructions may be due to septal deviation.
	<ol> <li>Gently rotate both swabs together and leave in nasopharynx for 15 to 30 seconds to</li> </ol>
	absorb mucus.
	5. Repeat procedure through other nostril using the same two (2) swabs.
	6. Place each swab into a separate tube of transport media, run the swab (streak) up
	the agar and then put the swab into the media.
	7. Label both transport tubes with patient's name and place each tube back into the
	ziploc bag.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Best results are obtained by transporting specimen at room temperature the same day
	taken. If delays are expected (not transported the same day), place inoculated tubes
	into an incubator at 35-37°C. Cooled transport of the specimen significantly decreases
	the number of bacteria.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate
	results and to avoid misleading information that might lead to misdiagnosis and
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	and clinically relevant information to support good patient care.
	<ul><li>and clinically relevant information to support good patient care.</li><li>Unlabeled or improperly labeled specimen</li></ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> </ul>
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Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Bordetella pertussis PCR
	PCR cannot be ordered independently of culture (See Bordetella pertussis
	culture). Both assays are performed in parallel
Synonym:	B. pertussis, pertussis, Whooping Cough
Laboratory/Phone:	Molecular Biology: 443-681-3924
Turnaround Time:	2-3 Business Days
Specimen Required:	Nasopharyngeal specimen on Dacron swab, submitted in Regan-Lowe transport media.
Specimen Identification:	Specimen should be labeled with patient's name, and date of collection. The
specimentaentineation.	specimen/sample must be properly labeled and match the test requisition or electronic
	test order.
Specimen Volume (Optimum):	N/A Nasopharyngeal swab
Specimen Volume (Optimum):	N/A Nasopharyngeal swab
Collect:	To order Pertussis PCR/culture kit, call 443-681-3777.
	Collect according to kit instructions. Use Dacron <sup>™</sup> -tipped swabs only.
	1. Remove swabs from sterile package.
	<ol> <li>Infants and young children should be supine. The infant/child's head must be held</li> </ol>
	immobile by an assistant.
	3. Pass two (2) swabs simultaneously through one nostril and gently along the floor of
	the nasopharyngeal cavity until it reaches the posterior nares. <b>NOTE: Do not force</b>
	swabs. Obstructions may be due to septal deviation.
	4. Gently rotate both swabs together and leave in nasopharynx for 15 to 30 seconds to
	absorb mucus.
	5. Repeat procedure through other nostril using the same two (2) swabs.
	6. Place each swab into a separate tube of transport media, run the swab (streak) up
	the agar and then put the swab into the media.
	7. Label both transport tubes with patient's name and place each tube back into the
	ziplock bag.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
	Specimens must be packaged in a triple packaging system to ensure that under normal
Packaging and Shipping*:	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	Page 9 & 10).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Best results are obtained by transporting specimen at room temperature the same day
	taken. If delays are expected (not transported the same day), place inoculated tubes into
	an incubator at 35-37°C. Cooled transport of the specimen significantly decreases the
	number of bacteria.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate
	results and to avoid misleading information that might lead to misdiagnosis and
	inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> <li>Illegible or no submitter information on the request form</li> </ul>
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	<ul> <li>Regan-Lowe media not used</li> </ul>
	<ul> <li>Media expired</li> </ul>
	<ul> <li>Specimen frozen</li> </ul>
	<ul> <li>Specified frozen</li> <li>Unlabeled specimen or name discrepancy between specimen and request label</li> </ul>
	<ul> <li>Prolonged delay in transport (usually more than 72 hours)</li> </ul>
Availability:	Monday through Friday
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Results and Interpretation:	Positive: B. pertussis DNA WAS DETECTED by real time PCR
	Negative: B. pertussis DNA WAS NOT DETECTED by real time PCR
Additional Information:	PCR cannot be ordered independent of culture. Both assays are performed in
	parallel
Purpose of Test:	Detect the presence of B. pertussis nucleic acid (DNA).
Method:	PCR: Polymerase chain reaction, real-time
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

TEST:	Bordetella Pertussis Toxin IgG Antibody
Synonym:	IgG Anti-pertussis toxin assay
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first Name, DOB, specimen type, date and time of
	collection. The specimen/sample must be properly labeled and match the test requisition
	or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
Form:	For outbreak investigation use only. Prior approval required. Specific specimen criteria
	applies for details call 443-681-3889
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
·	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Packaging and Shipping:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Specimen Rejection Criteria:	Specimen from patient vaccinated against B.pertussis toxin in < 6months or age under 11
	cannot be tested. Discrepancy between name on tube and name on form, unlabeled;
	hemolytic; gross bacterial contamination. Refer to serology guideline
Availability:	Monday through Friday
Results and Interpretation:	Results can be used for investigational use only
	Pertussis antitoxin IgG level:
	Positive: ≥94 IU/ml
	Negative:<49 IU/ml
	Equivocal: between 49-93 IU/ml
Additional Information:	For more information, see the CDC link at: <a href="http://www.cdc.gov/hepatitis/index.htm">http://www.cdc.gov/hepatitis/index.htm</a>
Purpose of Test:	Test is for detecting elevated antibody titers. This is designed to be used in adult and
	adolescent populations for epidemiological studies and outbreak response as these
	patients may not seek medical attention when the isolation of Bordatella pertussis by
	culture or PCR would be likely. At this time, the serologic test results should not be relied
	for case confirmation of pertussis infection. This assay should not be used to and assess
	susceptibility/immunity to pertussis or for clinical diagnosis. It is limited to surveillance
	purposes only.
Method:	ELISA
Interfering Substances:	Specimen from patient vaccinated against B.pertussis toxin in < 6months or under age
	11yrs cannot be tested.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	This test is used for surveillance purpose only.

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TEST:	Borrelia burgdorferi serology
Synonym:	Borrelia burgdorferi IgG/IgM Antibody, Lyme Disease
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml whole blood
Specimen Volume (Minimum):	1 ml whole blood
Collect:	Red-top vacutainer tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	<ul> <li>NON-REACTIVE. Indicates no detectable antibodies to Borrelia burgdorferi. A negative result does not exclude a Lyme disease infection. Patients with early stages of infection or who have undergone antibiotic therapy may not produce measurable IgG/IgM antibodies. Additional specimens should be submitted in 2-4 weeks if Borrelia burgdorfe exposure has not been ruled out.</li> <li>REACTIVE. Antibodies to Borrelia burgdorferi have been detected. Sera from individuals with other pathogenic spirochetal diseases, bacterial and viral infections, and individuals with connective tissue autoimmune diseases or anti-nuclear antibody may also have antibodies which cross-react with B. burgdorferi.</li> <li>EQUIVOCAL—Immunological status cannot be determined, please re-draw patient in 2-4 weeks.</li> </ul>
Additional Information:	http://www.cdc.gov/lyme/
Purpose of Test:	Detect antibody to B. burgdorferi
Methods:	ELFA (enzyme –linked fluorescent immunoassay, Western Blot
Interfering Substances:	Icteric, hemolyzed, lipemic
Testing Site:	DHMH Laboratories Administration, Central Laboratory
Comment:	1770 Ashland Avenue, Baltimore, MD 21205 Your health care provider has ordered a laboratory test for the presence of Lyme Disease
	for you. Current Laboratory testing for Lyme Disease can be problematic and standard laboratory tests often result in false negative and false positive results, and if done too early, you may not have produced enough antibodies to be considered positive because your immune response requires time to develop antibodies. If you are tested for Lyme Disease and the results are negative, this does not necessarily mean you do no have Lyme Disease. If you continue to experience unexplained symptoms, you should contact your health care provider and inquire about the appropriateness of retesting or initial or additional treatment. The Western blot test will be used to confirm the presence of B. burgdorferi specific antibodies detected by the ELFA screening test on all Positive & Equivocal specimens.

TEST:	Botulism ( <i>Clostridium botulinum</i> –Adult and <i>Clostridium botulinum</i> –Infant) Must have consent of the State Epidemiologist before sending specimen to the Laboratory (410-767-6685).
sSynonym:	Botulism; <i>Clostridium botulinum</i> : Refer to instructions for <i>Clostridium botulinum</i> –Adult and <i>Clostridium botulinum</i> –Infant.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response: <b>410-925-3121 (24/7 emergency contact number)</b> Select Agents Microbiology Laboratory: 443-681-3954 Division of Microbiology Laboratory: 443-681-3952

TEST:	Brucella serology (CDC Referral)
Synonym:	Bang's Disease, Undulant fever, Malta Fever
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	2 weeks (CDC Referral)
Specimen Required:	Serum
Specimen identification:	Label tube with patients first and last name. The specimen/sample must be properly
	labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	-20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/brucellosis/index.html
Purpose of Test:	Detect antibody to Brucella
Method:	Brucella microagglutination test (BMAT)
Interfering Substances:	No serology available for B. canis or RB51.
	May have poor sensitivity for chronic or complicated brucellosis.
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen
	submission. Required supplemental information: Exposure and travel history, include other relevant risk factors; clinical symptoms, treatment and relevant lab results.

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TEST:	Brucella species, culture
Synonym:	Bang's Disease, Undulant fever, Malta Fever, and Rock of Gibraltar Fever
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952
Turnaround Time:	5 - 30 days [from specimen receipt in the Laboratory]
Specimen Required:	1. Blood or bone marrow
	2. Spleen, liver or abscess
	3. Serum-acute and convalescent-phases
	4. Isolate
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	<ol> <li>Blood: Collect appropriate blood volume and number of sets per routine laboratory protocol. Specimens should be inoculated into appropriate culture media within two (2) hours of collection.</li> </ol>
	2. Biopsied Tissue: Collect per laboratory protocol. Tissues must be kept moist; add several drops of sterile saline if necessary.
	3. Serum: At least 1 ml of serum. Follow standard laboratory protocol. Preferably serum refrigerated.
	4. Isolate: Pure culture, 24 hours old, growing on a sheep blood agar plate or slant.
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Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	1. Blood Cultures: Transport at room temperature. Hold them at ambient
	temperature until they are incubated. DO NOT REFRIGERATE.
	2. Tissue: Transport at room temperature, adding several drops of sterile normal
	saline to keep tissues moist for immediate processing. Keep the specimen chilled if
	the processing of the specimen will be delayed.
	3. Serum: Keep serum on cold packs.
	<ol> <li>Isolates: Transport at room temperature on a sealed sheep blood agar plate or slant.</li> </ol>
Specimen Rejection Criteria	The following rejection criteria are designed to prevent the reporting of inaccurate
-	results and to avoid misleading information that might lead to misdiagnosis and
	inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
Availability:	24 hours/day, 7days/week
Results and Interpretation:	Brucella species isolated/detected
	Brucella species not found
Additional Information:	Call 410-925-3121 before sending specimen to the Laboratory.
Purpose of Test:	To confirm the diagnosis of Brucella species.
Method:	LRN protocols
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Brucella species are highly infectious. PLEASE use a biological safety cabinet when
	working with specimens suspected of being Brucella species.
	Call 410-925-3121 before sending to the laboratory.

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TEST:	Burkholderia mallei and Burkholderia pseudomallei
Synonym:	B. mallei is the causative agent of Glanders; and
	B. pseudomallei is the causative agent of Melioidosis
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952
Turnaround Time:	4 - 8 days [from specimen receipt in the Laboratory]
Specimen Required:	1. Blood: Collect blood specimens before antibiotics are administered.
	2. Urine
	3. Abscesses, tissue aspirates, body fluids: Collect tissues and fluids rather than swabs, when possible.
	4. Isolate
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must
	be properly labeled and match the test requisition or electronic test order.
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DHMH-Laboratories Administration	The J. Mehsen Joseph Public Health Laboratory
Specimen Volume (Optimum):	<ol> <li>Blood: Collect appropriate volume and number of sets per laboratory protocol.</li> <li>Urine: 5 ml.</li> </ol>
	3. Abscesses, tissues and body fluids: Collect per routine laboratory protocol.
Specimen Volume (Minimum):	N/A
Collect:	1. Blood: Collect appropriate blood volume and number of sets as per routine laboratory protocol.
	2. Urine: Collect 5 ml. of midstream clean-catch specimen or a cauterization specimen.
	3. Abscesses, tissues aspirates, body fluids: Collect tissues and body fluids rather than swabs.
	4. Isolate: Pure culture, 24 hours old, growing on a sheep blood agar plate or slant.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	1. Blood: Transport at room temperature. Hold them at ambient temperature until
•	they are incubated. DO NOT REFRIGERATE.
	2. Urine: Transport in a sterile, well sealed container chilled using wet ice or cold
	packs.
	3. Abscesses, tissues, and fluids: Transport the specimen at room temperature for
	immediate processing. Keep the specimen chilled if processing of the specimen will
	be delayed.
	4. Isolate: Transport the specimen at room temperature on a sealed sheep blood agar
	plate or slant.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate
	results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
Availability:	24 hours/day, 7 days/week
Results and Interpretation:	B. mallei/B. pseudomallei isolated/detected.
·	B. mallei/B. pseudomallei not found.
Additional Information:	Call 410-925-3121 before sending specimen to the Laboratory.
Purpose of Test:	To confirm the diagnosis of <i>B. mallei</i> and <i>B. pseudomallei</i> .
Method:	LRN Protocols
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Call 410-925-3121 before sending to the Laboratory.

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TEST:	C. difficile Toxin (A and B)
Synonym:	Clostridium difficile toxin, C. diff
Laboratory/Phone:	Microbiology: 443-681-3952
Turnaround Time:	Two (2) days [from specimen receipt in the Laboratory]
Specimen Required:	Fresh, unpreserved stool specimen
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Two (2) grams
Specimen Volume (Minimum):	N/A
Collect:	Stool in a clean, unpreserved stool transport vial
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Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Unpreserved, shipped in insulated container with freezer pack
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate
	results and to avoid misleading information that might lead to misdiagnosis and
	inappropriate therapy. A request for a new specimen will provide appropriate materials
	and clinically relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
	<ul> <li>Formed stool</li> </ul>
	<ul> <li>Stool preserved in 10% formalin, SAF, or PVA</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Positive (Toxin A and/or Toxin B present) or Negative (No Toxin A or Toxin B detected)
Reference Range:	Negative
Additional Information:	Clostridium difficile can be grown and isolated on a stool culture, but its presence does
	not indicate whether the strain present is a toxin producer. It also does not distinguish
	between C. difficile colonization and overgrowth/infection.
Purpose of Test:	The Clostridium difficile toxin test is used to diagnose antibiotic-associated diarrhea and
	pseudomembranous colitis that is caused by C. difficile. It may also be ordered to detect
	recurrent disease.
Method:	EIA (Enzyme Immunoassay)
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
-	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	This test does not differentiate between Toxin A and Toxin B.

TEST:	Campylobacter Culture Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins– producing E. coli)
Synonym:	Stool culture for enteric pathogens; enteric pathogens; stool culture and sensitivity; feces culture: Refer to instructions for Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins–producing <i>E. coli</i> ).
Laboratory/Phone:	Microbiology-Enterics: 443-681-4570

TEST:	CDC Referrals (Serology)
Synonym:	CDC's Infectious Diseases Laboratories provides an online Test Directory that allows you
	to identify the right test for your needs.
	http://www.cdc.gov/laboratory/specimen-submission/list.html#B
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	Refer to CDC Test Directory
	http://www.cdc.gov/laboratory/specimen-submission/list.html#B
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
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Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	See CDC specific transport requirements.
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	Call 443-681-3938/3931 before sending specimen to State lab.
Purpose of Test:	Detect antibodies which may be due to a particular infectious agent
Methods:	Varies
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
-	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen
	submission. Required supplemental information: Exposure and travel history, include
	other relevant risk factors; clinical symptoms, treatment and relevant lab results.

TEST:	Chagas disease
Synonym:	Trypanosoma cruzi
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	-20°C (frozen).
Specimen Rejection Criteria:	Hemolysis, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	NEGATIVE: Antibodies to T. cruzi have not been detected and there is a high probability
	of non-infection or an early infection with low level of antibody present.
	EQUIVOCAL: The presence or absence of antibody to T. cruzi cannot be established.
	POSITIVE: Antibodies to T. cruzi, the causative agent of Chagas' disease were detected.
Additional Information:	http://www.cdc.gov/parasites/chagas/
Purpose of Test:	Detect antibodies which may be due to T. cruzi
Methods:	EIA
Interfering Substances:	Hemolysis
Testing Site:	DHMH. Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the
	assessment of a patient's health. Clinical correlation is required. Positive and Equivocal
	results will be forwarded to CDC for confirmation.

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TEST:	Chancroid Culture (Hemophilus ducreyi)
Synonym:	Haemophilus ducreyi culture: Refer to instructions for Hemophilus ducreyi Culture.
Laboratory/Phone:	Microbiology: 443-681-4570

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TEST:	Chikungunya IgM EIA
1201.	(Arbovirus Travel-Associated Panel)
	Test available based on patient's travel history.
Synonym:	Arthropod-borne virus: Chikungunya Virus
Laboratory/Phone:	443-681-3937
Turnaround Time:	5-10 business days during Arbovirus Season (excluding PRNT Testing)
Specimen Required:	Serum (blood)
Specimen Identification:	The specimen/sample must be properly labeled and include:
specification	1. The patient's name or unique patient/sample identifier matching the test
	requisition or electronic test order,
	<ol> <li>If appropriate, the date and time of specimen/sample collection, and</li> </ol>
	<ol> <li>Any additional information relevant and necessary for the test.</li> </ol>
	The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml serum
Specimen Volume (Minimum):	1 ml serum
Collect:	Red-top vacutainer, transfer serum to sterile tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form. Write "S" for serum in the
	"Arbovirus Travel-Associated Panel".
	For testing to be initiated the following information MUST be provided: date of onset,
	<b>date specimen collected, travel history, and flavivirus vaccination history.</b> Also please provide: patient's date of birth, diagnosis, symptoms, fatality, and whether patient is
	immunocompromised.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48
	hours, specimen can be frozen at -20°C and shipped on dry ice.
Specimen Rejection Criteria:	Grossly hemolyzed specimen, unlabeled specimen, leaking container, mismatch between
	labeling of specimen and test request form/electronic test order, and does not meet
Availability:	epidemiological criteria required for testing (e.g. travel history, etc.)
	Monday-Friday
Results and Interpretation:	<b><u>Negative</u></b> : No detectable IgM antibody, The result does not rule out Chikungunya virus infection. An additional sample should be tested within 7-14 days if early infection is
	suspected.
	<b>Equivocal:</b> Chikungunya virus IgM antibody cannot be determined, further testing by
	PRNT (plaque reduction neutralization test) is required. <u>Positive:</u> Presence of detectable IgM antibody, presumptive infection with Chikungunya
	virus. Confirmatory testing by PRNT (plaque reduction neutralization test) is required. A
	positive IgM result may not indicate a recent infection because IgM may persist for
	several months after infection.
Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name give to
	viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).
	Arboviruses that cause human encephalitis are members of three virus families: The
	Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae.
	For more information, see the CDC link at:
	http://www.cdc.gov/ncidod/dvbid/arbor/arbdet.htm Patients with travel history supporting suspicion of other arboviruses will be sent to the
	CDC for testing.
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Purpose of Test:	For the presumptive detection of IgM antibody to Chikungunya Virus. Confirmatory testing by PRNT may be required.
Method:	EIA (Screening) & PRNT (Plaque Reduction Neutralization Test) referral to the Centers for Disease Control and Prevention (CDC) for confirmatory testing.
Interfering Substances:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue Baltimore, MD 21205
Comment:	

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TEST:	Chlamydia Cell Culture
Synonym:	
Laboratory/Phone:	443-681-3937
Turnaround Time:	Within 10 business days
Specimen Required:	Swab: Endocervix, urethra, conjunctiva, nasopharynx, throat, rectum, vagina. For other
	sources, call laboratory to discuss.
	Place swab in ChlamTrans <sup>™</sup> transport tube. (Check expiration date of transport media.)
Specimen identification:	The specimen/sample must be properly labeled and include:
•	1. The patient's name or unique patient/sample identifier matching the test requisition
	or electronic test order,
	2. If appropriate, the date and time of specimen/sample collection, and
	3. Any additional information relevant and necessary for the test.
	The specimen/sample must be properly labeled and match the test requisition or
	electronic test order.
Specimen Volume (Optimum):	2ml of media already in transport tube
Specimen Volume (Minimum):	2ml of media already in transport tube
Collect:	Swab placed in ChlamTrans™ Transport media, or other commercial media stating it is
	appropriate for Chlamydia isolation.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Chlamydia trachomatis located under Virus/Chlamydia heading. Indicate specimen type
	next to test requested using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Transport at 2-8°C
	Must reach the lab within 2 days of collection
Specimen Rejection Criteria:	Too old, No patient ID on specimen, leaked, quantity not sufficient, no swab, expired
	transport, out of temp. range, no specimen received, broken, improper swab or
	collection kit, improper collection site, thick mucus, gross contamination, illegible ID,
	missing or incomplete lab slip (no site, date, gender, patient info., submitter info.),
	mismatched patient ID.
Availability:	Monday-Friday
Results and Interpretation:	Chlamydia trachomatis Isolated in cell culture.
	Chlamydia trachomatis not Isolated in cell culture.
	Chlamydia trachomatis toxic in cell culture. Resubmit.
Additional Information:	This test is limited to medico-legal specimens: cervical, rectal, male urethral; and non-
	cervical, non-rectal, and non-male urethral specimens.
Purpose of Test:	Diagnostic, qualitative detection of Chlamydia
Method:	Cell culture
Interfering Substances/Limitations:	A negative result does not exclude the possibility of infection. Interpret results in
	conjunction with other information.
	Do not use ChlamTrans™ if leakage, evaporation, contamination or pH changes are
	apparent.
	Store ChlamTrans <sup>™</sup> refrigerated.
	Do not freeze unless <-50°C. If frozen, must transport on dry ice.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	
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TEST:	Chlamydia Serology
Synonym:	Chlamydia Group antigen antibody (IgG) EIA
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot, separated serum at 2-8°C
	(refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being
	frozen.
Availability:	Monday through Friday
Results and Interpretation:	<b>POSITIVE</b> —Detectable IgG Chlamydial antibodies. Suggest immunological exposure to
	one or more chlamydial species.
	<b>NEGATIVE</b> —No detectable IgG Chlamydial antibodies. Suggest no prior immunological
	exposure to chlamydial species. Does not rule out recent exposure and collection of
	sample prior to development of IgG antibodies.
	EQUIVOCAL—Immunological exposure cannot be assessed.
Additional Information:	This test is not intended to replace culture
Purpose of Test:	For the detection of antibody to Chlamydia group antigen
Method:	EIA
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	This test does not differentiate between different species of Chlamydia. Serologic results
	should not be used as a sole means for diagnosis, treatment, or for the assessment of a
	patient's health. Clinical correlation is required

TEST:	Chlamydia trachomatis and Neisseria gonorrhoeae
	Nucleic Acid Amplification Test (NAAT)
Synonym:	Becton Dickinson's ProbeTec <sup>™</sup> Q <sup>X</sup> , Amplified DNA Assay
Laboratory/Phone:	Chlamydia Laboratory / 410-767-6154
Turnaround Time:	Within 7 business days
Specimen Required:	Endocervical swab Male urethral swab Male and female urine (first of the void)
Specimen identification:	Label specimen with the full name exactly matching form, date of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Swab: Tube, Prefilled with 2 ml of diluent Urine: Optimal quality specimen is 20-60 ml of "first of the void" urine collected in a plastic collection cup. Swirl to mix. Using a sterile transfer pipette, transfer 2-3 ml from cup into labeled BD urine tube so volume falls between the two fill lines on the tube. Do not surpass the top fill line.
Specimen Volume (Minimum):	Swab: Tube, Prefilled with 2 ml of diluent Urine: Collect a minimum of 4ml (20-60 best) in a plastic collection cup. Using a sterile transfer pipette, transfer 2-3 ml from cup into labeled BD urine tube so volume falls between the two fill lines on the tube. Volume must reach the lower fill line.
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Indicate specimen type next to test requested using the "Specimen Code" on form.           Packaging and Shipping*:         Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).           **Refer to current Federal regulations for specific shipping requirements.           specimen Rejection Criteria:         Svab: 2-30°C. Must test within 30 days of collection.           Urine: 2-8°C. Must test within 7 days of collection site, hick mucus, illegible ID, missin or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Results and Interpretation: <i>Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           Nesseria gonorrhoeae was actected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           Nesseria gonorrhoeae was actected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           Nesseria gonorrhoeae was actected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           Neteseria gonorrhoeae was not detected by Nucleic Acid Amplification USA Placement Amplification (SDA) method.           Netseria gonorrhoeae was not detected by Nucleic Acid Amplification USA Placement Amplification (SDA) method.           Neteseria gonorrhoeaee was not detected by Nucleic Acid Amplification </i>	Collect:	Swab: BD ProbeTec <sup>™</sup> Q <sup>X</sup> Collection Kit for Endocervical and Lesion Specimens (part
Urine: Sterile, preservative-free, leakproot, plastic specime collection cup. The patient should not have urinated for at least 1 hour prior to specimen collection. Collect 20-60 ml of "first of the void urine." Transfer 2-3 ml of swirfed neat urine into the BD collection tube between the two fill lines. Replace cap tightly.           ••••••••••••••••••••••••••••••••••••		#441357)or BD ProbeTec <sup>™</sup> Q <sup>X</sup> CT/GC Amplified DNA Assay Collection Kit for Male
should not have urinated for at least 1 hour prior to specime collection. Collect 20-60 ml of "first of the void urine." Transfer 2-3 ml of swirled neat urine into the BD collection tube between the two fill lines. Replace cap tighty.           "orm:         DHIMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type next to test requested using the "Specimen Code" on form.           "Packaging and Shipping":         Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).           "Refer to current Federal regulations for specific shipping requirements.           Specimens Rejection Criteria:         Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp. range, no specimen received, broken improper swab or collection kt, improper collection site, thick mucus, lilegible ID, missin or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Nesseria goorn/honee was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Chlomydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria goorn/hoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria goorn/hoeae was not detected by Nucleic Acid Amplification using the Strand Displacem		Urethral Specimens (part #441358).
ml of "first of the void urine," Transfer 2-3 ml of swirled neat urine into the BD collection tube between the two fill lines. Replace cap tighty.         Form:       DHIMH Form #4676 infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type next to test requested using the "Specimen Code" on form.         Packaging and Shipping*:       Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).         "Refer to current feedaral regulations for specific shipping requirements.         Specimen Rejection Criteria:       Swab: 2-30°C. Must test within 30 days of collection.         Urine: 2-8°C. Must test within 2 days of collection in urine: 2-8°C. Must test within 2 days of collection in improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab silp (no site, date, gender, patient info., submitter info.), mismatched patient ID.         Vavailability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was of detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was of detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was of detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.<		Urine: Sterile, preservative-free, leakproof, plastic specimen collection cup. The patient
tube between the two fill lines. Replace cap tightly.           Form:         DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)           Indicate specimen type next to test requested using the "Specimen Code" on form.           Packaging and Shipping*:         Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).           "Refer to current Federal regulations for specific shipping requirements.           Fransport Conditions:         Swab: 2-30°C. Must test within 30 days of collection.           Urine: 2-8°C. Must test within 7 days of collection.         Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp, range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Results and Interpretation:         • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Specimen failed in assay.		should not have urinated for at least 1 hour prior to specimen collection. Collect 20-60
Form:         DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type next to test requested using the "Specimen Code" on form.           *ackaging and Shipping*:         Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).           *Refer to current Feedral regulations for specific shipping requirements.           fransport Conditions:         Swab: 2-30°C. Must test within 30 days of collection. Urine: 2-8°C. Must test within 7 days of collection. Urine: 2-8°C. Must test within 7 days of collection and the specific shipping requirements.           specimen Rejection Criteria:         Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp, range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missin or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Results and Interpretation:         • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was not detected by Nucl		ml of "first of the void urine." Transfer 2-3 ml of swirled neat urine into the BD collection
Indicate specimen type next to test requested using the "Specimen Code" on form.           Packaging and Shipping*:         Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).           "Refer to current Federal regulations for specific shipping requirements.           fransport Conditions:         Swab: 2-30°C. Must test within 30 days of collection.           Urine: 2-8°C. Must test within 7 days of collection.         Urine: 2-8°C. Must test within 7 days of collection.           specimen Rejection Criteria:         Too old, No patient ID on specimen, >60 ml of collectod urine, leaked, quantity not sufficient, no swab, expired transport, out of temp, range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Results and Interpretation:         • Chiamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Specimen failed in assay.           Not applicable.           Additional Information:           Reference Range:           Not applicable.           Additional Informatio		tube between the two fill lines. Replace cap tightly.
Packaging and Shipping*:         Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).           *Refer to current Federal regulations for specific shipping requirements.           Specimen Rejection Criteria:         Swab: 2:30°C. Must test within 30 days of collection.           Urine: 2-8°C. Must test within 7 days of collection.         Urine: 2-8°C. Must test within 7 days of collection.           Specimen Rejection Criteria:         Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp, range, no specime received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Results and Interpretation:         • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Specimen faile in assay.           Reference Range:         Not applicable.           Not applicable.         Strand Displacement Amplification (SDA)	Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).         "Refer to current Federal regulations for specific shipping requirements.         Fransport Conditions:       Swab: 2-30°C. Must test within 30 days of collection.         Urine: 2-8°C. Must test within 7 days of collection.       Urine: 2-8°C. Must test within 7 days of collection.         Specimen Rejection Criteria:       Too old, No patient 1D on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp. range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.         Availability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhaeee was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhaeee was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhaeee was dot detected by Nuclei Acid Amplification using the Strand Displacement Amplification (SDA)         Purpose of Test:       Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhaeea DNA.         Vethod:       Strand Displacement Amplification (SDA)         <		Indicate specimen type next to test requested using the "Specimen Code" on form.
pages 9 & 10 for triple packing guidance).         *Refer to current Federal regulations for specific shipping requirements.         fransport Conditions:	Packaging and Shipping*:	
*Refer to current Federal regulations for specific shipping requirements.           Fransport Conditions:         Swab: 2-30°C. Must test within 30 days of collection.           Urine: 2-8°C. Must test within 7 days of collection.         Specimen Rejection Criteria:           Specimen Rejection Criteria:         Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp, range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Results and Interpretation:         Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was on detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was on detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA)           • Neisseria gonorrhoeae was on detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA)           • Not applicable.           Additional Information:		conditions of transport they cannot break, be punctured or leak their contents (Refer to
Transport Conditions:       Swab: 2-30°C. Must test within 30 days of collection.         Urine: 2-8°C. Must test within 7 days of collection.         Specimen Rejection Criteria:       Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp. range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.         Availability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA)         • Not applicable.         Additional Information:       Restricted testing (preapproved submitters only, call 410-767-6154)         * Urros of Test:       Direct, qual		pages 9 & 10 for triple packing guidance).
Urine: 2-8°C. Must test within 7 days of collection.         Specimen Rejection Criteria:       Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp, range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.         Availability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Displacement Amplification (SDA) method.         • Specimen failed in assay.         Yuethod:         Not applicable.         Additional Information:         Restricted testing (preapproved submitters only, call 410-767-6154)         Purpose of Test:       Direct, qualitative detection of Chlamydia trachomatis.         Nuethod:       Interfering substances:         • swab - blood > 60%       • urine - blood > 1%         • urine - blood >		*Refer to current Federal regulations for specific shipping requirements.
Specimen Rejection Criteria:         Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not sufficient, no swab, expired transport, out of temp, range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.           Availability:         Monday-Friday           Results and Interpretation:         • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.           • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement falled in assay.           Reference Range:         Not applicable.           Variand Displacement Amplification (SDA)         Interfering substances/Limitations:           Interfering Substances/Limitations:         Interfering substances: • swab · blood > 60% • urine · blood > 1% Limitations:           Interfering Substances/Limitations:         Interfering subchares: • on ot detect plasmid free variants of Chlamydia trachomatis. Only cell culture isolation should be used when testing for the evaluation of suggested sexu	Transport Conditions:	Swab: 2-30°C. Must test within 30 days of collection.
sufficient, no swab, expired transport, out of temp. range, no specimen received, broken improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.         Availability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement failed in assay.         Reference Range:       Not applicable.         Additional Information:       Restricted testing (preapproved submitters only, call 410-767-6154)         Purpose of Test:       Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .         Method:       Strand Displacement Amplification (SDA)         Interfering Substances/Limitations:       Interfering substances:         • swab - b		Urine: 2-8°C. Must test within 7 days of collection.
improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID. Availability: Monday-Friday Results and Interpretation: - Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method. - Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method. - Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method. - Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method. - Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method. - Specimen failed in assay. Reference Range: Not applicable. Additional Information: Restricted testing (preapproved submitters only, call 410-767-6154) Purpose of Test: Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA . Strand Displacement Amplification (SDA) Interfering Substances/Limitations: Interfering substances: - swab - blood > 60% - urine - blood > 1% Limitations: Endocervical specime adequacy cannot be determined. A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information. Does not detect plasmid free variants of Chlamydia trachomatis. Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes. GC assay may cross react with <i>N. cinere</i> and <i>N. lactamica</i> . Festing Site: DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205	Specimen Rejection Criteria:	Too old, No patient ID on specimen, >60 ml of collected urine, leaked, quantity not
or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched patient ID.         Availability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Specimen failed in assay.         Reference Range:       Not applicable.         Additional Information:       Restricted testing (preapproved submitters only, call 410-767-6154)         Purpose of Test:       Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .         Method:       Strand Displacement Amplification (SDA)         Interfering Substances/Limitations:       Interfering substances:         • swab - blood > 60%       urine - blood > 1%         Limitations:       Endocervical specimen adequacy cannot be determined.         A negative test result does not exclude the possibility of infection. Interpret		sufficient, no swab, expired transport, out of temp. range, no specimen received, broken
patient ID.         Availability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Specimen failed in assay.         Reference Range:         Not applicable.         Additional Information:         Restricted testing (preapproved submitters only, call 410-767-6154)         Purpose of Test:       Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .         Method:       Strand Displacement Amplification (SDA)         Interfering Substances/Limitations:       Interfering substances:         Interfering substances/Limitations:       Swab - blood > 60%         Endocervical specimen adequacy cannot be determined.       A		improper swab or collection kit, improper collection site, thick mucus, illegible ID, missing
Availability:       Monday-Friday         Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Specimen failed in assay.         Reference Range:         Not applicable.         Additional Information:         Restricted testing (preapproved submitters only, call 410-767-6154)         Purpose of Test:         Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .         Method:         Interfering Substances/Limitations:         Interfering Substances/Limitations:         Interfering substances/Limitations:         Endocervical specimen adequacy cannot be determined.         A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.         Does not detect plasmid free variants of Chlamydia trachomatis.         Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.		or incomplete lab slip (no site, date, gender, patient info., submitter info.), mismatched
Results and Interpretation:       • Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Specimen failed in assay.         Reference Range:       Not applicable.         Additional Information:       Restricted testing (preapproved submitters only, call 410-767-6154)         Durpose of Test:       Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .         Method:       strand Displacement Amplification (SDA)         Interfering Substances/Limitations:       Interfering substances:         • swab - blood > 60%       • urine - blood > 1%         Limitations:       Endocervical specimen adequacy cannot be determined.         A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.         Does not detect plasmid free variants of Chlamydia trachomatis.         Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.         Ge assay may cross react with		patient ID.
the Strand Displacement Amplification (SDA) method.• Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.• Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.• Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.• Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.• Specimen failed in assay.Reference Range:Not applicable.Additional Information:Restricted testing (preapproved submitters only, call 410-767-6154)Purpose of Test:Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .Vethod:Strand Displacement Amplification (SDA)Interfering Substances/Limitations:Interfering substances:• swab - blood > 60%• urine - blood > 1%Limitations:Endocervical specimen adequacy cannot be determined.A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.Does not detect plasmid free variants of Chlamydia trachomatis.Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.Gassay may cross react with N. cinerea and N. lactamica.Festing Site:DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205	Availability:	Monday-Friday
<ul> <li>Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Specimen failed in assay.</li> <li>Reference Range:</li> <li>Not applicable.</li> <li>Additional Information:</li> <li>Restricted testing (preapproved submitters only, call 410-767-6154)</li> <li>Purpose of Test:</li> <li>Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .</li> <li>Method:</li> <li>Strand Displacement Amplification (SDA)</li> <li>Interfering Substances/Limitations:</li> <li>Inswab - blood &gt; 60%</li> <li>urine - blood &gt; 1%</li> <li>Limitations:</li> <li>Endocervical specimen adequacy cannot be determined.</li> <li>A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.</li> <li>Does not detect plasmid free variants of Chlamydia trachomatis.</li> <li>Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.</li> <li>GC assay may cross react with N. cinerea and N. lactamica.</li> <li>Testing Site:</li> <li>DHMH Laboratories Administration, Central Laboratory</li> <li>1770 Ashland Avenue, Baltimore, Maryland 21205</li> </ul>	Results and Interpretation:	Chlamydia trachomatis plasmid DNA was DETECTED by Nucleic Acid Amplification using
using the Strand Displacement Amplification (SDA) method.         Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         Specimen failed in assay.         Reference Range:         Not applicable.         Additional Information:         Restricted testing (preapproved submitters only, call 410-767-6154)         Durpose of Test:         Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .         Method:         Interfering Substances/Limitations:         Interfering substances:         • symbol > blood > 60%         • urine - blood > 1%         Limitations:         Endocervical specimen adequacy cannot be determined.         A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.         Does not detect plasmid free variants of Chlamydia trachomatis.         Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.         GC assay may cross react with N. cinerea and N. lactamica.         Testing Site:       DHMH Laboratories Administration, Central Laboratory         1770 Ashland Avenue, Baltimore, Maryland 21205		the Strand Displacement Amplification (SDA) method.
<ul> <li>Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Specimen failed in assay.</li> <li>Not applicable.</li> <li>Additional Information:</li> <li>Restricted testing (preapproved submitters only, call 410-767-6154)</li> <li>Purpose of Test:</li> <li>Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA.</li> <li>Method:</li> <li>Strand Displacement Amplification (SDA)</li> <li>Interfering Substances/Limitations:</li> <li>Interfering substances:         <ul> <li>swab - blood &gt; 60%</li> <li>urine - blood &gt; 1%</li> <li>Limitations:</li> <li>Endocervical specimen adequacy cannot be determined.</li> <li>A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.</li> <li>Does not detect plasmid free variants of Chlamydia trachomatis.</li> <li>Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.</li> <li>GC assay may cross react with N. cinerea and N. lactamica.</li> </ul> </li> <li>Festing Site:</li> <li>DHMH Laboratories Administration, Central Laboratory</li> <li>1770 Ashland Avenue, Baltimore, Maryland 21205</li> </ul>		Chlamydia trachomatis plasmid DNA was not detected by Nucleic Acid Amplification
<ul> <li>Neisseria gonorrhoeae was detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.</li> <li>Specimen failed in assay.</li> <li>Not applicable.</li> <li>Additional Information:</li> <li>Restricted testing (preapproved submitters only, call 410-767-6154)</li> <li>Purpose of Test:</li> <li>Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA.</li> <li>Method:</li> <li>Strand Displacement Amplification (SDA)</li> <li>Interfering Substances/Limitations:</li> <li>Interfering substances:         <ul> <li>swab - blood &gt; 60%</li> <li>urine - blood &gt; 1%</li> <li>Limitations:</li> <li>Endocervical specimen adequacy cannot be determined.</li> <li>A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.</li> <li>Does not detect plasmid free variants of Chlamydia trachomatis.</li> <li>Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.</li> <li>GC assay may cross react with N. cinerea and N. lactamica.</li> </ul> </li> <li>Festing Site:</li> <li>DHMH Laboratories Administration, Central Laboratory</li> <li>1770 Ashland Avenue, Baltimore, Maryland 21205</li> </ul>		using the Strand Displacement Amplification (SDA) method.
Displacement Amplification (SDA) method.• Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.• Specimen failed in assay.Reference Range:Not applicable.Additional Information:Restricted testing (preapproved submitters only, call 410-767-6154)Purpose of Test:Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .Method:Strand Displacement Amplification (SDA)Interfering Substances/Limitations:Interfering substances:• swab - blood > 60%• urine - blood > 1%Limitations:Endocervical specimen adequacy cannot be determined.A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.Does not detect plasmid free variants of Chlamydia trachomatis. Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes. GC assay may cross react with N. cinerea and N. lactamica.Festing Site:DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205		
• Neisseria gonorrhoeae was not detected by Nucleic Acid Amplification using the Strand Displacement Amplification (SDA) method.         • Specimen failed in assay.         Reference Range:       Not applicable.         Additional Information:       Restricted testing (preapproved submitters only, call 410-767-6154)         Purpose of Test:       Direct, qualitative detection of Chlamydia trachomatis and Neisseria gonorrhoeae DNA .         Method:       Strand Displacement Amplification (SDA)         Interfering Substances/Limitations:       Interfering substances:         • swab - blood > 60%       urine - blood > 1%         Limitations:       Endocervical specimen adequacy cannot be determined.         A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.         Does not detect plasmid free variants of Chlamydia trachomatis.         Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.         GC assay may cross react with N. cinerea and N. lactamica.         Festing Site:       DHMH Laboratories Administration, Central Laboratory         1770 Ashland Avenue, Baltimore, Maryland 21205		Displacement Amplification (SDA) method.
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Reference Range:       Not applicable.         Additional Information:       Restricted testing (preapproved submitters only, call 410-767-6154)         Purpose of Test:       Direct, qualitative detection of <i>Chlamydia trachomatis</i> and <i>Neisseria gonorrhoeae</i> DNA .         Method:       Strand Displacement Amplification (SDA)         Interfering Substances/Limitations:       Interfering substances:         •       swab - blood > 60%         •       urine - blood > 1%         Limitations:       Endocervical specimen adequacy cannot be determined.         A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.         Does not detect plasmid free variants of <i>Chlamydia trachomatis</i> .         Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.         GC assay may cross react with <i>N. cinerea</i> and <i>N. lactamica</i> .         Testing Site:       DHMH Laboratories Administration, Central Laboratory         1770 Ashland Avenue, Baltimore, Maryland 21205		
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• urine - blood > 1%         Limitations:         Endocervical specimen adequacy cannot be determined.         A negative test result does not exclude the possibility of infection. Interpret result in conjunction with other information.         Does not detect plasmid free variants of <i>Chlamydia trachomatis</i> .         Only cell culture isolation should be used when testing for the evaluation of suggested sexual abuse or other medico-legal purposes.         GC assay may cross react with <i>N. cinerea</i> and <i>N. lactamica</i> .         Festing Site:       DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205	Interfering Substances/Limitations:	
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	resuing Site:	
Lomment:		1770 Ashiand Avenue, Baltimore, Maryland 21205
	Comment:	

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TEST:	<i>Clostridium botulinum</i> –Adult MUST HAVE CONSENT OF THE STATE EPIDEMIOLOGIST BEFORE SENDING SPECIMEN TO THE LABORATORY (410-767-6685).
Synonym:	Botulism
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952
Turnaround Time:	3-7 days [from specimen receipt in the Laboratory]
Continued Next Page>	

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Specimen Required:	Suspected foodborne botulism cases:	
	Suitable specimens for examination are: serum, feces, vomitus, gastric contents.	
	Suspected wound botulism cases:	
	Suitable specimens for examination are: serum, tissue, feces.	
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,	
	specimen type/source, and the date and time of collection. The specimen/sample must	
	be properly labeled and match the test requisition or electronic test order.	
Specimen Volume (Optimum):	Serum: At least 10 ml (obtained from using at least 20 ml of whole blood).	
Specimen Volume (Minimum):	N/A	
Collect:	Serum: Collect using routine laboratory protocol using the red top or separator type	
	tube (NO anticoagulants).	
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)	
	Indicate specimen type using the "Specimen Code" on form.	
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal	
· · · · · · · · · · · · · · · · · · ·	conditions of transport they cannot break, be punctured or leak their contents (Refer to	
	pages 9 & 10 for triple packing guidance).	
	*Refer to current Federal regulations for specific shipping requirements.	
Transport Conditions:	Serum: Transport to the Laboratory on wet ice or cold packs. If an unavoidable delay of	
	several days is anticipated, the specimen should be kept frozen and then packed in an	
	insulated container with dry ice and proper cushioning material for shipment.	
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate	
specificit rejection citteria.	results and to avoid misleading information that might lead to misdiagnosis and	
	inappropriate therapy. A request for a new specimen will provide appropriate materials	
	and clinically relevant information to support good patient care.	
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>	
	<ul> <li>Non-sterile or leaking container</li> </ul>	
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>	
	<ul> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> </ul>	
	broken specificity sumple container	
	The wrong specific for test request	
	<ul> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> </ul>	
	megiste of no patient mornation on the specifien	
	Expired transport media	
Availability:	24 hours/day, 7 days/week	
Results and Interpretation:	Clostridium botulinum toxin detected/not detected.	
Additional Information:	Must have consent of the State Epidemiologist before sending specimen to the	
	Laboratory (410-767-6685).	
Purpose of Test:	To confirm the presence of <i>Clostridium botulinum</i> toxins	
Method:	LRN Methods	
Interfering Substances:	If the patient has been taking any medication that might interfere with toxin assays or	
	culturing of the stool, the Laboratory should be notified. For example, it has been	
	demonstrated that anticholinesterase drugs given orally to patients for myasthenia gravi	
	can interfere with mouse botulinum toxin assays of stool extracts.	
Testing Site:	DHMH Laboratories Administration, Central Laboratory	
	1770 Ashland Avenue, Baltimore, Maryland 21205	
Comment:	PHYSICIAN MUST CALL FOR A CONSULT BEFORE SENDING SPECIMEN. SPECIMENS ARE NOT	
comment.	PROCESSED UNTIL THE CASE IS APPROVED FOR TESTING BY THE STATE EPIDEMIOLOGIST	

 TEST:
 Clostridium botulinum–Infant MUST HAVE CONSENT OF THE STATE EPIDEMIOLOGIST BEFORE SENDING SPECIMEN TO THE LABORATORY (410-767-6685).

 Synonym:
 Botulism

 Laboratory/Phone:
 Office of Laboratory Emergency Preparedness and Response: 410-925-3121 (24/7 emergency contact number) Select Agents Microbiology Laboratory: 443-681-3954 Division of Microbiology Laboratory: 443-681-3952

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Turnaround Time:	2.20 days. [from specimen receipt in the Laboratory]
	3-30 days [from specimen receipt in the Laboratory]
Specimen Required:	Suspected infant botulism cases: Suitable specimens: Stool, rectal swabs (not necessary to collect serum.)
Charling an Identification.	
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must be
	properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Stool: 10-50 grams (English walnut size)
Specimen Volume (Minimum): Collect:	N/A Stack Collect in a starile well cooled unbrook allo container. Shin on cold packs If
Collect:	Stool: Collect in a sterile, well-sealed, unbreakable container. Ship on cold packs. If
	delayed, freeze stool specimen and ship frozen.
	Enema (if needed): Use minimal amount of sterile water or non-bacteriostatic water,
-	place 20 ml of liquid into a sterile, well-sealed, unbreakable container.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Stool: Transport to the Laboratory on wet ice or cold packs. If an unavoidable delay of
	several days is anticipated, the specimen should be kept frozen and then packed in an
	insulated container with dry ice and proper cushioning material for shipment.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results
	and to avoid misleading information that might lead to misdiagnosis and inappropriate
	therapy. A request for a new specimen will provide appropriate materials and clinically
	relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	Expired transport media
Availability:	24 hours/day, 7 days/week
Results and Interpretation:	Clostridium botulinum toxin detected/not detected.
Additional Information:	Must have consent of the State Epidemiologist before sending specimen to the
	Laboratory (410-767-6685).
Purpose of Test:	To confirm the presence of Clostridium botulinum toxin in the specimen.
Method:	LRN Methods
Interfering Substances:	Glycerin Enema will interfere with the recovery of Clostridium botulinum toxin.
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	If the patient has been taking any medication that might interfere with toxin assays or
	culturing of the stool, the Laboratory should be notified. For example, it has been
	demonstrated that anticholinesterase drugs given orally to patients for myasthenia gravis
	can interfere with mouse botulinum toxin assays of stool extracts.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	PHYSICIAN MUST CALL FOR A CONSULT BEFORE SENDING SPECIMEN. SPECIMENS ARE NOT
	PROCESSED UNTIL THE CASE IS APPROVED FOR TESTING BY THE STATE EPIDEMIOLOGIST
	(410-767-6685).

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TEST:	Clostridium difficile toxin
Synonym:	C. diff, C. difficile Toxin (A and B): refer to instructions for C. diff Toxin
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:	Clostridium perfringens Culture	
Synonym:	Clostridium perfringens Culture: Refer to instructions for Foodborne Pathogens (Bacillus	
	cereus, Clostridium perfringens, Staph aureus).	
Laboratory/Phone:	Microbiology 443-681-3952	

TEST:	Corynebacterium diptheriae culture (Diptheria)	
Synonym:	Corynebacterium diptheriae culture: Refer to instructions for Diptheria Culture.	
Laboratory/Phone:	Microbiology / 443-681-3952	

#### TEST: Coxiella serology Synonym: Coxiella burnetii, Q fever Laboratory/Phone: 443-681-3938/3931 Turnaround Time: 5 business days Specimen Required: Serum Specimen identification: The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order. Specimen Volume (Optimum): 2 ml. (Whole Blood) Specimen Volume (Minimum): 1 ml. (Whole Blood) Collect: Red-top vacutainer tube Form: DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form. Packaging and Shipping\*: Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). \*Refer to current Federal regulations for specific shipping requirements. **Transport Conditions:** Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or 20°C (frozen). Specimen Rejection Criteria: Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being frozen Availability: Monday through Friday **Results and Interpretation:** Titer ≥ 1:16 in both Phase I and Phase II antigen suggests a C. burnetii infection. Phase I antibody titers of greater than or equal to Phase II antibody titers are consistent with a chronic infection or convalescent phase Q fever. Titers < 1:16 in Phase I with titers >1:256 in Phase II antigen suggests a C. burnetii infection. Titer < 1:16 in both Phase I and Phase II antigen. No antibody detected. This result is seen in persons with either no C. burnetii infection or with an early infection. If Q fever suspected, collect a second specimen in 2-3 weeks. A 4-fold IgG antibody endpoint titer increase is considered supportive evidence of current or recent acute infection. Additional Information: http://www.cdc.gov/qfever/ Purpose of Test: Detect IgG antibodies which may be due to Coxiella burnetii infections Methods: IFA Interfering Substances: Icteric, hemolyzed, lipemic specimen **Testing Site:** DHMH. Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205 Comment: Serologic responses are time dependent. Specimens obtained too early in the infection may not contain detectable antibody levels. If Q fever is suspected obtain a second specimen 2-3 weeks later.

TEST:	Coxsackie Virus, Virus Culture	
Synonym:	Coxsackie Virus: Refer to instructions for Virus Culture.	
Laboratory/Phone:	Virology: 443-681-3934	

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TEST:	Cryptococcal antigen	
Synonym:	Cryptococcus neoformans antigen	
Laboratory/Phone:	443-681-3938/3931	
Turnaround Time:	5 business days	
Specimen Required:	Serum or cerebrospinal fluid (CSF)	
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.	
Specimen Volume (Optimum):	2 ml. (Whole Blood & CSF)	
Specimen Volume (Minimum):	1 ml. (Whole Blood & CSF)	
Collect:	Red Top vacutainer tube (Whole blood); CSF (Sterile container)	
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)	
	Indicate specimen type using the "Specimen Code" on form.	
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal	
	conditions of transport they cannot break, be punctured or leak their contents (Refer to	
	pages 9 & 10 for triple packing guidance).	
	*Refer to current Federal regulations for specific shipping requirements.	
Transport Conditions:	Collect ASAP after onset. Ship promptly on cold packs. Do not freeze.	
Specimen Rejection Criteria:	Hemolysis; insufficient volume	
Availability:	Monday through Friday	
Results and Interpretation:	<b>POSITIVE</b> <i>Cryptococcus neoformans</i> antigen detected. Additional follow-up and culture strongly recommended.	
	<b>NEGATIVE</b> — <i>Cryptococcus neoformans</i> antigen not detected. If status of patient suggest a cryptococcal infection, subsequent specimens and culture strongly recommended.	
Additional Information:		
Purpose of Test:	For the detection of <i>Cryptococcus neoformans</i> capsular polysaccharide antigens in serum or CSF	
Method:	Latex agglutination	
Interfering Substances:	Macroglobulins (e.g. Rheumatoid factors), hemolysis, lipemic	
Testing Site:	DHMH Laboratories Administration, Central Laboratory	
-	1770 Ashland Avenue, Baltimore, Maryland 21205	
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.	

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TEST:	Cysticercosis serology (CDC Referral)	
Synonym:	Neurocysticercosis, Taenia solium, cysitcercus	
Laboratory/Phone:	443-681-3938/3931	
Turnaround Time:	18 business days (CDC Referral)	
Specimen Required:	Serum, plasma, CSF	
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order	
Specimen Volume (Optimum):	2 ml. (Whole Blood & CSF)	
Specimen Volume (Minimum):	0.5 ml. (Whole Blood & CSF)	
Collect:	Red-top vacutainer tube (serum); lavender- top vacutainer tube (plasma); sterile container (CSF)	
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)	
	Indicate specimen type using the "Specimen Code" on form.	
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.	
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or - 20°C (frozen).	
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Availability:	Monday through Friday	
Results and Interpretation:	Given on CDC report	
Additional Information:	http://www.cdc.gov/parasites/cysticercosis/	
Purpose of Test:	or the detection of an antibody response to cysticerci lesions.	
Method:	Immunoblot, Western blot, Antibody detection	
Interfering Substances: Substance known to interfere with immunoassays include: bilirubin, lipids,		
	hemoglobin	
Processing Site for CDC referral:	DHMH. Laboratories Administration, Central Laboratory	
	1770 Ashland Avenue, Baltimore, MD 21205	
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen	
	submission. Required supplemental information: Exposure and travel history, include	
other relevant risk factors; clinical symptoms, treatment and relevant lab re		

TEST:	Cytomegalovirus (CMV) Culture	
Synonym:	Cytomegalovirus (CMV): Refer to instructions for Virus Culture.	
Laboratory/Phone:	Virology: 443-681-3934	

TEST:	Cytomegalovirus Serology
Synonym:	CMV, Cytomegalovirus IgG antibody
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or 20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	POSITIVEIgG antibodies to CMV detected NEGATIVE—IgG antibodies to CMV not detected EQUIVOCAL—Immunological status cannot be assessed
Additional Information:	
Purpose of Test:	For the detection of antibody to CMV
Method:	ELFA – enzyme-linked fluorescent immunoassay
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

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TEST:	Deerfly fever
Synonym:	Francisella tularensis; Pasteurella tularensis, tularemia, rabbit fever, deerfly fever, Ohara's
	disease, Francis disease: Refer to instructions for Francisella tularensis Culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Dengue Fever IgM EIA
	(Arbovirus Travel-Associated Panel)
	Test available based on patient's travel history.
Synonym:	Arthropod-borne virus: Dengue Fever
Laboratory/Phone:	443-681-3937
Turnaround Time:	5-10 business days during Arbovirus Season (excluding PRNT Testing)
Specimen Required:	Serum (blood)
Specimen Identification:	The specimen/sample must be properly labeled and include:
	1. The patient's name or unique patient/sample identifier matching the test
	requisition or electronic test order,
	2. If appropriate, the date and time of specimen/sample collection, and
	3. Any additional information relevant and necessary for the test.
	The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml serum
Specimen Volume (Minimum):	1 ml serum
Collect:	Red-top vacutainer, transfer serum to sterile tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form. Write "S" for serum in the
	"Arbovirus Travel-Associated Panel".
	For testing to be initiated the following information MUST be provided: date of onset,
	date specimen collected, travel history, and flavivirus vaccination history. Also please
	provide: patient's date of birth, diagnosis, symptoms, fatality, and whether patient is
	immunocompromised.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48
	hours, specimen can be frozen at -20°C and shipped on dry ice.
Specimen Rejection Criteria:	Grossly hemolyzed specimen, unlabeled specimen, leaking container, mismatch between
	labeling of specimen and test request form/electronic test order, and does not meet
	epidemiological criteria required for testing (e.g. travel history, etc.).
Availability:	Monday-Friday
Results and Interpretation:	Negative: No detectable IgM antibody, The result does not rule out Dengue virus
	infection. An additional sample should be tested within 7-14 days if early infection is
	suspected.
	Equivocal: Dengue virus IgM antibody cannot be determined, further testing by PRNT
	(plaque reduction neutralization test) is required.
	Positive: Presence of detectable IgM antibody, presumptive infection with Dengue virus.
	Confirmatory testing by PRNT (plaque reduction neutralization test) is required. A
	positive IgM result may not indicate a recent infection because IgM may persist for
	several months after infection.
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Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name give to
	viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).
	Arboviruses that cause human encephalitis are members of three virus families: The
	Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae.
	For more information, see the CDC link at:
	http://www.cdc.gov/ncidod/dvbid/arbor/arbdet.htm
	Patients with travel history supporting suspicion of other arboviruses will be sent to the
	CDC for testing.
Purpose of Test:	For the presumptive detection of IgM antibody to Dengue Virus. Confirmatory testing by
	PRNT may be required.
Method:	EIA (Screening) & PRNT (Plaque Reduction Neutralization Test) referral to the Centers for Disease Control and Prevention (CDC) for confirmatory testing.
Interfering Substances:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue Baltimore, MD 21205
Comment:	

TEST:	Diptheria Culture
Synonym:	Corynebacterium diptheriae culture
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	48-72 hrs [from specimen receipt in the Laboratory]
Specimen Required:	Respiratory illness: Throat and nasopharyngeal swabs. Cutaneous diphtheria: Skin, throat and nasopharynx.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	Swab infected areas thoroughly, getting swab well into membranes or other lesions present. Inoculate stuart transport media and break off stick where handled. Leave swab in the tube and tighten cap.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Definitive identification of Corynebacterium diptheriae. Toxigenicity testing has to follow identification.
Reference Range:	Corynebacterium diphtheria <u>NOT</u> found.
Additional Information:	Take culture before starting antimicrobial therapy – if possible.
Purpose of Test:	Diagnosis of toxigenic strains of Corynebacterium diptheriae and antibiotic treatment are essential in limiting spread of infection.
Method:	Culture and smear

Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	When C. diptheriae is isolated, the isolate is forwarded to the Centers for Disease Control
	and Prevention (CDC) for detection of the toxin.
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TEST:	Disk Diffusion Susceptibility Testing
Synonym:	Disk Diffusion Susceptibility Testing: Refer to instructions for Antimicrobial Susceptibility
	Test
Laboratory/Phone:	Microbiology 443-681-3952

TEST:	E. coli O157 typing
Synonym:	Isolate for <i>E. coli</i> O157 serotyping (referral isolate); and other than O157 serotypes.
Laboratory/Phone:	Microbiology-Enterics, 443-681-4570
Turnaround Time:	4 – 10 days [from specimen receipt in the Laboratory]
Specimen Required:	Pure isolate of <i>E. coli</i>
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Sorbitol negative <i>E. coli</i> from culture.
Specimen Volume (Minimum):	N/A
Collect:	N/A
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store and ship at room temperature, ship as quickly as possible.
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	E. coli O157 identified and H7 antigens identified.
Reference Range:	No <i>E. coli</i> O157 detected
Additional Information:	Isolates submitted for <i>E. coli</i> O157 typing will be sub-cultured upon arrival and tested for shiga toxins, O157 antigen and biochemically identified as well as tested for H7 if needed.
Purpose of Test:	Detect the presence of <i>E. coli</i> O157
Method:	Culture and serotyping
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Eastern Equine Encephalitis Virus (EEEV)
	(Arbovirus Endemic Panel)
	Panel includes WNV IgM, SLE IgM, and EEEV IgM. LaCrosse (LAC) IgM testing available
	based on patient's travel history.
Synonym:	Arthropod-borne virus: WNV (West Nile Virus), EEEV (Eastern Equine Encephalitis Virus),
	SLEV (St. Louis Encephalitis Virus): Refer to instructions for Arbovirus Endemic Panel.
Laboratory/Phone:	Virology: 443-681-3937

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TEST:	Echinococcus serology (CDC Referral)
Synonym:	Echinococcosis, Hydatitd Disease, Echinococcus granulosus, parasite
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	18 business days (CDC Referral)
Specimen Required:	Serum, plasma
Specimen Identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2ml. (Whole Blood)
Specimen Volume (Minimum):	0.5ml. (Whole Blood)
Collect:	Red-top vacutainer tube (serum) Lavendat-top vacutainer (plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or - 20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/parasites/echinococcosis/
Purpose of Test:	Detect antibodies which may be due Echinococcus parasite infections
Methods:	Immunoblot, Western blot, Antibody detection
Interfering Substances:	Substance known to interfere with immunoassays include: bilirubin, lipids, and hemoglobin
Processing Site for CDC referral:	DHMH. Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen submission. Required supplemental information: Exposure and travel history, include other relevant risk factors; clinical symptoms, treatment and relevant lab results.

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TEST:	Echovirus Culture
Synonym:	Echovirus culture: Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934

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TEST:	Ehrlichia serology
Synonym:	Human Monocytic Ehrlichiosis (HME)
	Human Granulocytic Anaplasmosis (HGA)
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
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Collect:	Red-top vacutainer tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -
	20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	NEGATIVE—Titer < 1:80
	<b>POSITIVE</b> —Titer > 1:320 probable recent infection
	INDETERMINATE—Titer >1:80 but <1:320, possible early infection/past exposure with
	falling titers or cross-reactivity with related organism
Additional Information:	
Purpose of Test:	Detect antibodies to HME & HGA
Method:	IFA
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the
	assessment of a patient's health. Clinical correlation is required.
	Cross reaction between E. chaffeensis, E. canis & E. ewingii by IFA can occur.

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TEST:	Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins–producing <i>E. coli</i> )
Synonym:	Stool culture for enteric pathogens; enteric pathogens; stool culture and sensitivity; feces
	culture.
Laboratory/Phone:	Microbiology - Enterics 443-681-4570
Turnaround Time:	Usually four (4) days to several weeks [from specimen receipt in the Laboratory].
Specimen Required:	Stool in stool culture transport media (Para Pak for Enteric pathogens [orange cap]).
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be
	properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	1-2 grams fresh stool; 5-10 ml if liquid
Specimen Volume (Minimum):	Rectal swab (less effective than stool specimen).
	NOTE: Campylobacter cannot be tested for on specimens submitted on a rectal swab.
Collect:	Fresh stool in Para Pak for enteric pathogens (Cary-Blair transport media), select portion of
	stool containing pus, blood or mucous; rectal swab inserted one (1) inch beyond anal
	sphincter, rotate carefully, withdraw and place in Cary-Blair transport medium.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)-
	Check Enteric Routine culture
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Orange top Para-Pak Transport Media: store and ship refrigerated (2-8°C) temperature.
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Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> <li>Specimen received after prolonged delay (usually more than 96 hours)</li> <li>Dry specimen</li> <li>Specimen contaminated with urine or water</li> <li>Stool containing barium</li> <li>Insufficient quantity</li> </ul>
Availability:	Specimen frozen Monday through Friday
Results and Interpretation:	Identification of pathogenic enteric organisms and determination of antimicrobial susceptibilities, if clinically appropriate.
Reference Range:	Normal stool flora
Additional Information:	Enteric culture screens routinely for Salmonella, Shigella, Campylobacter, and Shiga toxin – producing <i>E. coli</i> . Yersinia culture and Vibrio culture must be specifically indicated as they are not part of routine testing. Same transport media will support the growth and detection of these organisms. Collect specimens early in the course of enteric disease and prior to antimicrobial therapy. Collect 2 or 3 stools on separate days to increase the likelihood of isolating enteric pathogens. DO NOT COLLECT SPECIMEN FROM THE TOILET. AVOID CONTAMINATION WITH URINE.
Purpose of Test:	Isolation, identification and if clinically appropriate, antimicrobial susceptibilities of potentially pathogenic organisms.
Method:	Culture on selective media, staining, biochemical testing, antimicrobial susceptibility testing; EIA (Enzyme Immuno Assay) for <i>E. coli</i> O157.
Interfering Substances/Limitations:	Administration of antibiotics, barium
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Enterohemorrhagic Escherichia coli (EHEC)
Synonym:	E. coli O157 typing; Isolate for E. coli O157 serotyping (referral isolate): Refer to
	instructions for <i>E. coli</i> O157 typing.
Laboratory/Phone:	Microbiology-Enterics 443-681-3952

TEST:	Enteroinvasive Escherichia coli (EIEC)
Synonym:	E. coli O157 typing; Isolate for E. coli O157 serotyping (referral isolate): Refer to
	instructions for <i>E. coli</i> O157 typing.
Laboratory/Phone:	Microbiology-Enterics 443-681-3952

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TEST:	Enterovirus Culture
Synonym:	Enterovirus (including Echovirus, Coxsackie, and Polio): Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934

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Epstein Barr Virus (EBV) serology
EBV, Epstein Barr Virus
443-681-3938/3931
5 business days
Serum
The specimen/sample must be properly labeled and include patient's name or unique
patient/sample identifier matching the test requisition or electronic test order.
2 ml. (Whole Blood)
1 ml. (Whole Blood)
Red-top vacutainer tube
DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
Indicate specimen type using the "Specimen Code" on form.
Specimens must be packaged in a triple packaging system to ensure that under normal
conditions of transport they cannot break, be punctured or leak their contents (Refer to
pages 9 & 10 for triple packing guidance).
*Refer to current Federal regulations for specific shipping requirements.
Ambient temperature for specimens on the blood clot (whole blood specimens
transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
20°C (frozen).
Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being
frozen
Monday through Friday
<b>POSITIVE</b> —Antibodies detected (EBNA-1(Epstein Barr Nuclear Antigen) denotes previous
infection, VCA (Viral Capsid Antigen)-IgM denotes current or reactivated infection, VCA-
IgG denotes current or previous infection, when EA (Early Antigen) & VCA-IgG positive may
denote chronic or recurrent illness.)
<b>NEGATIVE</b> —Antibodies not detected (EBNA-1, EA, VCA-IgG, presume susceptible to primary infection, VCA IgM presume no active infection)
<b>EQUIVOCAL</b> —Immunological status cannot be determined. Please resubmit another
specimen in 1-3 weeks.
Detect antibodies to EBV
EIA
Icteric, hemolyzed, lipemic or heat inactivation specimen
DHMH Laboratories Administration, Central Laboratory
1770 Ashland Avenue, Baltimore, Maryland 21205
This test aid in the diagnosis of Infectious mononucleosis. Serologic results should not be
used as a sole means for diagnosis, treatment, or for the assessment of a patient's health.
Clinical correlation is required.

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TEST:	Filariasis serology (CDC Referral)
Synonym:	Wuchereria bancrofti, Brugia malayi, Bancroftian filariasis
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	18 business days (CDC Referral)
Specimen Required:	Serum; plasma
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	0.5 ml. (Whole Blood)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
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Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or - 20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information: Purpose of Test:	Detect antibodies to filaria
Methods:	EIA, ELISA, Antibody Detection
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Processing Site for CDC referral:	DHMH. Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen submission. Required supplemental information: Exposure and travel history, include other relevant risk factors; clinical symptoms, treatment and relevant lab results.

TEST:	Foodborne Pathogens (Bacillus cereus, Clostridium perfringens, Staph aureus)
Synonym:	Foodborne Pathogenic Microorganisms, Stool Culture for Foodborne Pathogens
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	3 - 5 days [from specimen receipt in the Laboratory]
Specimen Required:	Stool, unpreserved
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must be
	properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	4 gm
Specimen Volume (Minimum):	N/A
Collect:	Fresh, unpreserved stool in a sterile screw-top jar. Submit within 48 hours.
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ship on wet ice
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results
	and to avoid misleading information that might lead to misdiagnosis and inappropriate
	therapy. A request for a new specimen will provide appropriate materials and clinically
	relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
	<ul> <li>Stool in preservative</li> </ul>
	<ul> <li>Specimen received after prolonged delay (usually more than 72 hours)</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Staph. aureus: Any amount is significant and is reported as rare, few, moderate, or many.
	Bacillus cereus and Clostridium perfringens: colony count of > 100,000 CFU/ml is
	considered significant.
Reference Range:	(Staph aureus: Bacillus cereus: Clostridium perfringens) not found after 48 hours
	incubation.
	Continued Next Page>

Additional Information:	<b>Bacillus cereus:</b> The symptoms of <i>B. cereus</i> diarrheal type food poisoning mimic those of
	<i>Clostridium perfringens</i> food poisoning. The onset of watery diarrhea, abdominal cramps, and pain occurs 6-15 hours after consumption of contaminated food. Nausea may accompany diarrhea, but vomiting (emesis) rarely occurs. Symptoms persist for 24 hours in most instances. The emetic type of food poisoning is characterized by nausea and vomiting within 0.5 to 6 hours after consumption of contaminated foods. Occasionally, abdominal cramps and/or diarrhea may also occur. Duration of symptoms is generally less
	than 24 hours. <b>Clostridium perfringens:</b> The common form of <i>C. perfringens</i> poisoning is characterized by intense abdominal cramps and diarrhea which begin 8-22 hours after consumption of foods containing large numbers of those <i>C. perfringens</i> bacteria capable of producing the food poisoning toxin. The illness is usually over within 24 hours but less severe symptoms may persist in some individuals for 1 or 2 weeks.
	<b>Staph. aureus:</b> The onset of symptoms in staphylococcal food poisoning is usually rapid and in many cases acute, depending on individual susceptibility to the toxin, the amount of contaminated food eaten, the amount of toxin in the food ingested, and the general health of the victim. The most common symptoms are nausea, vomiting, retching, abdominal cramping, and prostration. Some individuals may not always demonstrate all the symptoms associated with the illness. In more severe cases, headache, muscle cramping, and transient changes in blood pressure and pulse rate may occur. Recovery generally takes two (2) days; however, it is not unusual for complete recovery to take three (3) days and sometimes longer in severe cases.
Purpose of Test:	To detect the presence of bacteria that may be agents of food poisoning, since the presence of any amount of <i>Staph aureus</i> or the presence of large amounts (greater than 100,000 CFU/mI) of <i>Bacillus cereus</i> or <i>Clostridium perfringens</i> is consistent with a potential hazard to health.
Method:	Culture, isolation and identification of <i>Bacillus cereus</i> , <i>Clostridium perfringens</i> or <i>Staph aureus</i> . Colony count performed on specimens for <i>Bacillus cereus</i> and <i>Clostridium perfringens</i> .
Interfering Substances:	Stool preservative
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Francis disease
Synonym:	Francisella tularensis; Pasteurella tularensis, tularemia, rabbit fever, deerfly fever, Ohara's
	disease, Francis disease: Refer to instructions for Francisella tularensis Culture or
	Francisella tularensis Antibody.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

TEST:	Francisella tularensis Culture
Synonym:	Pasteurella tularensis, tularemia, rabbit fever, deerfly fever, Ohara's disease, Francis
	disease
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952
Turnaround Time:	2 -7 days [from specimen receipt in the Laboratory]
Specimen Required:	1. Blood Cultures
	2. Tissue samples
	3. Tissue aspirates (Including lymph node and bone marrow)
	4. Isolate
	5. Respiratory Specimens: Sputum, BAL, or pleural fluid.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection.
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Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Specimen Volume (Minimum): Collect: Form: Packaging and Shipping*:	<ul> <li>N/A</li> <li>1. Blood Culture: Collect appropriate blood volume and number of sets per routine laboratory protocol.</li> <li>2. Tissues or scraping of an ulcer is preferable. A swab of the ulcer is an acceptable alternative. Collect in a sterile container. For small amount tissue samples, add several drops of sterile normal saline to keep the tissue moist.</li> <li>3. Swabs: Collect a firm sample of the advancing margin of the lesion. If using a swab transport carrier, the swab should be reinserted into the transport package and the swab fabric moistened with the transport medium inside the packet.</li> <li>4. Aspirate of involved tissue: Collect per routine laboratory protocol.</li> <li>5. Isolate: Pick a pure culture to a chocolate agar plate or slant.</li> <li>DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.</li> <li>Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 &amp; 10 for triple packing guidance).</li> </ul>
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions: Specimen Rejection Criteria:	<ol> <li>Blood Cultures: Transport directly to the Laboratory at room temperature.</li> <li>Tissues: Transport in a sterile container. For small sample, add several drops of sterile saline to keep the tissue moist. Transport immediately to the Laboratory at room temperature. If transport is delayed, keep specimen chilled at 2-8°C.</li> <li>Swabs: Transport to the Laboratory using transport carrier at 2-8°C. Room temperature is acceptable.</li> <li>Aspirates: Transport directly to the Laboratory at room temperature. If transporting is delayed keep specimen chilled at 2-8°C.</li> <li>Isolates: Transport the specimen at room temperature on a sealed chocolate agar plate or slant.</li> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for request dest</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport and patient on the specimen</li> </ol>
Availability:	24 hrs/day, 7 days/week
Results and Interpretation:	Francisella tularensis isolated/detected. Francisella tularensis not found.
Additional Information:	Call 410-925-3121 before sending specimen to the Laboratory.
Purpose of Test:	To confirm diagnosis of tularemia by culture.
Method:	LRN Protocols
Interfering Substances:	Isolate must be inoculated unto media that contains cystine (e.g., chocolate agar plate or slant).
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Francisella tularensis is highly infectious. <i>PLEASE</i> use a biological safety cabinet when working with specimens suspected of harboring <i>F. tularensis. Call 410-925-3121 before sending to the Laboratory.</i>

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TEST:	Francisella tularensis Serology (CDC Referral)
Synonym:	Tularemia antibody; rabbit fever, deerfly fever, Ohara's disease, Francis disease
Laboratory/Phone:	443-681-3839/3931
Turnaround Time:	2 weeks (CDC Referral)
Specimen Required:	Serum
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Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/tularemia/index.html
Purpose of Test:	Detect antibodies to F. tularensis
Method:	Microagglutination
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimer
	submission. Required supplemental information: Please include submitting agency,
	contact name, address, phone number, specimen identifier, patient name, specimen
	source and type, sex and date of birth, symptoms of onset, sample collection date, and
	clinical information including type and date of treatment patient has received.

TEST:	Genital culture (Bacterial Culture, Routine)
Synonym:	Aerobic culture, routine culture, genital culture: Refer to instructions for <b>Bacterial Culture, Routine</b> .
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:	Giardia (Ova and Parasites Microscopic Examination)
Synonym:	Giardia, Parasitic identification: Refer to instructions for <b>Ova and Parasites Microscopic</b> <b>Examination</b> .
Laboratory/Phone:	Microbiology 443-681-3952 or 443-681-4570

TEST:	Glanders (Burkholderia mallei)
Synonym:	Glanders; Burkholderia (formerly Pseudomonas) mallei: Refer to instructions for
	Burkholderia mallei and Burkholderia pseudomallei.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

TEST:	Gonorrhea Culture
Synonym:	GC Culture, Neisseria gonorrhoeae Culture
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	2-3 days – minimum [from specimen receipt in the Laboratory]
Specimen Required:	Cervical, rectal, throat, urethral, vaginal
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OHMH-Laboratories Administration	The J. Mehsen Joseph Public Health Laboratory
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. Don't use china markers – their marking smudges and rubs off when wet or use permanent marker. Label bottom of plate (not lid). [Lot number and expiration date must remain visible on media.] The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	GC culture plate streaked with Dacron <sup>™</sup> swab immediately after collection.
Specimen Volume (Minimum):	N/A
Collect:	Materials*: GC culture plate, Dacron <sup>™</sup> swab, CO <sub>2</sub> tablet, resealable plastic bag. Roll swab directly on the medium in a large "Z" (1a) (to provide adequate exposure of the swab to the medium for transfer of organisms.) Cross-streak immediately with a sterile loop (1b).
	*Please do not use damaged plates or less than optimal media.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on the form and number of hours incubated (if any).
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	DO NOT REFRIGERATE after specimen is collected. When packing plates for transport,
	keep them inverted and place in a suitable container that will protect them from extreme
	heat or cold.
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> </ul>
	L Billion des selence de Existence
Availability:	Monday through Friday
Results and Interpretation:	Neisseria gonorrhea isolated and identified. Antibiotic susceptibilities reported.

Additional Information:	Store unused plates under refrigeration upside down (media facing down). Discard any
	plate(s) with an expired expiration date or that exhibit growth prior to use (never use
	contaminated plates). Always allow plates to warm to room temperature before using
	(cold kills Neisseria gonorrhea). Use Dacron™ tipped swabs with plastic shafts (do not use
	cotton-tipped swabs, as they may contain fatty acids that can interfere with the survival of some organisms. Also do not use calcium alginate-tipped swabs. They can be toxic for
	some strains of <i>N. gonorrhoeae.</i> ) Always allow the surface of plates to dry before using (a wet surface hampers isolated colony formation). DO NOT CRUSH OR ADD WATER TO THE CO <sub>2</sub> GENERATING TABLET (CAUSES LOSS OF CO <sub>2</sub> AND POSSIBLE CONTAMINATION BY WATER.) MOISTURE FROM THE MEDIUM WILL ACTIVATE THE CO <sub>2</sub> TABLET. Do not incubate inoculated plates in the clinic longer than 24 hours (over-incubation leads to more growth of contaminating normal flora). If incubated, indicate the number of hours
	on the test request form. If an incubator is not available, invert the inoculated plates and hold them at room temperature until picked up by the courier. Do not refrigerate after inoculating. When packing plates for transport, keep them inverted and place in a suitable container that will protect them from extreme heat or cold.
Purpose of Test:	Isolation, identification and antibiotic susceptibility testing for Neisseria gonorrhea.
Method:	Culture
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
-	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

TEST:	Group A Strep Culture
Synonym:	Beta Strep culture, Streptococcus pyogenes culture, throat culture for Group A Strep
Lab/Phone:	Microbiology 443-681-3952
Turnaround Time:	1-2 days [from specimen receipt in the Laboratory]
Specimen Required:	Throat swab
Specimen identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	One (1) throat swab
Specimen Volume (Minimum):	N/A
Collect:	Culturette tube with transport medium
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store and ship at room temperature, ship as quickly as possible.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care. Unlabeled or improperly labeled specimen Non-sterile or leaking container Inappropriate specimen transport conditions Illegible, or no submitter information on the request form Mismatched form and specimen Broken specimen/sample container The wrong specimen for test request Inappropriate outfit for requested test Illegible or no patient information on the specimen Expired transport media
Availability:	Monday through Friday
Results and Interpretation:	Group A Strep isolated and identified
Reference Range:	No Group A Strep detected
Additional Information:	N/A
Purpose of Test:	Detect the presence of Group A Strep
Method:	Culture
Interfering Substances:	N/A
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Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	Group A streptococcus (ABCs (previously BIDS))
Synonym:	Active Bacterial Core Surveillance (ABCs) (Bacterial Invasive Disease Surveillance) Group A
	streptococcus: Refer to instructions for ABCs (previously BIDS).
Lab/Phone:	Microbiology 443-681-3952

TEST:	Group B Strep Screen
Synonym:	Prenatal screen for Group B Strep; Group B Strep culture; Genital Culture
Lab/Phone:	Microbiology 443-681-3952
Turnaround Time:	2-3 days [from specimen receipt in the Laboratory]
Specimen Required:	Vaginal/rectal swab
Specimen identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must b
	properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	One (1) vaginal/rectal swab
Specimen Volume (Minimum):	N/A
Collect:	Culturette tube with transport medium (Amies or Stuart's)
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
-	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store and ship at room temperature, ship as quickly as possible.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate result
	and to avoid misleading information that might lead to misdiagnosis and inappropriate
	therapy. A request for a new specimen will provide appropriate materials and clinically
	relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
	<ul> <li>Specimen received after prolonged delay (usually more than 72 hours)</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Group B Strep isolated and identified
Reference Range:	No Group B Strep detected
Additional Information:	Prenatal screening for Group B Strep at 35-37 weeks gestation. If patient is allergic to
	penicillin, add note to this effect and request antimicrobial susceptibility testing to
	clindamycin and erythromycin.
	Gardnerella vaginalis isolation done on request for routine genital cultures.
Purpose of Test:	Detect the presence of Group B Strep
Method:	Culture
Interfering Substances:	N/A
Testing Site:	DHMH Laboratories Administration, Central Laboratory
i coung one.	1770 Ashland Avenue, Baltimore, Maryland 21205
Commont:	
Comment:	N/A

TEST:	Group B Streptococcus (ABCs (previously BIDS))
Synonym:	Active Bacterial Core Surveillance (ABCs) (Bacterial Invasive Disease Surveillance) Group B
	Streptococcus: Refer to instructions for ABCs (previously BIDS).
Lab/Phone:	Microbiology 443-681-3952

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TEST:	Haemophilus ducreyi Culture
Synonym:	Chancroid Culture; Haemophilus ducreyi culture
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	Seven (7) days [from specimen receipt in the Laboratory]:
Specimen Required:	Ulcer scrapings
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must be
	properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	Collect prior to antimicrobial treatment.
	Clean the surface of the lesion with 0.85% NaCl. If there is a crust on the lesion remove it.
	Moisten swab with saline and collect specimen by vigorously rubbing the base of the
	lesion, put the swab in Amies transport medium or scrape the base of the ulcer with a
	sterile scalpel blade, irrigate with sterile saline. Then rub the base vigorously with a sterile
	swab and put it in Amies transport medium or aspirate fluid with a flamed smoothed
	Pasteur pipette or needle and syringe, put it in sterile container.
	For abscess disinfect skin with alcohol and iodine. Aspirate fluid with a needle and syringe
	and put it in a sterile container. NOTE: Intact bubo aspirates are rarely positive for the
	organisms unless they have ruptured.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	After collection, place specimen immediately on ice or in the refrigerator and transport on
	ice to the laboratory.
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results
	and to avoid misleading information that might lead to misdiagnosis and inappropriate
	therapy. A request for a new specimen will provide appropriate materials and clinically
	relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and encines</li> </ul>
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	<ul> <li>Broken specimen/sample container</li> <li>The unexpected provides the test request</li> </ul>
	<ul> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	<ul> <li>Expired transport media</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Positive Culture: Haemophilus ducreyi present. A positive culture indicates infection in a
Results and interpretation.	patient with an ulcerative lesion. Mixed infections with other agents known to cause
	ulcerative sexually transmitted diseases are not uncommon. The presence of
	Haemophilus ducreyi does not rule out these other infections which should be considered
	in the evaluation of the patient.
Reference Range:	Haemophilus ducreyi not found
Additional Information:	False-Negative cultures can result from prior antimicrobial therapy, strain growth
	variability, and sample and transport techniques
Purpose of Test:	Diagnosis of chancroids
Method:	Culture
Interfering Substances:	Prior antimicrobial therapy
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Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

TEST:	Haemophilus influenzae (ABCs (previously BIDS))
Synonym:	Active Bacterial Core Surveillance (ABCs) (Bacterial Invasive Disease Surveillance) Haemophilus influenzae: Refer to instructions for ABCs (previously BIDS).
Laboratory/Phone:	Microbiology 443-681-3952

TEST:	Hantavirus serology (CDC Referral)
Synonym:	Hanta, HPS, HFRS, Hantaan
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	10 business days (CDC Referral)
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or - 20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/hantavirus/index.html
Purpose of Test:	Detect IgG & IgM antibody to the SNV
Method:	ELISA
Interfering Substances:	None
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen submission. Required supplemental information: Exposure and travel history, include other relevant risk factors; clinical symptoms, treatment and relevant lab results.
	Required supplemental form at: http://www.cdc.gov/ncezid/dhcpp/vspb/pdf/specimen-submission.pdf

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TEST:	Helminths
Synonym:	Helminths are worm-like parasites that include the flukes (Trematodes); tapeworms
	(Cestodes); and roundworms (Nematodes): Refer to instructions for Ova and Parasites
	Microscopic Examination.
Lab/Phone:	Microbiology 443-681-3952

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TEST:	Hepatitis A IgM Antibody (Hepatitis A Screen)
Synonym:	Hepatitis A IgM Antibody, HAV IgM, HAVAB-M.
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum; plasma
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time o
	collection. The specimen/sample must be properly labeled and match the test requisition
	or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum or Plasma)
Specimen Volume Minimum:	3 ml. (Whole blood) or 2 ml. (Serum or Plasma)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
Form.	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
Transport Conditions.	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic;
Specimen Rejection Criteria:	
A veile bilite v	gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Monday to Friday. Call laboratory for prior approval.
Results and Interpretation:	Assay results should be interpreted only in the context of other clinical laboratory findings
	and the total clinical status of the individual. It has been shown that a viremic window
	exists with individuals infected with HAV, where the individual may be symptomatic for
	hepatitis but IgM anti-HAV nonreactive.
	Negative: IgM anti-HAV not detected. Does not exclude the possibility of exposure to or
	infection with HAV. Levels of IgM anti-HAV may be below the cut-off in early infection.
	Equivocal/Gray zone: HAV IgM antibody may or may not be present. Patients exhibiting
	gray zone test results should be closely monitored by redrawing and retesting
	approximately one week intervals. Monitoring the level of IgM anti-HAV by redrawing and
	retesting at approximately one week intervals will distinguish rapidly rising IgM anti-HAV
	levels associated with early acute hepatitis A infection from gradually decreasing or
	unchanging IgM anti-HAV levels often associated with late acute stage of HAV infection.
	Positive: HAV IgM antibody detected. Presumptive evidence of HAV infection. A reactive
	IgM anti-HAV result does not rule out other hepatitis infections.
Additional Information:	For more information, see the CDC link at: <a href="http://www.cdc.gov/hepatitis/index.htm">http://www.cdc.gov/hepatitis/index.htm</a>
Purpose of Test:	HAVAB-M assay is for the qualitative detection of IgM antibody to hepatitis A virus (IgM
	anti-HAV) in human serum or plasma. IgM anti-HAV is indicated for testing of specimens
	from individuals who have signs and symptoms consistent with acute hepatitis. Test
	results are used in conjunction with other laboratory results and clinical information as an
	aid in the diagnosis of acute or recent hepatitis A viral infection. During the acute phase of
	HAV infection, IgM anti-HAV appears in the patient's serum and is nearly always
	detectable at the onset of symptoms. In most cases, IgM anti-HAV response peaks within
	the first month of illness and can persist for up to six months. It is not intended for use in
	the mist month of miless and can persist for up to six months. It is not interfued for use in
Method:	screening blood, plasma, or tissue donors.
	screening blood, plasma, or tissue donors. Chemiluminescent micro particle immunoassay (CMIA)
Method: Interfering Substances:	screening blood, plasma, or tissue donors.         Chemiluminescent micro particle immunoassay (CMIA)         May not detect a recent infection, or infection in a person with severely compromised
Interfering Substances:	screening blood, plasma, or tissue donors.         Chemiluminescent micro particle immunoassay (CMIA)         May not detect a recent infection, or infection in a person with severely compromised immune system.
	screening blood, plasma, or tissue donors.         Chemiluminescent micro particle immunoassay (CMIA)         May not detect a recent infection, or infection in a person with severely compromised

Comment:	LIMITATIONS: Specimens from individuals who have received preparations of mouse
	monoclonal antibodies for diagnosis or therapy may contain human anti-mouse antibodies
	(HAMA). Such specimens may show either falsely elevated or depressed values when
	tested with assay kits (such as HA; HAVAB-M) reagents contain a component that reduces
	the effect of HAMA reactive specimens. Additional clinical or diagnostic information may
	be required to determine patient status. A reactive IgM anti-HAV result does not should
	be used and interpreted only in the context of the overall clinical picture. A negative test
	result does not exclude the possibility of exposure to hepatitis A virus. Levels of IgM anti-
	HAV may be below the cut-off in early infection and late acute infection. Heterophilic
	antibodies in human serum can react with reagent immunoglobulins, interfering with in
	vitro immunoassays. Patients routinely exposed to animals or to animal serum products
	can be prone to this interference and anomalous values may be observed. Additional
	information may be required for diagnosis. Specimens from individuals with Non-
	Hodgkin's Lymphoma may cross-react with this assay.

TEST:	Hepatitis A IgG Antibody.
Synonym:	HAV IgG, HAVAB-G
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum; Plasma
Specimen identification:	Label container with patient's last name, first Name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum or Plasma)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum or Plasma)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or 20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic; gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Service available only to state and local health departments Monday to Friday.
Results and Interpretation:	Negative: No detectable IgG antibody to hepatitis A virus.           Positive: Presence of detectable IgG antibody to HAV. It indicates past HAV infection or confers immunity by HAV vaccination.
Additional Information:	For more information, see the CDC link at: <u>http://www.cdc.gov/hepatitis/index.htm</u>
Purpose of Test:	HAVAB-G assay is for the qualitative detection of IgG antibody to hepatitis A virus (IgG anti-HAV) in human serum or plasma. Detectable levels above the assay cut-off suggest immunity to HAV infections.
Method:	Chemiluminescent micro particle immunoassay (CMIA)
Interfering Substances:	May not detect a recent infection, or infection in a person with severely compromised immune system.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
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Comment:	LIMITATIONS: If the Architect HAVAB-G results are inconsistent with clinical evidence,
	additional testing is suggested to confirm the results. Specimens from patients who have
	received preparations of mouse monoclonal antibodies for diagnosis or therapy may
	contain human anti-mouse antibodies (HAMA). Such specimens may show either falsely
	elevated or depressed values when tested with assay kits (such as ARCHITECT HAVAB-G)
	that employ mouse monoclonal antibodies. Specimens from individuals with anti-E.coli,
	anti-CMV, or hemodialysis patients may cross react with this assay. Heterophilic
	antibodies in human serum can react with reagent immunoglobulins, interfering with in
	vitro immunoassays. Patients routinely exposed to animals or to animal serum products
	can be prone to this interference and anomalous values may be observed. Additional
	information may be required for diagnosis. Specimens containing low antibody
	concentrations (near the cutoff) assayed after a freeze/thaw may exhibit elevated values
	that may be false positives.

TEST:	Hepatitis B Core Antibody IgM (Hepatitis B surface antigen Positive reflex test)
Synonym:	HBc IgM Ab; anti-HBc IgM, CORE-M
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum; plasma
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum or Plasma)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum or Plasma)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic; gross bacterial contamination.
Availability:	Monday through Friday.
Results and Interpretation:	<ul> <li>Negative: IgM anti-HBc not detected. Does not exclude the possibility of exposure to or infection with HBV.</li> <li>Equivocal/Gray zone: IgM anti-HBc may or may not be present. Patients with specimens exhibiting grayzone test results should be retested at approximately one-week intervals. Monitoring the level of IgM anti-HBc by retesting at approximately one week intervals will distinguish rapidly rising IgM anti-HBc levels associated with early acute hepatitis B infection from gradually decreasing or unchanging IgM anti-HBc levels often associated with late acute stage of HBV infection, six to nine months from the appearance of HBsAg.</li> <li>Positive: Presumptive evidence of IgM anti-HBc antibodies.</li> </ul>
Additional Information:	For more information, see the CDC link at: <u>http://www.cdc.gov/hepatitis/index.htm</u>
Purpose of Test:	The CORE-M assay is for the qualitative detection of IgM antibody to hepatitis B core antigen in human serum or plasma. A test for IgM anti-HBc is indicated as an aid in the diagnosis of acute or recent hepatitis B virus (HBV) infection in conjunction with other laboratory results and clinical information. It is not intended for use in screening blood, plasma, or tissue donors.
Method:	Chemiluminescent micro particle immunoassay (CMIA)
Interfering Substances:	May not detect a recent infection, or infection in a person with severely compromised immune system.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
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Comment:	Limitations: Current methods for the detection of IgM anti-HBc may not detect all
	infected individuals. A non-reactive test result does not exclude the possibility of
	exposure to or infection with HBV. CORE-M assay is limited to the detection of IgM anti-
	HBc in human serum or plasma. It can be used to determine whether a patient has, or
	has recently had, acute or subclinical hepatitis B infection. Supportive clinical
	information, including other hepatitis B markers, should also be evaluated. The test
	cannot determine a patient's immune status to hepatitis B. Specimens from patients
	with high levels of IgM (e.g., specimens from patients with multiple myeloma) may show
	depressed values when tested with assay kits (such as CORE-M) that use reagents
	containing anti-human IgM. Specimens from patients who have received preparations of
	mouse monoclonal antibodies for diagnosis or therapy may contain human anti-mouse
	antibodies (HAMA). Such specimens may show either falsely elevated or depressed
	values when tested with assay kits (such as CORE-M) that employ mouse monoclonal
	antibodies. Heterophilic antibodies in human serum can react with reagent
	immunoglobulins, interfering with in vitro immunoassays. Patients routinely exposed to
	animals or to animal serum products can be prone to this interference and anomalous
	values may be observed. Additional information may be required for diagnosis.

TEST: **Hepatitis B Core Antibody Total** HBclgMAb Synonym: Laboratory/Phone: Vaccine Preventable Disease/443-681-3889 Turnaround Time: 2-5 business days Specimen Required: Serum; plasma Label container with patient's last name, first name, DOB, specimen type, date and time Specimen identification: of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order. Specimen Volume (Optimum): 5 ml. (Whole blood) or 4 ml. (Serum or Plasma) Specimen Volume (Minimum): 3 ml. (Whole blood) or 2 ml. (Serum or Plasma) Collect: Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma) Form: DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form. Packaging and Shipping\*: Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). \*Refer to current Federal regulations for specific shipping requirements. Transport Conditions: Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen). Specimen Rejection Criteria: Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic; gross bacterial contamination. Availability: Monday through Friday. **Results and Interpretation:** Negative: Hepatitis B core antibodies not detected. Positive: Hepatitis B core antibodies were detected. The presence of anti-HBc antibodies does not differentiate between acute or chronic hepatitis B infections. Additional Information: For more information, see the CDC link at: http://www.cdc.gov/hepatitis/index.htm Purpose of Test: The CORE assay is for the qualitative detection of antibodies to hepatitis B core antigen in human serum or plasma. It is intended as an aid in the diagnosis of acute, chronic, or resolved hepatitis B virus (HBV) infection in conjunction with other laboratory results and clinical information. It is not intended for use in screening blood, plasma, or tissue donors. Method: Chemiluminescent microparticle immunoassay (CMIA) **Interfering Substances:** May not detect a recent infection, or infection in a person with severely compromised immune system. Testing Site: DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205 Continued Next Page>

Comment:	<b>LIMITATIONS:</b> A nonreactive test result does not exclude the possibility of exposure to or infection with HBV. Specimens from patients who have received preparations of mouse
	monoclonal antibodies for diagnosis or therapy may contain human anti-mouse
	antibodies (HAMA). Such specimens may show either falsely elevated or depressed
	values when tested with assay kits (such as CORE) that employ mouse monoclonal
	antibodies. Heterophilic antibodies in human serum can react with reagent
	immunoglobulins, interfering with in vitro immunoassays. Patients routinely exposed to
	animals or to animal serum products can be prone to this interference and anomalous
	values may be observed. Additional information may be required for diagnosis.

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TEST:	Hepatitis B Suface Antibody (Hepatitis B Panel, Hepatitis B post vaccine)
Synonym:	HBsAb, Hepatitis B surface Antibody, AUSAB.
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum; plasma
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time
	of collection. The specimen/sample must be properly labeled and match the test
	requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum or Plasma)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum or Plasma)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	-20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic;
	gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Monday through Friday.
Results and Interpretation:	<b>Negative:</b> < 8.00 mIU/mL Individual is considered not immune to HBV infection.
	<b>Equivocal/Grayzone:</b> ≥ 8.00 mIU/mL to < 12.00 mIU/mL The immune status of the
	individual should be further assessed by considering other factors, such as clinical status
	follow-up testing, associated risk factors, and the use of additional diagnostic
	information.
Defense Dense	<b>Positive:</b> ≥12.00 mIU/mL. Individual is considered immune to HBV infection.
Reference Range: Additional Information:	Patient's with a titer ≥12.00 mIU/mL is considered immune to Hepatitis B Virus infection
	For more information, see the CDC link at: <u>http://www.cdc.gov/hepatitis/index.htm</u>
Purpose of Test:	AUSAB assay is for the quantitative determination of antibody to hepatitis B surface
	antigen in human serum or plasma. It is intended for measurement of antibody response
	following hepatitis B virus (HBV) vaccination, determination of HBV immune status, and
	for the laboratory diagnosis of HBV disease associated with HBV test results and clinical
Mathad	information. It is not intended for use in screening blood, plasma, or tissue donors.
Method:	Chemiluminescent microparticle immunoassay (CMIA)
Interfering Substances:	May not detect a recent infection, or infection in a person with severely compromised
	immune system. DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Testing Site:	1770 Asmanu Avenue, Baltinore, Iviaryianu 21205

LIMITATIONS: For diagnostic purposes, results should be used in conjunction with
patient history and other hepatitis markers for diagnosis of acute and chronic infection.
A non-reactive test result does not exclude the possibility of exposure to hepatitis B
virus. Results obtained with the AUSAB assay may not be used interchangeably with
values obtained with different manufacturers' assay methods. Assay does not
differentiate between vaccines and natural infection. Results from immune suppressed
patients should be interpreted with caution. Performance characteristics have not been
established for therapeutic monitoring. A reactive anti-HBs result does not exclude co-
infection by another hepatitis virus.

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TEST:	Hepatitis B Surface Antigen (Hepatitis B Panel; Hepatitis B Screen)
Synonym:	HBsAg, Hepatitis B surface Antigen Qualitative; HBsAg Qual.
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum; plasma
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum or Plasma)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum or Plasma)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic; gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Monday through Friday.
Results and Interpretation:	<b>Negative:</b> HBsAg not detected. <b>Positive:</b> Presumptive evidence of HBsAg.
Additional Information:	For more information, see the CDC link at: <u>http://www.cdc.gov</u>
Purpose of Test:	HBsAg Qualitative assay is for the qualitative detection of hepatitis B surface antigen in human serum or plasma. The assay may also be used to screen for HBV infection in pregnant women to identify neonates who are at risk for acquiring hepatitis B during the perinatal period. Assay results in conjunction with other laboratory results and clinical information, may be used to provide presumptive evidence of infection with the hepatitis B virus (HBV) (state of infection or associated disease not determined) in persons with signs and symptoms of hepatitis and in persons at risk for hepatitis B infection. Not intended for use in screening blood, plasma, or tissue donors.
Method:	Chemiluminescent microparticle immunoassay (CMIA)
Interfering Substances:	May not detect a recent infection, or infection in a person with severely compromised immune system.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
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Comment:	LIMITATIONS: The effectiveness of the HBsAg Qualitative assay for use in screening
comment.	blood, plasma, or tissue donors has not been established. Assay performance
	characteristics have not been established when the HBsAg Qualitative assay is used in
	conjunction with other manufacturers' assays for specific HBV markers. Current methods
	for the detection of hepatitis B surface antigen may not detect all potentially infected
	individuals. A nonreactive test result does not exclude the possibility of exposure to or
	infection with hepatitis B virus. A nonreactive test result in individuals with prior
	exposure to hepatitis B may be due to antigen levels below the detection limit of this
	assay or lack of antigen reactivity to the antibodies in this assay. If the HBsAg Qualitative
	results are inconsistent with clinical evidence, additional testing is suggested to confirm
	the result for diagnostic purposes. Results should be used in conjunction with patient
	history and other hepatitis markers for diagnosis of acute and chronic infection.
	Heterophilic antibodies in human serum can react with reagent immunoglobulins,
	interfering with in vitro immunoassays. Patients routinely exposed to animals or to
	animal serum products can be prone to this interference and anomalous results may be
	observed. Additional information may be required for diagnosis. Specimens from
	patients who have received preparations of mouse monoclonal antibodies for diagnosis
	or therapy may contain human anti-mouse antibodies (HAMA). Specimens containing
	HAMA may produce anomalous values when tested with assay kits such as HBsAg
	Qualitative that employ mouse monoclonal antibodies. A reactive HBsAg result does not
	exclude co-infection by another hepatitis virus.

TEST:	Hepatitis B Surface Antigen Confirmation (Hepatitis B Positive Reflex Test)
Synonym:	HBsAg Qualitative Conformation; HBsAg Qualitative Neutralization assay
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum, plasma
Specimen identification:	Label container with patient's last name, first Name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum or Plasma)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum or Plasma)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	Test cannot be requested, it is a reflex test for Hepatitis B surface antigen Positive
	specimens.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or - 20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic;
	gross bacterial contamination. Specimens collected >7 days prior to arrival without being
	frozen.
Availability:	Monday through Friday
Results and Interpretation:	<b>Confirmed:</b> Presence of HBs Antigen confirmed. Confirmed result may indicate acute or
	chronic HBV infection, depending on presence of other HBV serological markers.
	<b>Not Confirmed:</b> Indicates the presence of HBsAg cannot be confirmed via neutralization.
	The repeatedly reactive result obtained with the HBsAg Qualitative assay may be the
	result of a nonspecific reaction (false reactive). As the presence of nonspecific binding may
	obscure low levels of HBsAg in the specimen due to early infection or early recovery, it is
	recommended that the patient be evaluated for other serologic markers of HBV infection
	( <i>i.e.</i> , total anti-HBc or IgM antiHBc) and that the patient be retested for HBsAg in 4 to 6
	weeks.

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Additional Information:	For more information, see the CDC link at: <a href="http://www.cdc.gov/hepatitis/index.htm">http://www.cdc.gov/hepatitis/index.htm</a>
Purpose of Test:	The HBsAg Qualitative confirmation assay is for the qualitative confirmation of the
	presence of hepatitis B surface antigen (HBsAg) in human serum or plasma by specific
	antibody neutralization. Assay results, in conjunction with other laboratory results and
	clinical information, may be used to provide presumptive evidence of infection with the
	hepatitis B virus (HBV) (state of infection or associated disease not determined) in persons
	with signs and symptoms of hepatitis and in persons at risk for hepatitis B infection. It is
	not intended for use in screening blood, plasma, or tissue donors.
Method:	Chemiluminescent microparticle immunoassay (CMIA)
Interfering Substances:	May not detect a recent infection, or infection in a person with severely compromised
	immune system.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Limitations: The effectiveness of HBsAg Qualitative Confirmatory assay for use in
	screening blood, plasma, or tissue donors has not been established. Assay performance
	characteristics have not been established when HBsAg Qualitative Confirmatory assay is
	used in conjunction with other manufacturers' assays for specific HBV serological markers.
	If HBsAg Qualitative Confirmatory results are Inconsistent with clinical evidence,
	additional testing is suggested to confirm the result. For diagnostic purposes, results
	should be used in conjunction with patient history and other hepatitis markers for
	diagnosis of acute and
	Chronic infection. Heterophilic antibodies in human serum can react with reagent
	immunoglobulins, interfering with in vitro immunoassays. Patients routinely exposed to
	animals or to animal serum products can be prone to this interference and anomalous
	results may be observed. Additional information may be required for diagnosis.
	Specimens from patients who have received preparations of mouse monoclonal
	antibodies for diagnosis or therapy may contain human anti-mouse antibodies (HAMA).
	Specimens containing HAMA may produce anomalous values when tested with assay kits
	such as HBsAg Qualitative Confirmatory that employ mouse monoclonal antibodies.
	Although there is an association between the presence of HBsAg infectivity and a reactive
	result, it is recognized that presently available methods for HBsAg confirmation may not
	detect all possible cases of HBV infection.

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Hepatitis C Antibody (Hepatitis C Screen)
HCV Ab; anti-HCV; Hepatitis C Screen
Vaccine Preventable Disease/443-681-3889
2-5 business days
Serum; plasma
Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
5 ml. (Whole blood) or 4 ml. (Serum or Plasma)
3 ml. (Whole blood) or 2 ml. (Serum or Plasma)
Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
Indicate specimen type using the "Specimen Code" on form next to Hepatitis C Screen.
Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).

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Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled; hemolytic; gross
	bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Monday through Friday.
Results and Interpretation:	<b>Negative:</b> Antibodies to HCV not detected; does not exclude the possibility of exposure HCV.
	Equivocal/Grayzone: Antibodies to HCV may or may not be present; another specimen
	should be obtained from the individual for further testing or follow CDC recommendatio
	for supplemental testing.
	<b>Positive:</b> Presumptive evidence of antibodies to HCV; follow CDC recommendations for
	supplemental testing.
Additional Information:	For more information, see the CDC link at: <u>http://www.cdc.gov/hepatitis/index.htm</u>
Purpose of Test:	Anti-HCV assay is for the qualitative detection of antibody to Hepatitis C Virus in human
	serum or plasma. Assay results, in conjunction with other laboratory results and clinical
	information, may be used to provide presumptive evidence of infection with HCV (state
	of infection or associated disease not determined) in persons with signs and symptoms
	of hepatitis and in persons at risk for hepatitis C infection. It is not intended for use in
	screening blood, plasma, or tissue donors.
Method:	Chemiluminescent microparticle immunoassay (CMIA)
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
-	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: For diagnostic purposes, results should be used in conjunction with
	patient history and other hepatitis markers for diagnosis of acute and chronic infection.
	Current methods for the detection of antibodies to HCV may not detect all infected
	individuals. A nonreactive test result does not exclude the possibility of exposure to HCV
	Nonreactive test results in individuals with prior exposure to HCV may be due to
	antibody levels being below the detection limit of this assay or to lack of antibody
	reactivity to the recombinant antigens used in this assay. Immunocompromised patients
	who have HCV may produce levels of antibody below the sensitivity of this assay and
	may not be detected as positive. The affinity or avidity differences of anti-human
	IgG/IgM for anti-HCV have not been determined with this assay. Therefore, there may
	not be a demonstration of a significant increase in antibody level between acute and
	convalescent specimens for a patient in the late acute stage of infection when IgM
	antibodies are decreasing. Results obtained with Anti-HCV assay may not be used
	interchangeably with values obtained with different manufacturers' assay methods.
	Assay performance characteristics have not been established for newborns, infants,
	children, or populations of immunocompromised or immunosuppressed patients.
	Heterophilic antibodies in human serum can react with reagent immunoglobulins,
	interfering with in vitro immunoassays. Patients routinely exposed to animals or to
	animal serum products can be prone to this interference and anomalous values may be
	observed. Additional information may be required for diagnosis. A reactive anti-HCV
	result does not exclude co-infection by another hepatitis virus. The magnitude of an
	Anti-HCV assay result cannot be correlated to an end point titer.

TEST:	Herpes Simplex Virus (HSV Types 1 & 2) Virus Culture	
Synonym:	Herpes Simplex Virus (HSV Types 1 & 2): Refer to instructions for Virus Culture.	
Laboratory/Phone:	Virology: 443-681-3934	

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TEST:	Herpes Simplex Virus Serology	
Synonym:	Herpes simplex virus (HSV) type 1 & 2 IgG serology	
Laboratory/Phone:	443-681-3938/3931	
Turnaround Time:	5 business days	
Specimen Required:	Serum	
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique	
	patient/sample identifier matching the test requisition or electronic test order.	
Specimen Volume (Optimum):	2 ml. (Whole Blood)	
Specimen Volume (Minimum):	1 ml. (Whole Blood)	
Collect:	Red-top vacutainer	
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Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	-20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	<b>POSITIVE</b> —Presumptive evidence of IgG antibodies to HSV-1/HSV-2
	NEGATIVE—No IgG antibodies to HSV-1/HSV-2 detected
	EQUIVOCAL—Immunological status cannot be determined, please re-draw patient in 4-
	12 weeks.
Additional Information:	The performance of this assay has not been established for use in a pediatric population
	or for neonatal screening.
Purpose of Test:	Detect IgG antibodies to HSV I and HSV II
Method:	EIA
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the
	assessment of a patient's health. Clinical correlation is required.

TEST:	HIV-1 p24 Antigen and HIV-1/HIV-2 Antibody Combination Assay
Synonym:	HIV Ag/Ab Combo Assay
Laboratory/Phone:	443-681-3877
Turnaround Time:	3-7 working days
Specimen Required:	Serum from whole blood
Specimen identification:	Label container with patient's name and date of collection. (CTR# if applicable)
Specimen Volume (Optimum):	7 ml (Whole Blood)
Specimen Volume (Minimum):	5 ml (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	2-8°C (cold packs) DO NOT EXCEED STORAGE TIME LIMITATIONS
Specimen Rejection Criteria:	Must comply with proper labeling, storage, and transport requirements.
Availability:	Testing is performed routinely
Results and Interpretation:	Non-reactive = HIV-1 p24 antigen and HIV-1/HIV-2 antibodies not detected Reactive = Presumptive evidence of HIV-1 p24 antigen and/or HIV-1/HIV-2 antibodies; perform confirmatory/ supplemental assays
Reference Range:	Signal to cutoff (S/CO) values $\geq$ 1.00 are presumptive reactive for HIV-1 p24 antigen or HIV-1/ HIV-2 antibodies.
Additional Information:	Confirmatory assays may be performed to confirm presence of HIV antibody or HIV-1 RNA; Supplemental assay may be performed to differentiate HIV-1 and HIV-2 infections.
Purpose of Test:	Aid in the diagnosis of HIV-1 / HIV-2 infection including primary or acute HIV-1 infection.
Method:	Chemiluminescence microparticle immunoassay (CMIA)
Interfering Substances:	Fibrin, red blood cells, or other particulate matter
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Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Store at room temperature no more than 3 days, or 7 days if stored 2-8°C following specimen collection.

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TEST:	Infectious Mononucleosis (IM Serology)
Synonym:	Heterophile Antibody Assay
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	POSITIVE—Infectious Mono heterophile antibody detected NEGATIVE—Infectious Mono heterophile antibody not detected
Additional Information:	Further EBV testing can aid in the clinical diagnosis
Purpose of Test:	Detect antibody in patients with infectious mononucleosis
Method:	Slide agglutination
Interfering Substances:	Specimens showing gross hemolysis, lipemia, turbidity, or contamination can cause false positive reactions and therefore should not be used.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

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TEST:Influenza Virus (Types A & B) Viral CultureSynonym:Influenza Virus (Types A & B): Refer to instructions for Virus Culture.Laboratory/Phone:Virology: 443-681-3934

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TEST: Japanese Encephalitis (CDC Referral) CDC test available based on patient's travel history. Synonym: Arthropod-borne virus: Japanese Encephalitis (JE) Laboratory/Phone: Virology: 443-681-3937 Turnaround Time: 3 weeks (CDC Referral) Specimen Required: Serum (blood) Specimen identification: Label container with patient's last name, first name, DOB, specimen type, date and time of collection. Specimen Volume (Optimum): 2 ml serum Specimen Volume (Minimum): 1 ml serum Continued Next Page>

Collect:	Red-top vacutainer tube, transfer serum to sterile tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
	Write "S" for serum in the "Other Tests Request" and indicate Japanese Encephalitis.
	For testing to be initiated, the following information MUST be provided: date of onset, date specimen collected, travel history, and flavivirus vaccination history. Also please provide: patient's date of birth, diagnosis, symptoms, fatality, and whether patient is immunocompromised.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48
	hours, specimen can be frozen at -20°C and shipped on dry ice.
Specimen Rejection Criteria:	Grossly hemolyzed specimen, unlabeled specimen, leaking container, mismatch between
	labeling of specimen and test request form/electronic test order, and does not meet
	epidemiological criteria required for testing (e.g. travel history, etc.)
Availability:	Specimens shipped to the CDC Monday-Wednesday.
Results and Interpretation:	Serum that tests positive for IgM and negative for IgG is consistent with acute Japanese
	Encephalitis infection. A positive Japanese Encephalitis EIA is confirmed by PRNT (plaque
	reduction neutralization). A positive IgG antibody and a negative IgM antibody are
	consistent with infection in the distant past and are not consistent with acute infection.
Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name give to
	viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).
	Arboviruses that cause human encephalitis are members of three virus families: The
	Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae. For more information,
	see the CDC link at: <a href="https://www.cdc.gov/ncezid/dvbd/">https://www.cdc.gov/ncezid/dvbd/</a>
	Patients with travel history supporting suspicion of other arboviruses will be sent to the
	CDC for testing.
Purpose of Test:	For the presumptive detection of antibodies to Japanese Encephalitis Virus.
	Confirmatory testing by PRNT may be required.
Method:	EIA (Screening) & PRNT (Plaque Reduction Neutralization Test) referral to the Centers for
	Disease Control and Prevention (CDC).
Interfering Substances:	
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Other Arboviral testing not available at the state lab will be forwarded to the CDC based
	on patient's travel history and onset date.

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TEST:	Legionella Antigen Detection
Synonym:	Legionella Urinary Antigen
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	urine
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml urine, first void preferred
Specimen Volume (Minimum):	0.5 ml urine
Collect:	Sterile container
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Leak proof containers shipped at 2-8°C (refrigerated)
Specimen Rejection Criteria:	Insufficient specimen volume; bloody specimen
Availability:	Monday through Friday
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Results and Interpretation:	<b>POSITIVE</b> – Presumptive evidence of L. pneumophila serogroup 1 antigen in urine,
	suggesting current or past infection.
	NEGATIVE—No evidence of L. pneumophila serogroup 1 antigen in urine suggesting no
	recent or current infection. Legionnaires' disease cannot be ruled out since other
	serogroups and species may also cause disease.
Additional Information:	Only detects L. pneumophila serogroup 1. All other serogroups and other Legionella
	species must be detected by culture.
	Refer to CDC website: http://www.cdc.gov/legionella/index.html
Purpose of Test:	Detect presence of L. pneumophila serogroup 1 antigen in urine.
Method:	EIA
Interfering Substances:	Specimens may produce a false positive result from patients with bacteremia
	(Streptococcus pneumonia) pulmonary conditions and urinary tract infection (Escherichia
	coli, Enterobacter cloacae).
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Some individuals have been shown to excrete antigen for an extended period of time, so
	a positive ELISA reaction may reflect a recent but not active infection.
	Early treatment with appropriate antibiotics may also decrease antigen excretion in
	some individuals.
	Serologic results should not be used as a sole means for diagnosis, treatment, or for the
	assessment of a patient's health. Clinical correlation is required.

TEST:	Legionella Culture
Synonym:	Legionella pneumophila culture isolation/identification
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	10-14 days from receipt in the laboratory
Specimen Required:	Sputum, lung tissue, other body tissue, pleural fluid, transtracheal aspiration, lung
	exudate, lung biopsy/autopsy, lung abscess material.
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	1 ml sputum; trans tracheal aspirate, biopsy; 1 gram lung tissue; 1 ml lung exudate; 1 cc
	lung biopsy; 50 ml bronchoalveolar lavage (BAL); 1 ml lung abscess material; 7 ml blood
	in an isolator tube; collect in sterile container.
Specimen Volume (Minimum):	Half of the optimum amount
Collect:	Specimen in sterile screw capped container. Prevent specimen from drying. DO NOT
	USE SALINE IN SPECIMEN COLLECTION. BAL specimens containing saline are acceptable.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Refrigerate and ship within 48 hours; if delayed, freeze for a maximum of a week at -20°C
	and transport frozen. Transport Isolator at 2-8°C. Place each specimen in a separate,
Creative or Dejection Criterio	individually sealed bag.
Specimen Rejection Criteria:	Specimen received after prolonged delay (more than 48 hours after collection), Swab
	specimen, improper labeling; specimen received in grossly leaking transport container; urine, stool, wounds or other culture material from non-respiratory sites.
Availability:	Monday through Friday.
Results and Interpretation:	Presence of Legionella pneumophila or Legionella spp.
Results and interpretation.	Legionella not isolated
Reference Range:	Culture negative for Legionella species.
Additional Information:	http://www.cdc.gov/legionella/index.html
Purpose of Test:	Isolation and identification of Legionella species.
Method:	Culture, staining, biochemical testing.
Interfering Substances/Limitations:	Avoid contamination with normal respiratory flora.
	DHMH Laboratories Administration, Central Laboratory
Testing Site:	1770 Ashland Avenue, Baltimore, Maryland 21205
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TEST:	Legionella Serology
Synonym:	Legionella pneumophila serogroup 1-6 assay
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	Specimen identification: The specimen/sample must be properly labeled and include
	patient's name or unique patient/sample identifier matching the test requisition or
	electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	-20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 7 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	POSITIVEFour-fold rise in titer between acute and convalescent specimens indicates a
	recent infection
	<b>NEGATIVE</b> — Single titer < 1:256. In paired sera less than a four-fold increase in titer or
	<128 in the convalescent phase serum.
	INCONCLUSIVE—Single or sustained titer ≥256 may indicate past infection or exposure
	to Legionella species, diagnostic relevance cannot be determined
Additional Information:	http://www.cdc.gov/legionella/index.html
Purpose of Test:	Detect antibody to Legionella pneumophila serogroup 1-6
Method:	IFA
Interfering Substances:	Icteric, hemolyzed, lipemic
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Acute phase sera should be collected within the first week after onset of illness, and
	convalescent phase sera, 3-9 weeks after onset. Serologic results should not be used as a
	sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical
	correlation is required.

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TEST:	Leishmaniasis Serology (CDC Referral)
Synonym:	Leishmania Kala azar, Leishmania donovoni, Leishmania major
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	18 business days (CDC Referral)
Specimen Required:	Serum, plasma
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	0.5 ml. (Whole Blood)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
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Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/parasites/leishmaniasis/index.html
Purpose of Test:	Detect antibodies which may be due to Leishmania parasite infections.
Methods:	Antibody detection
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen submission. Required supplemental information: Exposure and travel history, include other relevant risk factors; clinical symptoms, treatment and relevant lab results.

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TEST:	Leptospira culture
Synonym:	Leptospira culture isolation and identification
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	Usually within 4 -6 weeks from receipt in the lab.
Specimen Required:	Urine, heparinized whole blood, CSF depending on stage of illness
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	0.1ml urine; 5 ml heparinized whole blood; 1 ml CSF. Collection in Leptosira transport
	media
Specimen Volume (Minimum):	Half of the optimum amount
Collect:	Blood (Heparin),CSF and Urine in first week of Leptospira infection. Submit urine, after
	7days of illness. Specimen should be submitted in transport media as per instruction
	provided. Transport media can be requested from DHMH lab by calling 443-681-3777.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Transport specimen at room temperature and protect from exposure to light. Do not refrigerate or freeze specimen.
Specimen Rejection Criteria:	Specimen not collected in transport media, non-sterile or leaking container, cold or
	frozen specimen. Urine held for more than 2 hours from collection or in preservative.
Availability:	Monday through Friday.
Results and Interpretation:	Presence of spirochete by darkfield microscopy may indicate positive results. Further
	confirmation test for Identification of Leptospira is required.
Additional Information:	http://www.cdc.gov/leptospirosis/
Purpose of Test:	Isolation and identification of Leptospira species.
Method:	Culture, darkfield microscopy
Interfering Substances/Limitations:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	

TEST:Leptospira SerologySynonym:Leptospira Antibody, LeptospirosisLaboratory/Phone:443-681-3938/3931Turnaround Time:5 business daysSpecimen Required:Serum, plasmaContinued Next Page>

Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	-20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 7 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	Reactive: Indicates presence of IgM antibodies. Antibody presence alone cannot be
	used for diagnosis as antibodies from prior exposure may circulate for a prolong period
	of time.
	<b>Non-reactive:</b> IgM antibody is not present in the sample or is below the detection level.
	Borderline: A second specimen should be collected in 14 days.
Additional Information:	Titers generally fall below detectable levels within 9 months to 1 year.
Purpose of Test:	Detect antibodies to Leptospira species
Method:	ImmunoDOT
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Antibody titers to leptospirosis may be delayed or substantially decreased by early and
	intensive antibiotic treatment. Serologic results should not be used as a sole means for
	diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

TEST:	Listeria monocytogenes (ABCs (previously BIDS))
Synonym:	Active Bacterial Core Surveillance (ABCs) (Bacterial Invasive Disease Surveillance) <i>Listeria monocytogenes</i> is handled as an ABCs isolate and evaluated by the National Antimicrobial Resistance Monitoring Systems (NARMS) Program. Refer to instructions for <b>ABCs (previously BIDS)</b> .
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:Lyme SerologySynonym:Borrelia burgdorferi: Refer to instructions for Borrelia burgdorferi serology.Laboratory/Phone:443-681-3938/3931

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TEST:	Lymphogranuloma venereum (LGV)
Synonym:	
Laboratory/Phone:	Virology: 443-681-3937
Turnaround Time:	Chlamydia trachomatis culture within 10 business days; CDC LGV Send-out 2-6 months
Specimen Required:	Swab: Endocervix, urethra, conjunctiva, nasopharynx, throat, rectum, vagina. For other sources, call laboratory to discuss optimum specimen. Place swab in ChlamTrans™ transport tube.
Specimen identification:	Label specimen with a minimum of the full name exactly matching slip, date of collection, source of specimen
Specimen Volume (Optimum):	2ml of media already in transport tube
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Specimen Volume (Minimum):	2ml of media already in transport tube
Collect:	Swab placed in ChlamTrans™ Transport media, or other commercial media stating it is
	appropriate for Chlamydia
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Request Chlamydia trachomatis and in the Other Tests for Infectious Agents box, write in LGV
	Indicate specimen type next to test requested using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Transport at 2-8°C
	Must reach the lab within 2 days of collection.
Specimen Rejection Criteria:	Too old, No patient ID on specimen, leaked, quantity not sufficient, no swab, expired
	transport, out of temp. range, no specimen received, broken, improper swab or
	collection kit, improper collection site, thick mucus, gross contamination, illegible ID,
	missing or incomplete lab slip (no site, date, gender, patient info., submitter info.),
	mismatched patient ID.
Availability:	Monday-Friday
Results and Interpretation:	Chlamydia trachomatis Isolated in cell culture.
	Chlamydia trachomatis not Isolated in cell culture.
	Chlamydia trachomatis toxic in cell culture. Resubmit.
Additional Information:	Cell culture will be performed at the MD State Laboratory and if Chlamydia is isolated,
	the specimen remnant will be forwarded to CDC for LGV testing. Negative cultures are
	not sent to CDC.
Purpose of Test:	Diagnostic, qualitative detection of Chlamydia followed up with LGV testing at CDC if
	positive.
Method:	Cell culture followed by Nucleic Acid Amplification/ompA gene sequencing for LGV
Interfering Substances/Limitations:	A negative result does not exclude the possibility of infection. Interpret results in
	conjunction with other information.
	Do not use ChlamTrans if leakage, evaporation, contamination or pH changes are
	apparent.
	Store ChlamTrans refrigerated.
	Do not freeze unless <-50°C. If frozen, must transport on dry ice.
Testing Site:	Cell culture:
	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
	LGV: CDC
Comment:	

TEST:	Malta Fever
Synonym:	Bang's Disease; Undulant fever; Malta Fever; Rock of Gibraltar Fever: Refer to
	instructions for Brucella serology or Brucella species, culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

TEST:	Malaria serology (CDC Referral)
Synonym:	Plamodium falciparum, Plasmodium vivax, Plasmodium malariae, parasite
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	18 business days (CDC Referral)
Specimen Required:	Serum, plasma
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	0.5 ml. (Whole Blood)
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Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
	Specimens must be packaged in a triple packaging system to ensure that under normal
Packaging and Shipping*:	conditions of transport they cannot break, be punctured or leak their contents (Refer to
Packaging and Shipping .	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
	Ambient temperature for specimens on the blood clot (whole blood specimens
Transport Conditions:	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or
	-20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/malaria/
Purpose of Test:	Detect antibodies which may be due to Plasmodium infections.
Methods:	IFA, Antibody Detection
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
Processing Site for CDC referral:	1770 Ashland Avenue, Baltimore, MD 21205
	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen
Comment:	submission. Required supplemental information: Exposure and travel history, include
	other relevant risk factors; clinical symptoms, treatment and relevant lab results.

TEST:	Measles Virus Culture
Synonym:	Measles Virus culture: Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934

TEST:	Measles IgG Antibody–Measles Immunity Screen
Synonym:	Anti Rubeola IgG; Measles IgG antibody; Rubeola / Measles immunity test
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form next to Rubeola (Measles) Immunity Screen.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or - 20°C (frozen).
Specimen Rejection:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic; lipemic; gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Service available only to state and local health departments Monday to Friday.
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Results and Interpretation:	<b>Negative:</b> Indicates no detectable IgG antibody to Measles virus. A negative result
	indicates no current or previous infection with Measles virus. Such individuals are
	presumed to be susceptible to primary infection .However, specimen taken too early
	during a primary infection may not have detectable levels of IgG antibody. If primary
	infection is suspected, another specimen (convalescent) should be taken in 8-14 days and
	tested concurrently in the same assay with the original (acute) specimen to look for
	seroconversion. If acute specimen is negative and convalescent specimen is positive,
	seroconversion has taken place and a primary Measles virus infection is indicated.
	Equivocal: Equivocal results are indeterminate. Patient may or may not have immunity to
	Measles Virus. This result is not acceptable proof of immunity.
	Positive: Indicates evidence of Measles IgG antibodies. This suggests past or current
	infection with Measles virus, via acquired immunity or immunization and probable
	protection from clinical infection. (Immunity).
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/meas.pdf
Purpose of Test:	For detection of IgG antibodies to Measles virus. The test can be used to evaluate single
	sera for immune status.
Method:	ELISA
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: A diagnosis should not be made on the basis of anti-Measles results alone.
	Test results should be interpreted in conjunction with the clinical evaluation and the
	results of other diagnostic procedures. The antibody titer of a single serum specimen
	cannot be used to determine a recent infection. Paired samples (acute and convalescent)
	should be collected and tested concurrently to demonstrate seroconversion. Samples
	collected too early in the course of an infection may not have detectable levels of IgG. In
	such cases, a second sample may be collected after 2-7 weeks and tested concurrently
	with the Original sample to look for seroconversion. A positive Measles IgG test in
	neonates should be interpreted with caution since passively acquired maternal antibody
	can persist for up to 6 months.

TEST:	Measles IgM EIA
Synonym:	Anti-Measles IgM; Rubeola/Measles IgM antibody.
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time
	of collection. The specimen/sample must be properly labeled and match the test
	requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form. Prior approval by DHMH
	Epidemiology (410-767-6628) required.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens
	transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -
	20°C (frozen).
Specimen Rejection:	Discrepancy between name on tube and name on form; unlabeled specimen; hemolytic;
	lexemic; gross bacterial contamination.
Availability:	Monday to Friday. Test available only to DHMH epidemiologists for outbreak
	investigations. Prior approval by DHMH Epidemiology (410-767-6628) required.
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Results and Interpretation:	Negative: No detectable Measles IgM antibodies. A negative result indicates no current
Results and interpretation.	infection with Measles virus. However, specimens taken too early during a primary
	infection may not have detectable levels of IgM antibody. If a primary infection is
	suspected, another specimen should be taken within 7 days and tested concurrently in the
	same assay with the original specimen to look for seroconversion.
	<b>Equivocal:</b> Equivocal specimens are indeterminate. Another specimen should be collected
	after 7 days and retested.
	Positive: Indicates evidence of Measles IgM antibodies. This suggests primary or
	reactivated infection with Measles virus.
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/meas.pdf
Purpose of Test:	For detection of IgM antibodies to measles virus. Test available only to DHMH
	epidemiologists for outbreak investigations. Prior approval by DHMH Epidemiology
	(410-767-6628) required.
Method:	ELISA
Interfering Substances:	Test results from immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: Results of the Measles IgM ELISA are not by themselves diagnostic and
	should be interpreted in light of the patient's clinical condition and results of other
	diagnostic procedures. Measles virus specific IgG antibody may compete with IgM for
	binding sites and cause false negative results. Rheumatoid factor, if present along with
	specific IgG, will cause false positive results. The Serum Diluent plus contains an absorbent
	which will remove IgG from the test specimen, and significantly reduce the possibility of
	false positive or negative results. Heterotypic IgM antibody responses may occur in
	patients infected with Epstein-Barr virus, and sera from patients with infectious
	mononucleosis may have false positive results in the Measles IgM ELISA. Samples taken
	too early during the course of a primary infection may not have detectable levels of
	Measles specific IgM. A negative result does not rule out a primary infection with virus.
	The Measles IgM ELISA cannot distinguish the difference between vaccine-induced
	antibody and antibody resulting from a natural infection. False positive IgM results may be
	obtained from patients with autoimmune disease. The performance of the Measles IgM
	ELISA has not been validated using neonatal samples.
	LLISA has not been valuated using neonatal samples.

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TEST:	Melioidosis (Burkholderia pseudomallei)
Synonym:	Burkholderia (formerly Pseudomonas) pseudomallei; B. pseudomallei; Melioidosis: Refer to
	instructions for Burkholderia mallei and Burkholderia pseudomallei.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

TEST:	Methicillin Resistant Staph aureus (MRSA) culture	
Synonym:	MRSA (rule out), Methicillin Resistant Staph aureus (MRSA) culture	
Laboratory/Phone:	Microbiology 443-681-3952	
Turnaround Time:	2-3 days [from specimen receipt in the Laboratory]	
Specimen Required:	Nasal swab; nasopharyngeal swab, tissue	
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.	
Specimen Volume (Optimum):	n Volume (Optimum): One (1) swab	
Specimen Volume (Minimum): N/A		
Collect:	Culturette tube with transport medium	
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.	
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Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal	
	conditions of transport they cannot break, be punctured or leak their contents (Refer to	
	pages 9 & 10 for triple packing guidance).	
	*Refer to current Federal regulations for specific shipping requirements.	
Transport Conditions:	Store and ship at room temperature, ship as quickly as possible.	
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Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results	
	and to avoid misleading information that might lead to misdiagnosis and inappropriate	
	therapy. A request for a new specimen will provide appropriate materials and clinically	
	relevant information to support good patient care.	
	Unlabeled or improperly labeled specimen	
	Non-sterile or leaking container	
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>	
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>	
	<ul> <li>Mismatched form and specimen</li> </ul>	
	<ul> <li>Broken specimen/sample container</li> </ul>	
	<ul> <li>The wrong specimen for test request</li> </ul>	
	<ul> <li>Inappropriate outfit for requested test</li> </ul>	
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>	
	<ul> <li>Expired transport media</li> </ul>	
	<ul> <li>Specimen received after prolonged delay (usually more than 72 hours)</li> </ul>	
Availability:	Monday through Friday	
Results and Interpretation:	MRSA isolated and identified	
Reference Range:	MRSA was not detected	
Additional Information:	N/A	
Purpose of Test:	Detect the presence of MRSA	
Method:	Broth amplification, plate culture, isolation and identification, Cefoxitin disc screen to	
	identify methicillin resistance.	
Interfering Substances:	N/A	
Testing Site:	DHMH Laboratories Administration, Central Laboratory	
-	1770 Ashland Avenue, Baltimore, Maryland 21205	
Comment:	N/A	

TEST:	MRSA (rule out)
Synonym:	Methicillin Resistant Staph aureus (MRSA) culture: Refer to instructions for Methicillin
	Resistant Staph aureus (MRSA) culture.
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:	Mumps Virus Culture
Synonym:	Mumps Virus culture: Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934
Specimens:	1 Buccal swab in VTM and urine collected in sterile container with a requisition for each specimen. Refer to instructions for Virus Culture.

TEST:	Mumps Antibody IgG EIA (Mumps Immunity Screen)
Synonym:	Anti-Mumps IgG; Mumps immunity test
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
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Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form next to Mumps Immunity
	Screen.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection:	Discrepancy between name on tube and name on form; unlabeled specimen; hemolytic; lipemic;
	gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Service available only to state and local health departments Monday to Friday.
Results and Interpretation:	<b>Negative:</b> Indicates no detectable IgG antibody to Mumps virus. A negative results indicate no current or previous infection with Mumps. Virus. Such individuals are presumed to be susceptible to primary infection .However, specimen taken too early during a primary
	infection may not have detectable levels of IgG antibody. If primary infection is suspected,
	another specimen (convalescent) should be taken in 8-14 days and tested concurrently in
	the same assay with the original (acute) specimen to look for seroconversion. If acute
	specimen is negative and convalescent specimen is positive, seroconversion has taken
	place and a primary Mumps virus infection is indicated.
	Equivocal: Equivocal results are indeterminate. Patient may or may not have immunity to
	Mumps Virus. It is not acceptable proof of immunity.
	Positive: Indicates evidence of Mumps IgG antibodies
	This suggests past or current infection with Mumps virus, via acquired immunity or
	vaccination and probable protection from clinical infection (Immunity).
Additional Information:	For more information, see the CDC link at:
	https://www.cdc.gov/vaccines/pubs/pinkbook/chapters.html
Purpose of Test:	For detection of IgG antibodies to Mumps virus, the test can be used to evaluate single sera
	for immune status.
Method:	ELISA
Interfering Substances:	Test results from an immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
-	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: A diagnosis should not be made on the basis of anti- Mumps results alone.
	Test results should be interpreted in conjunction with the clinical evaluation and the results
	of other diagnostic procedures. The antibody titer of a single serum specimen cannot be
	used to determine a recent infection. Paired samples (acute and convalescent) should be
	collected and tested concurrently to demonstrate seroconversion. Samples collected too
	early in the course of an infection may not have detectable levels of IgG. In such cases, a
	second sample may be collected after 2-7 weeks and tested concurrently with the Original
	sample to look for seroconversion. A positive Mumps IgG test in neonates should be
	interpreted with caution since passively acquired maternal antibody can persist for up to 6
	months.

Anti-Mumps IgM; antibody. Mumps IgM IFA	
Vaccine Preventable Disease/443-681-3889	
2-5 business days	
Serum	
Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.	
ecimen Volume (Optimum): 5 ml. (Whole blood) or 4 ml. (Serum)	
Specimen Volume (Minimum): 3 ml. (Whole blood) or 2 ml. (Serum)	
Red-top vacutainer	
DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)	
Indicate specimen type using the "Specimen Code" on form.	

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Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
Transport Conditions.	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic;
Specifien Rejection Criteria.	
	gross bacterial contamination. Specimens collected >7 days prior to arrival without being
	separated and frozen.
Availability:	Monday to Friday. Test available only to DHMH epidemiologists for outbreak
	investigations. Prior approval by DHMH Epidemiology (410-767-6628) required.
Results and Interpretation:	Negative: No significant level of Mumps IgM antibodies detected. A negative result
	indicates no current infection with Mumps virus. However, specimens taken too early
	during a primary infection may not have detectable levels of IgM antibody. If a primary
	infection is suspected, another specimen should be taken within 7 days and tested
	concurrently in the same assay with the original specimen to look for seroconversion
	Positive: Evidence of Mumps IgM antibodies detected and indicative of current or recent
	infection.
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/mumps.pdf
Purpose of Test:	For the detection of IgM antibodies to Mumps virus. Test available only to DHMH
	epidemiologists for outbreak investigations. Prior approval by DHMH Epidemiology (410-
	767-6628) required.
Method:	IFA
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution. IgM
	anticell antibodies, if present in the serum, may interfere with the Mumps IgM test.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: Results of the Mumps IgM ELISA are not by themselves diagnostic and
	should be interpreted in light of the patient's clinical condition and results of other
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with specific IgG, will cause false positive results. The Sample diluent contains an absorbent
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with specific IgG, will cause false positive results. The Sample diluent contains an absorbent which will remove IgG from the test specimen, and significantly reduce the possibility of false positive or negative results Heterotypic IgM antibody responses may occur in patients
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with specific IgG, will cause false positive results. The Sample diluent contains an absorbent which will remove IgG from the test specimen, and significantly reduce the possibility of false positive or negative results Heterotypic IgM antibody responses may occur in patients infected with Epstein-Barr virus. Samples taken too early during the course of a primary
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with specific IgG, will cause false positive results. The Sample diluent contains an absorbent which will remove IgG from the test specimen, and significantly reduce the possibility of false positive or negative results Heterotypic IgM antibody responses may occur in patients infected with Epstein-Barr virus. Samples taken too early during the course of a primary infection may not have detectable levels of mumps specific IgM. A negative result does not
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with specific IgG, will cause false positive results. The Sample diluent contains an absorbent which will remove IgG from the test specimen, and significantly reduce the possibility of false positive or negative results Heterotypic IgM antibody responses may occur in patients infected with Epstein-Barr virus. Samples taken too early during the course of a primary

Mycobacterium tuberculosis culture
AFB culture, Acid Fast Bacteria Identification (Acid Fast Bacilli)
Microbiology - Mycobacteriology / 443-681-3942
AFB smear: 24 hours [Note all times are from specimen receipt in the Laboratory] Nucleic Acid Amplification (MTD): 48 hours Positive culture: 14-21 days. Reported as soon as detected.
Negative culture: 8 weeks Susceptibility Testing: up to 28 days
Preferred: Sputum Other Acceptable: respiratory aspirate, bronchial wash, bronchoalveolar lavage (BAL), body fluids, CSF, tissue, urine, lymph node.
Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Sputum, aspirate or CSF: 5-10 mls Body fluid: < 50 mls
Sputum aspirate or CSF: > 1 ml Body Fluid: > 5 mls

OHMH-Laboratories Administration	The J. Mehsen Joseph Public Health Laboratory
Collect:	In a sterile, leak-proof container, e.g., a 50 ml conical tube, collection of early morning sputum specimens on each of three (3) consecutive days is optimum.
	For optimal pulmonary specimens, collect sputum from the lung after a deep, productive cough. Do not pool specimens. Label induced sputum specimens as "induced" since they resemble saliva.
	Gastric lavage specimens should be collected in a hospital and sent to the Central Laboratory immediately for processing. If specimen transport is delayed, recovery of mycobacteria is severely compromised (since mycobacteria die rapidly in gastric washing). Indicate source of specimen on the lab form. Note: If > 1 hour delay, neutralize specimen with 100 mg sodium carbonate.
	Tissue: Submit skin lesions or other tissue; keep moistened with sterile saline.
	Store refrigerated. Do not use waxed container. Keep blood and CSF at room temperature. Blood in SPS (yellow top) or Heparin (green top vacutainer.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions*:	Should be received by Central Laboratory within 24 hours after collection
*Blood and CSF should be kept at	Preferred: Refrigerate, 2-8°C
room temperature	Other Acceptable: Ambient temperature
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> </ul>
Availability:	Monday through Friday, 8:00 A.M. to 4:30 P.M.
Results and Interpretation:	AFB Smear: Acid-fast bacilli seen on smears from this specimen. The acid-fast stain does not differentiate <i>M. tuberculosis</i> from other non-tuberculous mycobacteria.
	AFB Culture: Positive culture – Mycobacterial identification given.
	Negative culture – No mycobacteria were recovered from this specimen by culture.
	Client is notified of positive smear/culture, MTD or first positive M. tuberculosis complex culture.
Referred isolate for identification:	Provide specimen collection body site and date collected.
Reference Range:	Complete identification of clinically significant isolates. Antimicrobial susceptibilities performed on all initial isolates of <i>M. tuberculosis</i> complex. Drug resistant isolates will be tested for susceptibility to second-line anti-mycobacterial drugs. Anti-microbial susceptibilities performed on Mycobacterium other than <i>M. tuberculosis</i> complex isolated by request with justification for testing (immunocompromised patient, multiple site isolates, HIV patient, etc.).
Additional Information:	DNA probes (cultures only) available for M. tuberculosis complex, M. avium-intracellulare complex, M. gordonae and M. kansasii as indicated.
Purpose of Test:	The AFB smear can determine the presence of mycobacteria in clinical specimens by microscopic examination. AFB smears are made from the sediments of specimens that have been decontaminated and concentrated by centrifugation for culture. Special solid and liquid growth media are inoculated with the concentrated specimen for isolation and
Method:	identification of mycobacteria. Standard reference procedures for stain and culture. Biochemical standard reference
	procedures are used for rapid growers.
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Interfering Substances:	Propylene glycol, waxed containers, tap water (may contain saprophytic mycobacteria), antimicrobial therapy, food particles, mouthwash.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Nucleic Acid Amplification Assay:	Will be done on all new smear positive patient specimens or referred specimen concentrates on patients with a high suspicion for active tuberculosis. Patient must be on treatment < seven (7) days or not at all. Test should not be requested routinely. In our experience, the sensitivity and specificity of the test on smear positive specimens is 98.7% and 97.8%, respectively. On smear negative specimens, the sensitivity and specificity is 62.2% and 98.9%, respectively. (Chest 2007; 132: 946-951)

TEST:	Mycoplasma Serology
Synonym:	Mycoplasma pneumoniae
Laboratory/Phone:	443-681-3938
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer tube
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	<b>NEGATIVE</b> —No significant amount of IgG/IgM antibodies detected, no presumptive evidence of current/previous infection
	<b>POSITIVE</b> —IgG/IgM antibodies detected, evidence of a past/recent infection
	EQUIVOCAL—Immunological status cannot be determined. Please redraw patient in 1-3
	weeks
Additional Information:	http://www.cdc.gov/pneumonia/atypical/mycoplasma/
Purpose of Test:	Detect antibodies to M. pneumoniae
Methods:	EIA
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	May not detect a recent infection. If suspicion of a Mycoplasma infection, take a second
	sample at least 14 days later for additional testing.
	Serologic results should not be used as a sole means for diagnosis, treatment, or for the
	assessment of a patient's health. Clinical correlation is required.

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 TEST:
 Neisseria gonorrhoeae Culture

 Synonym:
 GC Culture; Gonorrhea Culture; N. gonorrhoeae Culture: Refer to instructions for Gonorrhea Culture.

 Laboratory/Phone:
 Microbiology 443-681-3952

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The J. Mehsen Joseph Public Health Laboratory

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Neisseria meningitidis (ABCs - previously BIDS))
Active Bacterial Core Surveillance (ABCs) (Bacterial Invasive Disease Surveillance) Neisseria
<i>meningitidis</i> : Refer to instructions for ABCs (previously BIDS).
Microbiology 443-681-3952
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TEST:	Ohara's disease
Synonym:	Francisella tularensis, Pasteurella tularensis, tularemia, rabbit fever, deerfly fever, Ohara's
	disease, Francis disease: Refer to instructions for Francisella tularensis culture or
	Francisella tularensis Antibody.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Ova and Parasites Microscopic Examination
Synonym:	Amebiasis, Giardia, Parasitic identification, worm identification
Laboratory/Phone:	Microbiology 443-681-3952 or 443-681-4570
Turnaround Time:	5 business days [Note time is from specimen receipt in the Laboratory]
Specimen Required:	Feces: Minimum of three (3) specimens collected over a 7-10 day period.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Please refer to the directions available with stool collection kit. There is no maximum limit on the amount of stool collected.
Specimen Volume (Minimum):	Please refer to the directions available with stool collection kit. As a minimum amount, collect several grams (or teaspoon amounts).
Collect:	Please refer to the directions available with stool collection kit.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Send the specimen to the laboratory as soon as possible at room temperature.
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Genus and species
Reference Range:	No Ovajor Parasites found
Additional Information:	Collect all fecal specimens prior to the administration of antibiotics or anti-diarrheal agents Avoid contamination with urine or water from the toilet.
Purpose of Test:	Diagnosis of intestinal parasite
Method:	Microscopic: Wet mount and permanent stain using Eco-fix and Eco-stain.
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Interfering Substances:	Avoid the use of mineral oil, bismuth and barium prior to fecal collection since all of these substances may interfere with detection or identification of intestinal parasites.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

TEST:	Parainfluenza Virus (Types 1, 2, and 3) Viral Culture
Synonym:	Parainfluenza Virus (Types 1, 2, and 3): Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934

TEST:	Parasitic examination (Ova and Parasites Microscopic Examination)
Synonym:	Amebiasis, Giardia, Entamoeba, Parasite identification, worm identification: Refer to
	instructions for Ova and Parasites Microscopic Examination.
Laboratory/Phone:	Microbiology 443-681-3952

TEST:	Pasteurella tularensis (Francisella tularensis) culture
Synonym:	Pasteurella tularensis, tularemia, rabbit fever, deerfly fever, Ohara's disease, Francis
	disease: Refer to instructions for Francisella tularensis culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

TEST:	Pertussis (Bordetella pertussis) PCR & Culture
Synonym:	B. pertussis, pertussis, Whooping Cough Refer to instructions for Bordetella pertussis PCR and culture.
Laboratory/Phone:	Molecular Biology: 443-681-3924 Microbiology 443-681-3952

TEST:	Pertussis Serology (Bordetella pertussis)
Synonym:	IgG Anti-Bordetella pertussis toxin assay. Refer to instructions for Bordetella Pertussis
	Toxin IgG Antibody
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889

TEST:	Pinworm Examination
Synonym:	Cellulose tape preparation for Enterobius vermicularis
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	24 hrs [from specimen receipt in the Laboratory] Monday through Friday
Specimen Required:	Cellulose tape preparation from the skin of the perianal area.
Specimen identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,
	specimen type/source, and the date and time of collection. The specimen/sample must be
	properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	N/A
Specimen Volume (Minimum):	N/A
Collect:	To obtain a sample from the perianal area, peel back the tape by gripping the labeled end, and, with the tape looped (adhesive side outward) over a wooden tongue depressor that is held firmly against the slide and extended about 2-5 cm beyond it, press the tape firmly several times against the right and left perianal folds. Smooth the tape back on the slide, adhesive side down. Label with patient's name and date.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
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Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Illegible or no patient information on the specimen</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Organism and stage
Reference Range:	Enterobius vermicularis NOT found
Additional Information:	Pinworm eggs are usually infectious. The female pinworm deposits eggs on the perianal skin only sporadically, without multiple tapes (taken consecutively, each morning), it is not possible to determine if the patient is positive or negative for the infection.
Purpose of Test:	Detection of human pinworm infections
Method:	Microscopic
Interfering Substances:	Opaque tape
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

TEST:	Plague (Yersinia pestis)
Synonym:	Plague; Yersinia pestis; Pasteurella pestis: Refer to instructions for Yersinia pestis culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Polio Virus, Virus Culture
Synonym:	Polio Virus Culture (Enterovirus, including Echovirus, Coxsackie, and Polio): Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934
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TEST:	Q-fever serology
Synonym:	Coxiella burnetii, Q-fever: Refer to instructions for Coxiella Serology.
Laboratory/Phone:	443-681-3938/3931

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TEST:	Rabbit fever
Synonym:	Francisella tularensi; Pasteurella tularensis, tularemia, deerfly fever, Ohara's disease,
	Francis disease: Refer to instructions for Francisella tularensis culture or Francisella
	tularensis Antibody.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Rabies Antibody Titer (RFFIT)
Synonym:	RFFIT Test
Laboratory/Phone:	Division of Virology and Immunology/Rabies Lab 443-681-3771
Turnaround Time:	15 working days
Specimen Required:	Serum/Blood
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier, date of birth, and specimen collection date matching the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml whole blood or 2 ml of serum
Specimen Volume (Minimum):	2 ml whole blood or 1 ml serum
Collect:	Red-top vacutainer or Zebra-top serum separator vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Whole blood specimens transported on ice packs;separated serum at 2-8°C (refrigerated)
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled tube; insufficient quantity of serum for testing; hemolysis; lipemia; gross bacterial contamination.
Availability:	Monday through Friday
Results and Interpretation:	Positive 0.5 IU/mL or greater (immunity) Negative indicates no detectable antibody to the rabies virus or the presence of detectable antibody < 0.5 IU/mL.
Reference Range:	Patient's with a titer > 0.5 IU/mL. is considered to have adequate immune response.
Additional Information:	Provide patient's rabies vaccination history.
Purpose of Test:	For detection of rabies antibody
Method:	Rapid Fluorescent Focus Inhibition Test (RFFIT)
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Restricted Test: Services provided to State and Local government employees (e.g. animal control, etc.). Maryland residents requiring testing refer to the Rabies Laboratory website: <u>http://dhmh.maryland.gov/laboratories/Pages/Rabies.aspx</u>

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TEST:	Rat Bite Fever
Synonym:	Streptobacillus moniliformis Culture; Haverhill Fever: Refer to instructions for
	Streptobacillus moniliformis Culture.
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:	Respiratory Syncytial Virus (RSV) Virus Culture
Synonym:	Respiratory Syncytial Virus (RSV): Refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934

TEST:	Rock of Gibraltar Fever
Synonym:	Brucellosis, Bang's Disease, Undulant fever, Malta Fever: Refer to instructions for Brucella
	serology or <i>Brucella</i> species culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Rocky Mountain Spotted Fever (RMSF) Antibody
Synonym:	Rickettsia rickettsii serology
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	Label tube with patients first and last name
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or - 20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	Titers ≥ 1:64 are suggestive of possible early infection, declining titers due to past exposure, or cross-reactivity with a related organism.
Additional Information:	http://www.cdc.gov/rmsf/ A second specimen will usually demonstrate a diagnostic four fold rise in titer for patients with active disease.
Purpose of Test:	Detect antibodies to R. rickettsii
Methods:	IFA
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Results are for epidemiological purposes only. Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

TEST:	Rubella IgG (Rubella Immunity Screen).
Synonym:	Anti-Rubella IgG; German Measles IgG antibody; Rubella immunity test
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form next to Rubella Immunity Screen.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or 20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic; lexemic; gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Service available only to state and local health departments Monday to Friday.
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Results and Interpretation:	<ul> <li>Negative: Indicates no detectable IgG antibody to Rubella virus. A negative results indicates no current or previous infection with Rubella virus. Such individuals are presumed to be susceptible to primary infection .However, specimen taken too early during a primary infection may not have detectable levels of IgG antibody. If primary infection is suspected, another specimen (convalescent) should be taken in 8-14 days and tested concurrently in the same assay with the original (acute) specimen to look for seroconversion. If acute specimen is negative and convalescent specimen is positive, seroconversion has taken place and a primary rubella virus infection is indicated.</li> <li>Equivocal: Equivocal results are indeterminate. Patient may or may not have immunity to Rubella Virus. It is not acceptable proof of immunity.</li> <li>Positive: Indicates evidence of Rubella IgG antibodies</li> <li>This suggests past or current infection with Rubella virus, via acquired immunity or</li> </ul>
	vaccination and probable protection from clinical infection (Immunity).
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/rubella.pdf
Purpose of Test:	For detection of IgG antibodies to Rubella virus. The test can be used to evaluate single
	sera for immune status or paired sera to demonstrate seroconversion.
Method:	ELISA
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: A diagnosis should not be made on the basis of anti-Rubella results alone. Test results should be interpreted in conjunction with the clinical evaluation and the results of other diagnostic procedures. The antibody titer of a single serum specimen cannot be used to determine a recent infection. Paired samples (acute and convalescent) should be collected and tested concurrently to demonstrate seroconversion. Samples collected too early in the course of an infection may not have detectable levels of IgG. In such cases, a second sample may be collected after 2-7 weeks and tested concurrently with the Original sample to look for seroconversion. A positive rubella IgG test in neonates should be interpreted with caution since passively acquired maternal antibody can persist for up to 6 months.

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TEST:	Rubella IgM Antibody
Synonym:	Anti-Rubella IgM; Rubella IgM antibody for Rubella/ German Measles - acute infection
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form. Prior approval by DHMH Epidemiology (410-767-6628) required.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic; lipemic; gross bacterial contamination.
Availability:	Monday to Friday. Test available only to DHMH epidemiologists for outbreak investigations. Prior approval by DHMH Epidemiology (410-767-6628) required.
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Desults and Internations	Negetive: Indicates as detected to Dubelle Intel entities dies. A second in the distance of
Results and Interpretation:	<b>Negative:</b> Indicates no detectable Rubella IgM antibodies. A negative result indicates no
	current infection with rubella virus. However, specimens taken too early during a primary
	infection may not have detectable levels of IgM antibody. If a primary infection is
	suspected, another specimen should be taken within 7 days and tested concurrently in the
	same assay with the original specimen to look for seroconversion
	<b>Equivocal:</b> Equivocal specimens are indeterminate. Another specimen should be collected
	after 7 days and retested.
	Positive: Indicates evidence of Rubella IgM antibodies.
	This suggests primary or reactivated infection with Rubella.
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/rubella.pdf
Purpose of Test:	Test available only to DHMH epidemiologists for outbreak investigations. Prior approval
	by DHMH Epidemiology (410-767-6628) required.
Method:	ELISA
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: Results of the Rubella IgM ELISA are not by themselves diagnostic and
	should be interpreted in light of the patient's clinical condition and results of other
	diagnostic procedures. Rubella virus specific IgG antibody may compete with IgM for
	binding sites and cause false negative results. Rheumatoid factor, if present along with
	specific IgG, will cause false positive results. The Sample Diluent contains an absorbent
	which will remove IgG from the test specimen, and significantly reduce the possibility of
	false positive or negative results. Heterotypic IgM antibody responses may occur in
	patients infected with Epstein-Barr virus, and sera from patients with infectious
	mononucleosis may have false positive results in the rubella IgM ELISA .Samples taken
	too early during the course of a primary infection may not have detectable levels of
	rubella specific IgM. A negative result does not rule out a primary infection. It cannot
	distinguish the difference between vaccine-induced antibody and antibody resulting from
	a natural infection. False positive anti-rubella IgM results may be obtained from patients
	with autoimmune disease. The performance of the Rubella IgM EIA has not been
	validated using neonatal samples.
	valuated using neonatal samples.

TEST:	Salmonella Culture Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins– producing <i>E. coli</i> )
Synonym:	Stool culture for enteric pathogens; enteric pathogens; stool culture and sensitivity; feces culture: Refer to instructions for Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins–producing <i>E. coli</i> ).
Laboratory/Phone:	Microbiology-Enterics 443-681-4570

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TEST:	Salmonella typing
Synonym:	Salmonella isolate for typing (referral isolate)
Laboratory/Phone:	Microbiology-Enterics 443-681-4570
Turnaround Time:	For epidemiological purposes only. CDC TAT: 8 weeks. For additional questions, contact the laboratory 443-681-4570
Specimen Required:	Pure culture on agar slant in screw cap tube.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Salmonella isolated from culture
Specimen Volume (Minimum):	N/A
Collect:	N/A
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
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Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	At room temperature. Do not freeze or refrigerate.
Specimen Rejection Criteria:	<ul> <li>The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care.</li> <li>Unlabeled or improperly labeled specimen</li> <li>Non-sterile or leaking container</li> <li>Inappropriate specimen transport conditions</li> <li>Illegible, or no submitter information on the request form</li> <li>Mismatched form and specimen</li> <li>Broken specimen/sample container</li> <li>The wrong specimen for test request</li> <li>Inappropriate outfit for requested test</li> <li>Illegible or no patient information on the specimen</li> <li>Expired transport media</li> <li>Specimen frozen</li> </ul>
Availability:	Monday through Friday
Results and Interpretation:	Salmonella somatic and flagellar antigens identified.
Reference Range:	N/A
Additional Information:	SUBCULTURE TO AGAR SLANT BEFORE TRANSPORTING. DO NOT SEND CULTURE PLATES. MAKE SURE CULTURE IS GROWING/VIABLE.
Purpose of Test:	Salmonella serotyping
Method:	Isolate is subcultured to confirm purity. Salmonella serological testing is performed by slide agglutination and tube agglutination tests using somatic (O) and flagella (H) antisera. Biochemical identification also.
Interfering Substances/ Limitations:	Submission of isolate on inhibitory media.
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

TEST: Schistosoma serology Synonym: Schistosomiasis, Schistosoma mansoni, Schistosoma haematobium, Schistosoma japonicum Bilharzia Laboratory/Phone: 443-681-3938/3931 Turnaround Time: 5 business days Specimen Required: Serum Specimen identification: The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order. Specimen Volume (Optimum): 2 ml. (Whole Blood) Specimen Volume (Minimum): 1 ml. (Whole Blood) Collect: Red-top vacutainer Form: DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form. Packaging and Shipping\*: Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). \*Refer to current Federal regulations for specific shipping requirements. Ambient temperature for specimens on the blood clot (whole blood specimens Transport Conditions: transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or 20°C (frozen). Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being Specimen Rejection Criteria: frozen Availability: Monday through Friday Reactive: IgG antibodies to a Schistosoma species were detected. **Results and Interpretation:** Non-Reactive: IgG antibodies to a Schistosoma species were NOT detected. For CDC Referral see CDC interpretations on report. Additional Information: http://www.cdc.gov/parasites/schistosomiasis/disease.html Purpose of Test: Detects antibodies to Schistosoma. Methods: EIA Continued Next Page>

Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Testing/Processing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Specimens can be referred to the CDC upon request. Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen submission. Required supplemental information: Exposure and travel history, include other relevant risk factors; clinical symptoms, treatment and relevant lab results. CDC Turnaround Time is 21 business days. Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

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TEST:	Shiga toxins-producing E. coli Culture
Synonym:	Stool culture for enteric pathogens; enteric pathogens; stool culture and sensitivity; feces culture: Refer to instructions for Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins–producing <i>E. coli</i> ).
Laboratory/Phone:	Microbiology-Enterics 443-681-4570

TEST:	Shigella Culture
Synonym:	Stool culture for enteric pathogens; enteric pathogens; stool culture and sensitivity; feces culture: Refer to instructions for Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins–producing <i>E. coli</i> ).
Laboratory/Phone:	Microbiology - Enterics 443-681-4570

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TEST:	Shigella typing	
Synonym:	Shigella isolate for typing (referral isolate)	
Laboratory/Phone:	Microbiology - Enterics / 443-681-4570	
Turnaround Time:	Usually 3-5 days [from receipt in the Laboratory]. CDC TAT: 8 weeks	
Specimen Required:	Pure culture on agar slant in screw cap tube.	
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB,	
	specimen type/source, and the date and time of collection. The specimen/sample must be	
	properly labeled and match the test requisition or electronic test order.	
Specimen Volume (Optimum):	Shigella isolated from culture	
Specimen Volume (Minimum):	N/A	
Collect:	N/A	
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)	
	Indicate specimen type using the "Specimen Code" on form.	
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal	
	conditions of transport they cannot break, be punctured or leak their contents (Refer to	
	pages 9 & 10 for triple packing guidance).	
	*Refer to current Federal regulations for specific shipping requirements.	
Transport Conditions:	At room temperature. Do not freeze or refrigerate.	
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results	
	and to avoid misleading information that might lead to misdiagnosis and inappropriate	
	therapy. A request for a new specimen will provide appropriate materials and clinically	
	relevant information to support good patient care.	
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>	
	<ul> <li>Non-sterile or leaking container</li> </ul>	
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>	
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>	
	<ul> <li>Mismatched form and specimen</li> </ul>	
	<ul> <li>Broken specimen/sample container</li> </ul>	
	<ul> <li>The wrong specimen for test request</li> </ul>	
	<ul> <li>Inappropriate outfit for requested test</li> </ul>	
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>	
	<ul> <li>Expired transport media</li> </ul>	
	Specimen frozen	
Availability:	Monday through Friday	

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Results and Interpretation:	Shigella somatic antigens identified
Reference Range:	N/A
Additional Information:	SUBCULTURE TO AGAR SLANT BEFORE TRANSPORTING. DO NOT SEND CULTURE PLATES.
	MAKE SURE CULTURE IS VIABLE/GROWING.
Purpose of Test:	Shigella serotyping
Method:	Isolate is subcultured to confirm purity. Shigella serological testing is performed by a slide agglutination test using somatic (O) antisera. Biochemical analysis performed to verify Shigella identification.
Interfering	Submission of isolate on inhibitory media.
Substances/Limitations:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	N/A

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TEST:	St. Louis Encephalitis Virus (SLEV) (Arbovirus Endemic Panel)
	Panel includes WNV IgM, SLE IgM, and EEEV IgM. LaCrosse (LAC) IgM testing available
	based on patient's travel history.
Synonym:	Arthropod-borne virus: WNV (West Nile Virus), EEEV (Eastern Equine Encephalitis Virus),
	SLEV (St. Louis Encephalitis Virus): Refer to instructions for Arbovirus Endemic Panel.
Laboratory/Phone:	Virology: 443-681-3937

TEST:	Staph aureus Culture
Synonym:	Staph aureus Culture: Refer to instructions for Foodborne Pathogens, Foodborne
	Pathogenic Microorganisms, Stool Culture.
Laboratory/Phone:	Microbiology 443-681-3952

TEST:	Stool Culture Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins– producing <i>E. coli</i> )
Synonym:	Stool culture for enteric pathogens; enteric pathogens; stool culture and sensitivity; feces culture: Refer to instructions for Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins–producing <i>E. coli</i> )
Laboratory/Phone:	Microbiology-Enterics 443-681-4570

TEST:	Streptobacillus moniliformis Culture
Synonym:	Rat Bite Fever; Haverhill Fever.
Laboratory/Phone:	Microbiology 443-681-3952
Turnaround Time:	2-3 weeks [from specimen receipt in the Laboratory]
Specimen Required:	Blood is the specimen of choice. Joint fluid, abscess fluid, wound exudates and lymph node are also acceptable.
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	Draw enough blood into the blood culture bottle to make about 20% of the total volume. If citrated blood is collected, draw a total of 10 ml.
Specimen Volume (Minimum):	N/A
Collect:	Follow the blood culture kit instructions.
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Room temperature
	Continued Next Page>

Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results
	and to avoid misleading information that might lead to misdiagnosis and inappropriate
	therapy. A request for a new specimen will provide appropriate materials and clinically
	relevant information to support good patient care.
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>
	<ul> <li>Non-sterile or leaking container</li> </ul>
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>
	<ul> <li>Mismatched form and specimen</li> </ul>
	<ul> <li>Broken specimen/sample container</li> </ul>
	<ul> <li>The wrong specimen for test request</li> </ul>
	<ul> <li>Inappropriate outfit for requested test</li> </ul>
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>
	Expired transport media
Availability:	Monday through Saturday
Results and Interpretation:	S. moniliformis present
Reference Range:	S. moniliformis NOT found.
Additional Information:	Because special enrichment of media is necessary, the laboratory needs to know that an
	infection with S. moniliformis is suspected.
Purpose of Test:	Cultural confirmation of rat bite fever is very helpful for diagnosis, since the disease is not
•	commonly seen.
Method:	Culture, convention and biochemicals.
Interfering Substances:	SPS in blood culture broth.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
-	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Serological tests are not readily available

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TEST:	Streptococcus pneumoniae (ABCs - previously BIDS))
Synonym:	Active Bacterial Core Surveillance (ABCs) (Bacterial Invasive Disease Surveillance)
	Streptococcus pneumoniae: Refer to instructions for ABCs (previously BIDS)
Laboratory/Phone:	Microbiology 443-681-3952

TEST:	Streptococcus pyogenes culture
Synonym:	Group A Strep culture; Throat culture for Group A Strep Beta; Strep culture; Streptococcus
	pyogenes culture: Refer to instructions for Group A Strep Culture.
Laboratory/Phone:	Microbiology 443-681-3952
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TEST:	Strongyloides serology
Synonym:	Strongyloidiasis; Strongloides stercoralis
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
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Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
-	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	Reactive: IgG antibodies to Strongyloides stercoralis were detected
	Non-Reactive: IgG antibodies to Strongyloides stercoralis were NOT detected.
	For CDC Referral see CDC interpretations on report.
Additional Information:	http://www.cdc.gov/parasites/strongyloides/
Purpose of Test:	Detects antibodies to Strongyloides.
Methods:	EIA
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Testing/Processing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Specimens can be referred to the CDC upon request.
	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen
	submission. Required supplemental information: Exposure and travel history, include
	other relevant risk factors; clinical symptoms, treatment and relevant lab results.
	CDC Turnaround Time is 21 business days.
	Results are for epidemiological purposes only. Serologic results should not be used as a sole
	means for diagnosis, treatment, or for the assessment of a patient's health. Clinical
	correlation is required.

TEST:	Syphilis EIA
Synonym:	Treponema pallidum
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum or plasma
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 7 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	<b>NEGATIVE</b> —Very low or now antibody is present in the sample. Does not rule out a recent
	or current infection
	<b>POSITVE</b> —Antibody is present as a result of previous or current infection with T. pallidum
	EQUIVOCAL—Suspect for infection with T. pallidum. Please submit another specimen in 2
	weeks for retesting.
Additional Information:	http://www.cdc.gov/std/syphilis/
Purpose of Test:	Detect antibodies (IgG & IgM) to treponema pallidum.
Methods:	EIA
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
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Comment:	All treponemal tests tend to remain reactive following treponemal infection; therefore, they should not be used to evaluate response to therapy. Because of the persistence of reactivity, probably for the life of the patient, the treponemal tests are of no value to the clinician in determining relapse or re-infection in a patient who has had a reactive result. Serologic results should not be used as a sole means for diagnosis, treatment, or for the
	assessment of a patient's health. Clinical correlation is required.

TEST:	Syphilis-RPR Serology
Synonym:	Rapid Plasma Reagin, Detect reagin antibodies associated with syphilis
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	3 business days
Specimen Required:	Serum/Plasma
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 7 days prior to arrival without being
	frozen
Availability:	Monday through Friday
Results and Interpretation:	REACTIVE- Non-Treponemal antibodies detected.
	NON-REACTIVE- Non-Treponemal antibodies not detected. False negatives occur in
	incubating primary and in latent syphilis
Additional Information:	Detect antibodies which may be due to syphilis or to quantify reagin antibodies associated
	with syphilis infections or to monitor response to treatment.
Purpose of Test:	Detect non-treponemal antibodies which may be due to syphilis
Method:	RPR (Rapid Plasma Reagin)
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	RPR tests may be non-specifically reactive in other conditions. Absence of reaginic antibody
	does not necessarily indicate inactive infection.
	Reactive specimens are quantitatively tested and reflexed to Syphilis IgG/IgM EIA and
	further serological study.
	Serologic results should not be used as a sole means for diagnosis, treatment, or for the
	assessment of a patient's health. Clinical correlation is required.

TEST:	Syphilis Serology -VDRL
Synonym:	Venereal Disease Research Laboratory
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Cerebrospinal fluid (CSF)
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml.
Specimen Volume (Minimum):	1 ml.
Collect:	Sterile CSF
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Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	NON-REACTIVE indicates that the patient does not have detectable non-treponemal
	antibody.
	REACTIVE indicates past or present syphilis infection of the central nervous system.
Additional Information:	This test is only performed on Cerebrospinal fluid (CSF)
Purpose of Test:	Detect antibodies which may be due to syphilis
Methods:	Slide flocculation test
Interfering Substances:	Blood
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	The VDRL is a non-treponemal test to detect lipoidal antigen to T. pallidum. VDRL is run on
	spinal fluid specimens only, for suspected neurosyphilis.

TEST:	Throat Culture (Group A Strep Culture)
Synonym:	Throat culture for Group A Strep Beta; Strep culture; <i>Streptococcus pyogenes</i> culture: Refer
	to instructions for Group A Strep Culture.
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:	Throat culture (Bacterial Culture, Routine)
Synonym:	Aerobic culture, routine culture, throat culture: Refer to instructions for <b>Bacterial Culture</b> , <b>Routine</b> .
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:	Tick identification/Ectoparasite
Synonym:	Arthropod Identification; Tick identification/Ectoparasite: refer to instructions for
	Arthropod Identification.
Laboratory/Phone:	Microbiology 443-681-3952

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TEST:	Toxocara serology (CDC Referral)
Synonym:	Toxocara canis, Toxacara cati, Toxocariasis, Larva migrans, parasite
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	18 business days (CDC Referral)
Specimen Required:	Serum, plasma
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
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Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/parasites/toxocariasis/
Purpose of Test:	Detect antibodies which may be due Toxocara canis infections.
Methods:	EIA, ELISA, Antibody Detection
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen
	submission. Required supplemental information: Exposure and travel history, include
	other relevant risk factors; clinical symptoms, treatment and relevant lab results.

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TEST:	Toxoplasma Serology
Synonym:	Toxoplasma gondii
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	NEGATIVE—No detectable IgG/IgM antibody to Toxoplasma gondii           POSITIVE—Detectable IgG/IgM antibody to Toxoplasma gondii indicating current or previous infection           EQUIVOCAL—Immunological status cannot be determined. Please submit a new specimen within 3 weeks for retesting
Additional Information:	
Purpose of Test:	Detect T. gondii antibodies (IgM & IgG) .
Methods:	EIA, ELFA
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required. The presence of IgG antibody against a particular virus or organism may not assure protection from that disease.

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TEST:	Trichinellosis Serology (CDC Referral)
Synonym:	Trichinosis, Trichnella spiralis
Laboratory/Phone:	443-681-3938/3931
Turnaround Time:	18 business days (CDC Referral)
Specimen Required:	Serum, plasma
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique
Specimen identification.	patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	0.5 ml. (Whole Blood)
Collect:	Red-top vacutainer (Serum) or Lavender-top vacutainer (Plasma)
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
FOIIII.	Indicate specimen type using the "Specimen Code" on form.
	Specimens must be packaged in a triple packaging system to ensure that under normal
Packaging and Shipping*:	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
Transport conditions.	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume
Availability:	Monday through Friday
Results and Interpretation:	Given on CDC report
Additional Information:	http://www.cdc.gov/parasites/trichinellosis/
Purpose of Test:	Detect antibodies which may be due Trichinella infections.
Methods:	EIA, ELISA, Antibody Detection
Interfering Substances:	Icteric, hemolyzed, lipemic specimen
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
Processing Site for CDC referral.	1770 Ashland Avenue, Baltimore, MD 21205
Comment:	Contact the MD DHMH Epidemiologist at (410)767-6700 for prior approval of specimen
	submission. Required supplemental information: Exposure and travel history, include
	other relevant risk factors (consumption of raw or undercooked pork or game meat);
	clinical symptoms, treatment and relevant lab results.

TEST:	Tuberculosis Bacteriology Culture (AFB/Mycobacterium Identification)
Synonym:	Acid Fast Bacteria Identification (Acid Fast Bacilli); M. Tuberculosis culture: Refer to
	instructions for Mycobacterium tuberculosis culture.
Laboratory/Phone:	Microbiology - Mycobacteriology 443-681-3942

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TEST:	Tularemia
Synonym:	Francisella tularensis culture, Pasteurella tularensis, rabbit fever, deerfly fever, Ohara's
	disease, Francis disease: Refer to instructions for Francisella tularensis culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Typhus Fever Serology
Synonym:	(Murine typhus); Typhus Fever Antibody; <i>R. typhi</i> serology
Laboratory/Phone:	443-681-3938
Turnaround Time:	5 business days
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include patient's name or unique patient/sample identifier matching the test requisition or electronic test order.
Specimen Volume (Optimum):	2 ml. (Whole Blood)
Specimen Volume (Minimum):	1 ml. (Whole Blood)
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Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Hemolysis; insufficient volume, specimen collected > 5 days prior to arrival without being frozen
Availability:	Monday through Friday
Results and Interpretation:	Titers $\geq$ 1:64 are suggestive of possible early infection, declining titers due to past exposure, or cross-reactivity with a related organism.
Additional Information:	A second specimen will usually demonstrate a diagnostic four fold rise in titer for patients with active disease
Purpose of Test:	Detect R. typhii antibodies (IgG).
Methods:	IFA
Interfering Substances:	Icteric, hemolyzed, lipemic or heat inactivation of specimen
Testing Site:	DHMH Laboratories Administration, Central Laboratory 1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Results are for epidemiological purposes only. Serologic results should not be used as a sole means for diagnosis, treatment, or for the assessment of a patient's health. Clinical correlation is required.

TEST:	Undulant fever
Synonym:	Brucellosis, Bang's Disease, Malta Fever, and Rock of Gibraltar Fever: Refer to instructions
	for Brucella serology or Brucella species, culture.
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

TEST: Urine culture (Bacterial Culture, Routine)	

TEST:	Urine culture (Bacterial Culture, Routine)
Synonym:	Aerobic culture, routine urine culture, urine culture: Refer to instructions for Bacterial
	Culture, Routine
Laboratory/Phone:	Microbiology 443-681-3952
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TEST:	Varicella Antibody IgG (Varicella Immunity Screen)
Synonym:	Anti-Varicella/ Varicella Zoster Virus (VZV)/Chickenpox IgG; Varicella immunity test.
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	2-5 business days
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form next to Varicella Immunity Screen.
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Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic;
	lipemic; gross bacterial contamination. Specimens collected > 7 days prior to submission.
Availability:	Service available only to state and local health departments Monday to Friday.
Results and Interpretation:	<b>Negative:</b> Indicates no detectable Varicella IgG antibodies. A negative results indicate no
	current or previous infection with Varicella virus. Such individuals are presumed to be
	susceptible to primary infection .However, specimen taken too early during a primary
	infection may not have detectable levels of IgG antibody. If primary infection is suspected,
	another specimen (convalescent) should be taken in 8-14 days and tested concurrently in
	the same assay with the original (acute) specimen to look for seroconversion. If acute
	specimen is negative and convalescent specimen is positive, seroconversion has taken
	place and a primary varicella virus infection is indicated.
	<b>Equivocal:</b> Equivocal results are indeterminate. Patient may or may not have immunity to
	Varicella Virus. It is not acceptable proof of immunity.
	<b>Positive:</b> Indicates evidence of Varicella IgG antibodies
	This suggests past or current infection with Varicella virus via acquired immunity or
	vaccination and probable protection from clinical infection (Immunity).
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/varicella.pdf
Purpose of Test:	For detection of IgG antibodies to Varicella virus. The test can be used to evaluate single
	sera for immune status.
Method:	ELISA
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
C C	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: A diagnosis should not be made on the basis of anti-Varicella results alone.
	Test results should be interpreted in conjunction with the clinical evaluation and the results
	of other diagnostic procedures. The antibody titer of a single serum specimen cannot be
	used to determine a recent infection. Paired samples (acute and convalescent) should be
	collected and tested concurrently to demonstrate seroconversion. Samples collected too
	early in the course of an infection may not have detectable levels of IgG. In such cases, a
	second sample may be collected after 2-7 weeks and tested concurrently with the Original
	sample to look for seroconversion. A positive Varicella IgG test in neonates should be
	interpreted with caution since passively acquired maternal antibody can persist for up to 6

TEST:	Varicella Antibody (IgM)
Synonym:	Anti-Varicella IgM; Varicella Zoster Virus/VZV antibody.
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Turnaround Time:	Serum
Specimen Required:	Serum
Specimen identification:	Label container with patient's last name, first name, DOB, specimen type, date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
Specimen Volume (Optimum):	5 ml. (Whole blood) or 4 ml. (Serum)
Specimen Volume (Minimum):	3 ml. (Whole blood) or 2 ml. (Serum)
Collect:	Red-top vacutainer
Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form. Prior approval by DHMH Epidemiology (410-767-6628) required.

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Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Ambient temperature for specimens on the blood clot (whole blood specimens transported
	on ice packs are acceptable), separated serum at 2-8°C (refrigerated) or -20°C (frozen).
Specimen Rejection Criteria:	Discrepancy between name on tube and name on form, unlabeled specimen; hemolytic;
	lipemic; gross bacterial contamination.
Availability:	Monday to Friday. Test available only to DHMH epidemiologists for outbreak
	investigations. Prior approval by DHMH Epidemiology (410-767-6628) required.
Results and Interpretation:	Negative: No detectable Varicella IgM antibodies. A negative result indicates no current
	infection with Varicella virus. However, specimens taken too early during a primary
	infection may not have detectable levels of IgM antibody. If a primary infection is
	suspected, another specimen should be taken within 7 days and tested concurrently in the
	same assay with the original specimen to look for seroconversion
	Equivocal: Equivocal specimens are borderline. Another specimen should be collected
	after 7 days and retested.
	Positive: Indicates evidence of Varicella IgM antibodies.
	This suggests primary or reactivated infection with Varicella.
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/varicella.pdf
Purpose of Test:	For detection of IgM antibodies to Varicella virus. Test available only to DHMH
•	epidemiologists for outbreak investigations. Prior approval by DHMH Epidemiology
	410-767-6628) required.
Method:	ELISA
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
Testing Site:	DHMH Laboratories Administration, Central Laboratory
5	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	LIMITATIONS: Results of the Varicella IgM ELISA are not by themselves diagnostic and
	should be interpreted in light of the patient's clinical condition and results of other
	diagnostic procedures. Varicella virus specific IgG antibody may compete with IgM for
	binding sites and cause false negative results. Rheumatoid factor, if present along with
	specific IgG, will cause false positive results. The Sample Diluent contains an absorbent
	which will remove IgG from the test specimen, and significantly reduce the possibility of
	false positive or negative results. Heterotypic IgM antibody responses may occur in patients
	infected with Epstein-Barr virus, and sera from patients with infectious mononucleosis.
	Samples taken too early during the course of a primary infection may not have detectable
	levels of Varicella specific IgM. A negative result does not rule out a primary infection with
	rubella virus. It cannot distinguish the difference between vaccine-induced antibody and
	antibody resulting from a natural infection. False positive anti-Varicella IgM results may be
	obtained from patients with autoimmune disease. The performance of the Varicella IgM
	ELISA has not been validated using neonatal samples.
	Leist has not seen valiaated asing neonatal samples.

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TEST:	Varicella Zoster Virus (VZV) Viral Culture
Synonym:	Varicella Zoster Virus (VZV) culture: refer to instructions for Virus Culture.
Laboratory/Phone:	Virology: 443-681-3934

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 Vibrio culture

 Synonym:
 Vibrio spp. Enteric Culture: Refer to instructions for Enteric Culture, Routine (Salmonella, Shigella, Campylobacter, and Shiga toxins-producing *E. coli*).

 Laboratory/Phone:
 Microbiology-Enterics 443-681-4570

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TEST:	Vibrio parahaemolyticus culture
Synonym:	Vibrio spp. Enteric Culture: Refer to instructions for Enteric Culture, Routine (Salmonella,
	Shigella, Campylobacter, and Shiga toxins-producing E. coli).
Laboratory/Phone:	Microbiology-Enterics 443-681-4570

TEST:	Virus Culture
Synonym:	Viral Culture, Virus isolation for: Adenovirus, Cytomegalovirus (CMV), Enterovirus (including Echovirus, Coxsackie, and Polio), Herpes Simplex Virus (HSV Types 1 & 2), Influenza (Types A & B), Measles, Mumps, Parainfluenza (Types 1,2 & 3), Respiratory Syncytial Virus (RSV), Varicella Zoster Virus (VZV)
Laboratory/Phone:	Virology: 443-681-3934
Turnaround Time:	3-28 business days
Specimen Required:	One specimen per test requested, collected during the acute phase of the disease: blood, cerebrospinal fluid (CSF), skin lesion, eye, genital, mucosal, oral, upper and lower respiratory tract, stool, tissue/biopsy, urine
Specimen identification:	Specify the source of the specimen. Label container with patient's last name, first name, DOB, specimen type, date and time of collection.
Specimen Volume (Optimum):	Fluid: ≥ 1 ml Swab/tissue in viral transport media (VTM) Unpreserved fresh stool: 4 grams in sterile container
Specimen Volume (Minimum):	
	Continued Next Page>

Collect:	Specimen	Collect	Container
	CSF	Collect <u>&gt;</u> 2 ml aseptically.	Sterile container with leak-proof screw top lid.
	Eye	Collect aseptically and leave swab in VTM.	Viral transport media (VTM)
	Nasopharyngeal aspirate	Aspirate using #8 French catheter and trap	Sterile container with leak-proof screw top lid.
	Oral	Swab inner side of both cheeks behind upper molars and floor of mouth, including any ulcerated areas. Leave swab in VTM.	Viral transport media (VTM)
	Buccal	Swab inner side of both cheeks. Leave swab in VTM	Viral transport media (VTM) Notify DHMH Epidemiology and send to laboratory ASAP after collection.
	Rectal	Insert swab at least 5 cm into orifice and rotate the swab. Leave swab in VTM.	Viral transport media (VTM)
	Stool	4-8 grams	Sterile container with leak-proof screw top lid.
	Throat	Swab tonsillar area and back of pharynx. Leave swab in VTM.	Viral transport media (VTM)
	Tissue	Collect biopsy and autopsy specimens aseptically	Sterile container with leak-proof screw top lid. If possible, add viral transport media.
	Urine	Clean catch, midstream urine	Sterile container with leak-proof screw top lid. For recovery of CMV, send to lab within 2-3 hours after collection on cold ice packs. DO NOT FREEZE!
	Urine for mumps	Collect entire stream.	Sterile container with leak-proof screw top lid. Notify DHMH
			Epidemiology and send to laboratory ASAP after collection.
Form:	Indicate the spec		ASAP after collection.
	Indicate the spec test. Provide clin collection date. Specimens must b conditions of tran pages 9 & 10 for t	ific virus suspected by placing a "S nical history, age of patient, relevan be packaged in a triple packaging sy	ASAP after collection. tion (Order Forms: 443-681-3777) pecimen Code' in the box next to the nt vaccination history, and specimen rstem to ensure that under normal ured or leak their contents (Refer to
Form: Packaging and Shipping*: Transport Conditions:	Indicate the spec test. Provide clin collection date. Specimens must b conditions of tran pages 9 & 10 for t *Refer to current For	ific virus suspected by placing a "S nical history, age of patient, relevan be packaged in a triple packaging sy nsport they cannot break, be punctu triple packing guidance). ederal regulations for specific shipping for enterovirus (Polio, Coxsackie, ar	tion (Order Forms: 443-681-3777) pecimen Code' in the box next to the nt vaccination history, and specimen rstem to ensure that under normal ured or leak their contents (Refer to requirements.
Packaging and Shipping*:	Indicate the spec test. Provide clin collection date. Specimens must b conditions of tran pages 9 & 10 for t *Refer to current F4 Stool specimens f refrigerated cold Specimens for CM after collection (v Zoster Virus, Influ Virus, and HSV cu specimen for viru	ific virus suspected by placing a "S nical history, age of patient, relevan be packaged in a triple packaging sy hsport they cannot break, be punctu triple packing guidance). ederal regulations for specific shipping for enterovirus (Polio, Coxsackie, ar packs. AV cultures should be delivered refu within 2-3 hours). DO NOT FREEZE s uenza, Parainfluenza, Adenovirus, M iltures should be shipped on cold pa is isolation other than those previou	ASAP after collection. tion (Order Forms: 443-681-3777) pecimen Code' in the box next to the nt vaccination history, and specimen restem to ensure that under normal ured or leak their contents (Refer to requirements. Id Echovirus) should be shipped on rigerated on cold packs immediately pecimens for CMV culture. Varicella- leasles, Mumps, Respiratory Syncytial acks or kept frozen using dry ice. Any usly listed should be shipped frozen in a
Packaging and Shipping*:	Indicate the spec test. Provide clin collection date. Specimens must b conditions of tran pages 9 & 10 for t *Refer to current F4 Stool specimens f refrigerated cold Specimens for CN after collection (v Zoster Virus, Influ Virus, and HSV cu specimen for viru dry ice outfit. Sea vapors. Whenever possib isolation tests are	ific virus suspected by placing a "S nical history, age of patient, relevan be packaged in a triple packaging sy hsport they cannot break, be punctu- triple packing guidance). ederal regulations for specific shipping for enterovirus (Polio, Coxsackie, ar packs. AV cultures should be delivered refu- within 2-3 hours). DO NOT FREEZE s uenza, Parainfluenza, Adenovirus, M iltures should be shipped on cold pa- is isolation other than those previou al the specimen container tightly to ele, submit both acute and convales be being requested.	ASAP after collection. tion (Order Forms: 443-681-3777) pecimen Code' in the box next to the nt vaccination history, and specimen restem to ensure that under normal ured or leak their contents (Refer to requirements. Id Echovirus) should be shipped on rigerated on cold packs immediately pecimens for CMV culture. Varicella- leasles, Mumps, Respiratory Syncytial acks or kept frozen using dry ice. Any usly listed should be shipped frozen in a prevent ingress of toxic carbon dioxide cent sera from patients for whom virus
Packaging and Shipping*:	Indicate the spec test. Provide clin collection date. Specimens must b conditions of tran pages 9 & 10 for t *Refer to current For Stool specimens for refrigerated cold Specimens for CM after collection (w Zoster Virus, Influ Virus, and HSV cu specimen for viru dry ice outfit. Sea vapors. Whenever possib isolation tests are Bacterial swab, du container, expired	ific virus suspected by placing a "S nical history, age of patient, relevan be packaged in a triple packaging sy hsport they cannot break, be puncto triple packing guidance). ederal regulations for specific shipping for enterovirus (Polio, Coxsackie, ar packs. AV cultures should be delivered refu- vithin 2-3 hours). DO NOT FREEZE s ienza, Parainfluenza, Adenovirus, M iltures should be shipped on cold pa- is isolation other than those previou al the specimen container tightly to hele, submit both acute and convales be being requested. ry swab, swab with wooden shaft, of d transport media, unlabeled specim	ASAP after collection. tion (Order Forms: 443-681-3777) pecimen Code' in the box next to the nt vaccination history, and specimen restem to ensure that under normal ured or leak their contents (Refer to requirements. Id Echovirus) should be shipped on rigerated on cold packs immediately pecimens for CMV culture. Varicella- leasles, Mumps, Respiratory Syncytial acks or kept frozen using dry ice. Any usly listed should be shipped frozen in a prevent ingress of toxic carbon dioxide cent sera from patients for whom virus ralcium alginate swab, leaking nen, mismatch between labeling of boom temperature more than 2 hours,

Results and Interpretation:	Positive: (Name of virus) isolated.
	Negative: No viruses isolated.
Additional Information:	
Purpose of Test:	Virus isolation to determine probable cause of infection and aid in the diagnosis of viral
	disease or to further characterization for epidemiological purposes.
Method:	Cell culture, viruses detected by cytopathic effect and/or antibody/fluorescent staining.
Interfering Substances:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	The two most important steps in viral isolation are specimen collection and specimen transportation. Since the detection of viruses is more likely to be achieved early in the illness, specimens for most viral diseases should be collected as soon as a viral infection is suspected and submitted to the laboratory as soon as possible.
	Submission of adequate specimen and patient history is essential. A blanket request for "Virus Study" should not be submitted. Information must specify the group of viruses suspected. Please indicate suspected infecting agent as well as additional information such as chief symptoms, clinical test results, epidemiology data, immunizations, etc. This will guide the laboratory in choosing which virological procedures and host systems should be inoculated. Since many viruses die rapidly once they have been separated from host tissue, specimens must be delivered to the Virology Laboratory immediately after collection.
	Isolation of a virus from clinical material does not establish an etiologic diagnosis per se. The significance of such a virus depends upon the source of the isolate. For example, isolation of a virus from the brain in encephalitis or from the spinal fluid in aseptic meningitis provides direct evidence of an etiological association. Likewise isolation of an influenza virus from throat washings of a patient ill with an influenza-like disease strongly suggests that the virus is the causative agent since this virus is only isolated from throat washings in acute influenza. In contrast, the isolation of an enteric virus from the stool of a patient suffering from aseptic meningitis does not by itself indicate an etiological relationship, as enteroviruses are sometimes found in the feces of healthy individuals. Occasionally a virus other than the one ordered is detected since any reaction in the host system is investigated.
	A negative viral culture report does not preclude the possibility of the suspect virus or another virus being involved in the patient's disease. The cultures may be negative because of specimen procurement problems, such as prolonged transportation or processing delays, procurement of sample too late in the course of the disease, or inability of some viruses or viral strains to adapt to growth in the tissue culture cell lines selected. For a more rapid diagnosis, Real-Time PCR detection tests for Influenza A virus, Influenza B virus, and Herpes simplex virus I and II are available.

Vancomycin-Resistant <i>Enterococcus</i> culture; rule out Vancomycin-Resistant <i>Enterococcus faecium</i> ; rule out Vancomycin-Resistant <i>Enterococcus faecalis</i>
Microbiology 443-681-3952
2-3 days
Rectal swab; perianal swab, stool
Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection. The specimen/sample must be properly labeled and match the test requisition or electronic test order.
One (1) swab
N/A
Culturette tube with transport medium
DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777) Indicate specimen type using the "Specimen Code" on form.
Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance). *Refer to current Federal regulations for specific shipping requirements.

Transport Conditions:	Store and ship at room temperature, ship as quickly as possible.	
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results and to avoid misleading information that might lead to misdiagnosis and inappropriate therapy. A request for a new specimen will provide appropriate materials and clinically relevant information to support good patient care. Unlabeled or improperly labeled specimen Non-sterile or leaking container Inappropriate specimen transport conditions Illegible, or no submitter information on the request form Mismatched form and specimen Broken specimen/sample container The wrong specimen for test request Inappropriate outfit for requested test Illegible or no patient information on the specimen Expired transport media Specimen received after prolonged delay (usually more than 72 hours)	
Availability:	Monday through Friday	
Results and Interpretation:	VRE isolated and identified, Vancomycin resistance confirmed.	
Reference Range:	No VRE detected	
Additional Information:	N/A	
Purpose of Test:	Detect the presence of VRE	
Method:	N/A	
Interfering Substances:	N/A	
Testing Site:	DHMH Laboratories Administration, Central Laboratory	
-	1770 Ashland Avenue, Baltimore, Maryland 21205	
Comment:	N/A	

TEST:	West Nile Virus IgM Equine EIA (Equine specimen)
Synonym:	Arthropod-borne virus: WNV (West Nile Virus)
Laboratory/Phone:	Virology: 443-681-3937
Turnaround Time:	7 business days
Specimen Required:	Serum (blood);CSF
Specimen identification:	Label container with horse's name, specimen type, date and time of collection.
Specimen Volume (Optimum):	2 ml serum; 2ml CSF
Specimen Volume (Minimum):	1 ml serum; 0.5 ml CSF
Collect:	Red top vacuum tube, transfer serum to sterile tube: CSF in sterile container with leak- proof cap.
Request Form:	Equine Arbovirus Testing Form
	[Order: 443-681-3776]
	For testing to be initiated, the ANIMAL INFORMATION box on the form must be filled out
	completely.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48
	hours, CSF must be frozen at -20°C and shipped on dry ice.
Specimen Rejection Criteria:	Grossly hemolyzed specimens, unlabeled specimen, leaking container, duplicate specimen
	type (e.g., two serum specimens collected on the same day-one tube will not be tested),
	and mismatch between labeling of specimen and test request form.
Availability:	Monday through Friday.
Results and Interpretation:	IgM: Negative, High Background, Equivocal, Positive
	Serum and CSF samples that tests positive for IgM is consistent with acute WNV infection
Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name given to
	viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc). For more
	information, see the CDC link at:
	http://www.cdc.gov/ncidod/dvbid/arbor/arbdet.htm
Purpose of Test:	
Method:	ELISA
Interfering Substances:	
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Testing Site:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	

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Arthropod-borne virus: Western Equine Encephalitis (WEE)	
Virology: 443-681-3937	
3 weeks	
Serum (blood)	
The specimen/sample must be properly labeled and include:	
1. The patient's name or unique patient/sample identifier matching the test	
requisition or electronic test order,	
2. If appropriate, the date and time of specimen/sample collection, and	
3. Any additional information relevant and necessary for the test.	
2 ml serum	
1 ml serum	
Red top vacutainer tube, transfer serum to sterile tube	
DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)	
Indicate specimen type using the "Specimen Code" on form. Write "S" for serum in the	
"Other Tests Request" and indicate Western Equine Enchephalitis.	
For testing to be initiated the following information MUST be provided: date of onset,	
date specimen collected, travel history, and flavivirus vaccination history. Also please	
provide: patient's date of birth, diagnosis, symptoms, fatality, and whether patient is	
immunocompromised.	
Specimens must be packaged in a triple packaging system to ensure that under normal	
conditions of transport they cannot break, be punctured or leak their contents (Refer to	
pages 9 & 10 for triple packing guidance).	
*Refer to current Federal regulations for specific shipping requirements.	
Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48	
hours, specimen can be frozen at -20°C and shipped on dry ice.	
Grossly hemolyzed specimen, unlabeled specimen, leaking container, mismatch between	
labeling of specimen and test request form/electronic test order, and does not meet	
epidemiological criteria required for testing (e.g. travel history, etc.)	
Specimens shipped to the CDC Monday-Wednesday.	
Serum that tests positive for IgM and negative for IgG is consistent with acute Western	
Equine Encephalitis infection. A positive Western Equine Encephalitis EIA is confirmed by	
PRNT (plaque reduction neutralization). A positive IgG antibody and a negative IgM	
antibody are consistent with infection in the distant past and are not consistent with acute infection.	
The term "Arbovirus" has no taxonomic significance, but is a shortened name give to	
viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).	
Arboviruses that cause human encephalitis are members of three virus families: The	
Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae. For more information, see	
the CDC link at: https://www.cdc.gov/ncezid/dvbd/	
Patients with travel history supporting suspicion of other arboviruses will be sent to the	
CDC for testing.	
For the presumptive detection of antibodies to Western Equine Encephalitis Virus.	
Confirmatory testing by PRNT may be required.	
EIA (Screening) & PRNT (Plaque Reduction Neutralization Test) referral to the Centers for	
Disease Control and Prevention (CDC).	
DHMH Laboratories Administration, Central Laboratory	
1770 Ashland Avenue, Baltimore, Maryland 21205	
Other Arboviral testing not available at the state lab will be forwarded to the CDC based on	
patient's travel history and onset date.	

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TEST:	Whooping Cough
Synonym:	B. pertussis, pertussis, Whooping Cough Refer to instructions for Bordetella pertussis PCR
	and Culture.
Laboratory/Phone:	Molecular Biology: 443-681-3924; Microbiology 443-681-3952

TEST:	Woolsorters' Disease
Synonym:	Bacillus anthracis, Cutaneous Anthrax: Refer to instructions for Anthrax, Cutaneous
	(Woolsorters' disease).
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:
-	410-925-3121 (24/7 emergency contact number)
	Select Agents Microbiology Laboratory: 443-681-3954
	Division of Microbiology Laboratory: 443-681-3952

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TEST:	Yellow Fever (CDC Referral)
	CDC test available based on patient's travel history.
Synonym:	Arthropod-borne virus: Bunyavirus
Laboratory/Phone:	Virology: 443-681-3937
Turnaround Time:	3 weeks (CDC Referral)
Specimen Required:	Serum
Specimen identification:	The specimen/sample must be properly labeled and include:
	1. The patient's name or unique patient/sample identifier matching the test
	requisition or electronic test order,
	2. If appropriate, the date and time of specimen/sample collection, and
	3. Any additional information relevant and necessary for the test.
Specimen Volume (Optimum):	2 ml serum
Specimen Volume (Minimum):	1 ml serum
Collect:	Red top vacutainer tube, transfer serum to sterile tube
Request Form:	DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)
	Indicate specimen type using the "Specimen Code" on form.
	Write "S" for serum in the "Other Tests Request" and indicate Yellow Fever.
	For testing to be initiated, the following information MUST be provided: date of onset,
	date specimen collected, travel history, and flavivirus vaccination history. Also please
	provide: patient's date of birth, diagnosis, symptoms, fatality, and whether patient is
	immunocompromised.
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal
	conditions of transport they cannot break, be punctured or leak their contents (Refer to
	pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48
	hours, specimen can be frozen at -20°C and shipped on dry ice.
Specimen Rejection Criteria:	Grossly hemolyzed specimen, unlabeled specimen, leaking container, mismatch between
	labeling of specimen and test request form/electronic test order, and does not meet
	epidemiological criteria required for testing (e.g. travel history, etc.)
Availability:	Specimens shipped to the CDC Monday-Wednesday.
Results and Interpretation:	Serum that tests positive for IgM and negative for IgG is consistent with acute Yellow Fever
	infection. All positive Yellow Fever EIA are confirmed by PRNT (plaque reduction
	neutralization). A positive IgG antibody and a negative IgM antibody are consistent with
	infection in the distant past and are not consistent with acute infection.
Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name give to
	viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).
	Arboviruses that cause human encephalitis are members of three virus families: The
	Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae. For more information, see
	the CDC link at: <u>https://www.cdc.gov/ncezid/dvbd/</u> Patients with travel history supporting suspicion of other arboviruses will be sent to the
	CDC for testing.
Purpose of Test:	Detection of Yellow Fever Virus antibodies.
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Method:	EIA (Screening) & PRNT (Plaque Reduction Neutralization Test) referral to the Centers for
	Disease Control and Prevention (CDC).
Interfering Substances:	
Processing Site for CDC referral:	DHMH Laboratories Administration, Central Laboratory
	1770 Ashland Avenue, Baltimore, Maryland 21205
Comment:	Other Arboviral testing not available at the state lab will be forwarded to the CDC based on
	patient's travel history and onset date.

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TEST:	Yersinia culture
Synonym:	Yersinia stool culture: Refer to instructions for Enteric Culture, Routine.
Laboratory/Phone:	Microbiology-Enterics 443-681-4570

TEST:	Yersinia enterocolitica
Synonym:	Yersinia enterocolitica culture: Refer to instructions for Enteric Culture, Routine.
Laboratory/Phone:	Microbiology-Enterics 443-681-4570

TEST:	Yersinia pestis	
Synonym:	Plague	
Laboratory/Phone:	Office of Laboratory Emergency Preparedness and Response:	
	410-925-3121 (24/7 emergency contact number)	
	Select Agents Microbiology Laboratory: 443-681-3954	
	Division of Microbiology Laboratory: 443-681-3952	
Turnaround Time [from specimen	3 -6 days	
receipt in the Laboratory]:		
Specimen Required:	1. Lower respiratory tract (pneumonic): Bronchial wash or transtracheal aspirate (>1 ml).	
	Sputum may be examined but this is not advised because of contamination by normal throat flora.	
	<ol> <li>Blood (septicemia): Collect appropriate blood volume and number of sets per established laboratory protocol. NOTE: In suspected cases of plague, an additional blood or broth culture (general nutrient broth) should be incubated at room temperature (22-28°C), the temperature at which <i>Y. pestis</i> grows faster.</li> </ol>	
	3. Aspirate of involved tissue (bubonic) or biopsied specimen: Liver, spleen, bone marrow, lung. NOTE: Aspirates may yield little material; therefore, a sterile saline flush may be needed to obtain an adequate amount of specimen. Syringe and needle of aspirated sample should be capped, secured by tape, and sent to the Laboratory.	
	4. Isolate	
Specimen Identification:	Specimen should be labeled with patient's last and first name, patient's address, DOB, specimen type/source, and the date and time of collection.	
Specimen Volume (Optimum):	N/A	
Specimen Volume (Minimum):	N/A	
Collect:	1. Respiratory/sputum: Bronchial wash or transtracheal aspirate (>1.0 ml).	
	2. Blood: Collect appropriate blood volume and number of sets as per routine laboratory protocol.	
	3. Tissue aspirate/biopsy specimen: Add several drops of sterile saline to keep tissue moist.	
	4. Isolate: Pure culture, 24 hours old, growing on a sheep blood agar plate or slant.	
Form:	DHMH Form #4676 Infectious Agents: Culture/Detection (Order Forms: 443-681-3777)	
	Indicate specimen type using the "Specimen Code" on form.	
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal	
	conditions of transport they cannot break, be punctured or leak their contents (Refer to	
	pages 9 & 10 for triple packing guidance).	
	*Refer to current Federal regulations for specific shipping requirements.	
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Transport Conditions:	1. Respiratory/sputum: Transport at room temperature. If it is known that the material	
transport conditions.	will be transported from 2-24 hours after collection, then store container and	
	transport at 2-8°C.	
	<ol> <li>Blood: Transport at room temperature. Hold them at ambient temperature until they</li> </ol>	
	are incubated. DO NOT REFRIGERATE.	
	3. Tissue aspirate/biopsy specimen: Transport the sample at room temperature for	
	immediate processing. Keep the specimen chilled if processing of the specimen will be	
	delayed.	
	<ol> <li>Isolate: Transport the specimen at room temperature on a sealed sheep blood agar</li> </ol>	
	plate or slant.	
Specimen Rejection Criteria:	The following rejection criteria are designed to prevent the reporting of inaccurate results	
specifien Rejection Chiena.	and to avoid misleading information that might lead to misdiagnosis and inappropriate	
	therapy. A request for a new specimen will provide appropriate materials and clinically	
	relevant information to support good patient care.	
	<ul> <li>Unlabeled or improperly labeled specimen</li> </ul>	
	<ul> <li>Non-sterile or leaking container</li> </ul>	
	<ul> <li>Inappropriate specimen transport conditions</li> </ul>	
	<ul> <li>Illegible, or no submitter information on the request form</li> </ul>	
	<ul> <li>Mismatched form and specimen</li> </ul>	
	<ul> <li>Broken specimen/sample container</li> </ul>	
	<ul> <li>The wrong specimen for test request</li> </ul>	
	<ul> <li>Inappropriate outfit for requested test</li> </ul>	
	<ul> <li>Illegible or no patient information on the specimen</li> </ul>	
	<ul> <li>Expired transport media</li> </ul>	
Availability:	24 hours/day, 7 days/week	
Results and Interpretation:	Yersinia pestis isolated/detected	
	Yersinia pestis not found	
Additional Information:	Call 410-925-3121 before sending to the Laboratory.	
Purpose of Test:	To confirm the diagnosis of plague.	
Method:	LRN Protocols	
Interfering Substances:	N/A	
Testing Site:	DHMH Laboratories Administration, Central Laboratory	
	1770 Ashland Avenue, Baltimore, Maryland 21205	
Comment:	Call 410-925-3121 before sending to the Laboratory.	

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TEST:	Zika Virus IgM EIA	
	Test available based on patient's travel history and risk assessment.	
Synonym:	Arthropod-borne virus: Zika Virus	
Laboratory/Phone:	443-681-3937	
Turnaround Time:	5-10 business days during Arbovirus Season (excluding PRNT Testing)	
Specimen Required:	Serum (blood)	
Specimen Identification:	<ul> <li>The specimen/sample must be properly labeled and include:</li> <li>1. The patient's name or unique patient/sample identifier matching the test requisition or electronic test order,</li> <li>2. If appropriate, the date and time of specimen/sample collection, and</li> <li>3. Any additional information relevant and necessary for the test.</li> </ul>	
Specimen Volume (Optimum):	2 ml serum	
Specimen Volume (Minimum):	1 ml serum	
Collect:	Red-top vacutainer, transfer serum to sterile tube	
Form:	<ul> <li>DHMH Form# 4677 Serological Testing (Order Forms: 443-681-3777)</li> <li>Indicate specimen type using the "Specimen Code" on form. Write "S" for serum in the "Arbovirus Travel-Associated Panel".</li> <li>For testing to be initiated the following information MUST be provided: date of onset, date specimen collected, travel history, and flavivirus vaccination history. Also please provide: patient's date of birth, diagnosis, symptoms, fatality, and whether patient is immunocompromised.</li> <li>Prior authorization for testing must be obtained by Maryland State or Local Health Department Epidemiologists before testing may proceed.</li> </ul>	
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OHMH-Laboratories Administration	The J. Mehsen Joseph Public Health Laboratory
Packaging and Shipping*:	Specimens must be packaged in a triple packaging system to ensure that under normal conditions of transport they cannot break, be punctured or leak their contents (Refer to pages 9 & 10 for triple packing guidance).
	*Refer to current Federal regulations for specific shipping requirements.
Transport Conditions:	Store refrigerated and ship on cold packs in a cooler. If shipping is delayed beyond 48
transport conditions.	hours, specimen can be frozen at -20°C and shipped on dry ice.
Specimen Rejection Criteria:	Grossly hemolyzed specimen, unlabeled specimen, leaking container, mismatch between
Specimen Rejection Criteria.	labeling of specimen and test request form/electronic test order, and does not meet
	epidemiological criteria required for testing (e.g. travel history, etc.)
Availability:	Monday-Friday
Results and Interpretation:	Negative: No detectable IgM antibody to Zika virus. This result does not rule-out Zika
	virus infection. Lack of serologic evidence of infection may reflect that the specimen was collected prior to the development of an antibody response. Virus-specific IgM antibodies can be detectable equal to or greater than four days after onset of illness. Serum collected within 7 days of illness onset might not have detectable virus-specific
	IgM antibodies. It has been reported that IgM antibodies persist for approximately 2-12 weeks. Tests of a single acute-phase specimen can be inconclusive. If indicated, please submit another serum specimen collected greater than 14 days after onset of illness for further testing.
	High Background: Results are uninterpretable due to high background reactivity. Please submit a new specimen for further testing.
	Equivocal: Specimen tested equivocal for IgM antibody to Zika virus. Further testing by
	PRNT (plaque reduction neutralization test) is required.
	Positive: Specimen tested presumptively positive for IgM antibody to Zika virus. Further
	testing by PRNT (plaque reduction neutralization test) is required. A positive IgM result
	may not indicate a recent infection because IgM may persist for several months after
	infection.
Reference Range:	IgM EIA: Negative, High Background, Equivocal, Positive
Additional Information:	The term "Arbovirus" has no taxonomic significance, but is a shortened name give to viruses that are transmitted by blood feeding arthropods (mosquitoes, ticks, etc).
	Arboviruses that cause human encephalitis are members of three virus families: The
	Togaviridae (genus Alphavirus), Flaviviridae, and Bunyaviridae.
	For more information, see the CDC link at:
	http://www.cdc.gov/zika/hc-providers/index.html.
	Patients with travel history supporting suspicion of other arboviruses will be sent to the
	CDC for testing.
	Additional Arbovirus testing may be performed as indicated by travel history, symptoms,
	or other epidemiological information to include but not limited to: Dengue Virus and
	Chikungunya Virus.
Purpose of Test:	For the presumptive detection of IgM antibody to Zika Virus. Confirmatory testing by
	PRNT may be required.
Method:	EIA (Screening) & PRNT (Plaque Reduction Neutralization Test) referral to the Centers for
	Disease Control and Prevention (CDC) for confirmatory testing.
Interfering Substances:	
Testing Site:	DHMH Laboratories Administration, Central Laboratory
Testing Site:	1770 Ashland Avenue Baltimore, MD 21205

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#### E. GUIDE TO INTERPRETATION OF RETROVIROLOGY SEROLOGICAL TESTS

# RETROVIRUSES

Human Immunodeficiency Viruses (HIV)

#### NORMAL/SIGNIFICANT RESULTS

Reactive results indicate presence of HIV antigen or antibody in serum/plasma. All screening test reactive specimens undergo testing using the Geenius HIV 1/2 Supplemental Assay for differentiation of HIV-1 and HIV-2 antibodies. An In-house developed HIV-1 NAAT assay is performed on the specimens that test reactive by the HIV antigen/ antibody screening test but are not confirmed as antibody positive in the Geenius assay.

# F. GUIDE TO INTERPRETATION OF HEREDITARY DISORDERS

#### F.1.TESTS

SIGNIFICANT RESULTS

#### F.1.a. Galactosemia - GALT

1.) < 7 days old

a.) Normal	Presence of fluorescence or enzyme activity
b.) Abnormal	Absence of fluorescence or enzyme activity
2.) ≥ 7 days old	
a.) Normal	Presence of fluorescence or enzyme activity
b.) Abnormal	Absence of fluorescence or enzyme activity

### F.1.b. Galactosemia – Total Galactose

1.) < 7 days old

a.) Normal	Less than 10 mg/dL
b.) Borderline	10 - 20 mg/dL
c.) Abnormal	20, 40, 60, 80, or greater mg/dL
	>40 mg/dL with abnormal GALT or
	>80 mg/dL = neonatal emergency

### 2.) ≥ 7 days old

a.) Normal	Less than 10 mg/dL
b.) Borderline	10 - 20 mg/dL
c.) Abnormal	20, 40, 60, 80, or greater mg/dL
	>40 mg/dL with abnormal GALT or
	>80 mg/dL = neonatal emergency

### F1.c. Biotinidase

- 1.) < 7 days old
  - a.) Normal Color change indicating enzyme activity
  - b.) Abnormal Lack of color change lack of enzyme activity

## 2.) ≥ 7 days old

- a.) Normal Color change indicating enzyme activity
- b.) Abnormal Lack of color change lack of enzyme activity

DHMH-Laboratories Administration

#### F.1.d. Thyroxine

a.) Normal	≥ 6.5 μg/dL
b.) Borderline	3.0 – 6.49 μg/dL
c.) Abnormal	2.0 – 2.9 or < 2.0 μg/dL

2.) ≥ 7 days old

a.) Normal	≥ 4.0 µg/dL
b.) Borderline	3.0 – 3.9 μg/dL
c.) Abnormal	2.0 – 2.9 or < 2.0 μg/dL

# F.1.e. TSH

1.) < 7 days old

a.) Normal	≤ 20 μIU/mL
b.) Borderline	21 - 40 μIU/mL
c.) Abnormal	≥ 40 μIU/mL

# 2.) ≥ 7 days old

a.) Normal	≤ 20 μIU/mL
b.) Borderline	21 - 40 μIU/mL
c.) Abnormal	≥ 40 μIU/mL

# F.1.f. Hemoglobin

1.) < 7 days old

a.) Normal	FA hemoglobins or AF
b.) Trait	FAS, FAC, FAV, ACF, ASF, AVF, FA(C), FA(S), FACV, FASV
c.) Disease	FS, FC, FSC, F, FV, FSV, FCV, FSA, FVA

# 1.) $\geq$ 7 days old

a.) Normal	FA hemoglobins or AF
b.) Trait	FAS, FAC, FAV, ACF, ASF, AVF, FA(C), FA(S), FACV, FASV
c.) Disease	FS, FC, FSC, F, FV, FSV, FCV, FSA, FVA

Same as above

### F.1.g. Congenital Adrenal Hyperplasia

1.) < 7 days old	
a.) Normal	Varies with weight. Call laboratory at 410-767-6099

2.) ≥ 7 days old

### F.1.h. GUIDE TO INTERPRETATION OF HEREDITARY DISORDERS

ANALYTE	NORM	AL RESULT	SIGNIFICANT RESULT		
	0-6 DAYS OLD	≥ 7 DAYS OLD	0-6 DAYS OLD	> 7 DAYS OLD	
Arginine	≤ 70 μM	≤ 80 µM	> 70 μM	> 80 µM	
Citrulline	≤ 40 μM	≤ 70 μM	> 40 μM	> 70 μM	
Valine	≤ 400 μM	≤ 400 μM	NA	NA	
Leucine	≤ 275 μM	≤ 300 μM	> 275 μM	> 300 μM	
Methionine	≤ 75 μM	≤ 80 µM	> 75 μM	> 80 µM	
Phenylalanine	≤ 120 μM	≤ 220 μM	> 150 μM	> 150 μM	
Tyrosine	≤ 300 μM	≤ 300 μM	> 300 μM	> 300 μM	
Acylcarnitine Profile (for 11 Organic Acidemias and 9 Fatty Acid Oxidation Disorders)	Contact Newborn Screening	Contact Newborn Screening	Contact Newborn Screening	Contact Newborn Screening	

F.2. CLINICAL AND HEMOTOLOGIC ASPECTS OF SOME HEMOGLOBINOPATHIES

TRAIT 1	HB TYPES	CLINICAL SEVERITY	RED-CELL MORPHOLOGY	ANEMIA	SICKLING
Hb-S trait	A + S	+	+ Normal		+
Hb-C trait	A + C	-	Normal	-	-
Hb-E trait	A + E	-	Normal	+	-
DISEASE 2	HB TYPES	CLINICAL SEVERITY	RED-CELL MORPHOLOGY	ANEMIA	SICKLING
Homozygous					
Sickle cell anemia	S + S	+ + +	Normocytic Normochromic	+ + +	+
HbC disease	C + C	+	+ Slightly microcytic		_
HbD disease	D + D	-	Microcrytic normochromic	-	-
HbE disease	E + E	+	+ Microcytic normochromic		-
Mixed Heterozygous					
Sickle Cell HbC Disease	C + S (F*)	- to + + +	Slightly microcytic, slightly hypochromic	- to + + +	+

Sickle Cell HbD Disease	D + S (F*)	+ +		+ + +	+
Thalassemia Syndrome					
Thalassemia major	A + F	++++	Microcytic hypochromic	+ + + +	-
Thalassemia HbS Disease	S + F + A	+ to + + + +	Microcytic hypochromic	+ + to + + + +	+
Thalassemia HbC Disease	A + C (F*)	+ to + +	Microcytic hypochromic	- to	-
Thalassemia HbE Disease	E + F	+ to + + + +	Microcytic hypochromic	+ to + + + +	-

References (to "Clinical and Hemotologic Aspects of Some Hemoglobinopathies")

<sup>1</sup> Nurembgerg, S.T. Electrophoreseis, F. A. David Co. Philadelphia. 1966. p. 127

<sup>2</sup> Modified from Chernoff (1958)

\* F may be present

# F.3. COMPARISON OF IRON-DEFICIENCY ANEMIA AND THALASSEMIA

PARAMETER	IRON-DEFICIENCY ANEMIA	BETA-THALASSEMIA MINOR
RBC	decreased	normal to increased
Hemoglobin	decreased	decreased
Hematocrit	decreased	decreased
Mean Corpuscular Volume (MCV) and Mean Corpuscular Hemoglobin (MCH)	decreased	decreased
Mean Corpuscular Hemoglobin Concentration (MCHC)	decreased	normal
Serum Iron	decreased	normal to increased
Total iron Binding Capacity (TIBC)	decreased	normal to increased
Response to parenteral iron administration	very rapid	negligible

### G. COMMON VIRAL AND RICKETTSIAL CLINICAL SYNDROMES

As a guide to the physician in submitting specimens for viral and rickettsial studies, the following chart has been included. It lists the common clinical syndromes, viruses which have been associated with each, and the clinical materials which should be collected. Every attempt should be made to obtain all of the materials listed for each illness, since this will greatly increase the chances of the laboratory in establishing an etiologic diagnosis.

	MANIFESTATION	AGENT	SOURCE OF SPECIMEN		
IVI	ANIFESTATION	AGENT	CLINICAL AUTOPSY		
G.	1. CARDIOVASCULAR				
	a. Myocarditis and Pericarditis	Enteroviruses: (including Coxsackie A), (types 4, 14, 16) B-1 – B-5	Throat swab/washing Feces Pericardial fluid	Blood Pericardial fluid	

	IFESTATION AGENT			
MANIFESTATION	AGENT	CLINICAL	AUTOPSY	
G.2. CENTRAL NERVOUS SYST	EM (CNS)	·		
a. Paralysis	Enteroviruses: Polioviruses types 1,2,3 Coxsackie A-7, A-9 ECHO types 2 and 9	Throat swab/washing CSF Feces	Brain Intestinal contents	
b. Aseptic meningitis and/or encephalitis	Enteroviruses: Poliovirus Coxsackie Group A and B ECHO viruses Herpes simplex	Throat swab/washing CSF Feces	Brain Intestinal contents	
		Mouth swab CSF	Brain	
	Mumps	Mouth swab of Swenson's ducts CSF Urine	Brain Parotid	
	Arboviruses	Blood Throat CSF	Brain	
	Lymphocytic choriomeningitis	Blood CSF	Brain	
	Lymphogranuloma venereum	CSF Primary Lesion site	Brain Liver Spleen	
	Rabies	See CDC Rabies Guidelines	See CDC Rabies Guidelines	

	Adenoviruses	Throat swab CSF Feces	Brain
	Measles (Rubeola)	Blood CSF	Brain
c. Guillain-Barré Syndrome	Coxsackie A ECHO viruses	Throat swab/washing CSF Feces	Brain cord
d. Subacute sclerosing Pan encephalitis (Dawson's encephalitis)	Measles (Rubeola)	CSF Blood	Brain

MANIFESTATION		AGENT	SOURCE OF SPECIMEN	
/IAN	IFESTATION	AGENT	CLINICAL	
i.3. E	EXANTHEMATOUS INFECTION			·
a.	Skin and Mucous Membrane			
	(1.) Smallpox	Vaccinia variola	Crusts	Liver
	(2.) Chickenpox	Varicella zoster	Throat swab/washing Vesicle fluid Scrapings from vesicle base	Spleen (Lung also for varicella)
	(3.) Fever blisters	Herpes simplex	Mouth swab Vesicle fluid and scrapings	CNS
	(4.) Herpangina	Enterovirus: Coxsackie A	Vesicle fluid Throat swab/washing Feces Vaginal swab	
	(5.) Hand, foot and mouth disease	Enterovirus Coxsackie A	Vesicle fluid Throat swab/washing (types 5, 10, 16)	Feces
	(6.) Dengue fever	Dengue virus (types 1-4)	Blood	Blood
b	. Maculopapular Rash			

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(1.) Enterovirus		Throat swab/washing Feces	
(2.) German measles	Rubella	Heparinized blood CSF Products of conception Throat swab/washing Urine	Lung Liver Spleen

	ACENT	SOURCE OF SPECIMEN			
MANIFESTATION	AGENT	CLINICAL	AUTOPSY		
G4. OCULAR (OPHTHALMIC DISEASE)					
a. Kerato-conjunctivitis	Adenoviruses (types 8, 19, and 37)	Eye swab	Throat swab/washing		
b. Ocular Herpes	Herpes Simplex	Eye swab	CNS		
c. Follicular Conjunctivitis	Adenoviruses (types 3, 7, and others)	Eye swab	Throat swab/washing Eye swab		
d. Conjunctivitis	New Castle Disease Virus		Conjunctival scrapings		
G.5. RESPIRATORY INFECTION					
a. Lower Tract					
(1.) Bronchitis Laryngotracheo bronchitis (Croup)	Influenza Parainfluenza Respiratory syncytial virus (infants)	Nasopharyngeal Aspirate Sputum	Lung Bronchial scrapings (for influenza, add spleen, liver, and/or kidney)		
	Chlamydia	Sputum Pleural fluid Throat swab/washing	Lung Liver Spleen		
	Adenoviruses	Sputum Nasopharyngeal Aspirate Feces	Lung Bronchial scrapings		
	Enteroviruses	Throat swab/washing Feces	Intestinal contents		
G.6. RICKETTSIAL INFECTIONS					
a. Rocky Mountain Spotted Fever	Rickettsia rickettsii	Blood	Liver Spleen		
b. Ehrlichiosis	Ehrlichia chaffeensis	Blood			
c. Epidemic typhus	Rickettsia prowazekii	Blood			

d. Murine typhus	Rickettsia typhi	Blood	
e. Q Fever	Coxiella burnetii	Sputum Urine CSF Blood	Liver Spleen
f. Rickettsial pox	Rickettsia akari	Blood	Liver Spleen

	ACENIT	SOURCE OF SPECIMEN			
MANIFESTATION	AGENT	CLINICAL	AUTOPSY		
G.7. SEXUALLY TRANSMITTED DISEASES (STD)					
a. Acquired Immuo-Deficiency	Human Immuno-Deficiency virus	Whole blood			
Syndrome (AIDS)	HIV1, HIV2				
b. Genitourinary tract infection	Herpes Simplex 2	Lesion scraping			
		Vaginal swab			
c. Vulvovaginitis	Coxsackie B	Vaginal swab			
	Herpes Simplex 2	Lesion scraping			
d. Lymphogranuloma	Chlamydia trachomatis	Fluid and pus			
venereum, cervicitis,		Cervical swab			
urethritis		Urethral swab			
		Rectal swab			
G.8. SYSTEMIC					
	Cytomegalovirus	Urine, Saliva	Kidney		
		Throat	Lung		
		swab/washing	Liver		
		Heparinized blood CSF	Brain		
		Lung Biopsy			
	Adenoviruses	Throat	Intestinal		
		swab/washing Sputum	contents		
		Feces	Lung		
		Urine	Brain		
		CSF	Liver		
			Kidney Heart		
	Coxsackie B	Throat	Brain		
		swab/washing	Heart		
		CSF	Lymph node		
		Feces, pleural, or as	Intestinal		
		indicated			

		SOURCE OF SPECIMEN	
IANIFESTATION	AGENT	CLINICAL	AUTOPSY
.9. MISCELLANEOUS			1
a. Infantile diarrhea	Coxsackie A (types 18, 20, 21, 22, 24)	Feces	
b. Hepatitis	Enteroviruses (including Coxsackie A) (types 4, 9)	Throat swab/washing Feces Live	Intestinal contents
c. Hemolytic-uremic Syndrome	Coxsackie A (type 4)	Throat swab/washing Feces	Lung Kidney Intestinal contents
d. T cell leukemia	HTLV I, II	Heparinized blood	
e. Gastroenteritis	ECHO Coxsackie B Rotaviruses Norovirus	Feces Throat swab/washing Vomitus	
f. Orchitis and Epididymitis	Mumps Coxsackie	Urine Throat swab/washing Feces	Parotid
g. Intussusception	Adenovirus	Feces Mesenteric lymph node	
h. Colorado Tick Fever	CTF virus	Blood	
i. Acute Infectious Lymphocytosis	Epstein-Barr virus (EB) Coxsackie-like virus	Blood	
j. Post Perfusion Syndrome	Cytomegalovirus Epstein-Barr virus	Blood	

# H. DIRECTORY OF LOCAL HEALTH DEPARTMENTS

HEALTH DEPARTMENT	ADDRESS	TELEPHONE	EMERGENCY/ AFTER HOURS PHONE#	FAX NO.
Allegany	P.O. Box 1745 12501-12503 Willowbrook Rd. Cumberland MD 21501-1745	301-759-5000	301-759-3060	301-777-5674
Anne Arundel	Health Services Buildings 3 Harry S. Truman Parkway Annapolis MD 21401	410-222-7375	410-222-7095	410-222-4436

			EMERGENCY/	
HEALTH DEPARTMENT	ADDRESS	TELEPHONE	AFTER HOURS PHONE#	FAX NO.
Baltimore City	1001 East Fayette Street Baltimore MD 21202	410-396-4387	410-396-3100	410-396-1617
Baltimore County	Drumcastle Government Center 6401 York Road, 3rd Floor Baltimore MD 21212	410-887-2243	410-832-7182	410-377-5397
Calvert	P.O. Box 980 975 Solomons Island Rd Prince Frederick MD 20678	410-535-5400	443-532-5973	410-535-5285
Caroline	403 South 7th Street Denton MD 21629	410-479-8030	Comm. Disease 443-786-1398 Rabies 410-479-2232	410-479-0554
Carroll	290 S. Center Street Westminister MD 21157	410-876-2152	410-386-2260	410-876-4988
Cecil	John M. Byers Health Center 401 Bow Street Elkton MD 21921	410-996-5550	410-996-5550	410-996-5179
Charles	4545 Crain Highway White Plains MD 20695-1050 Mailing Address: P.O. Box 1050 White Plains MD 20695	301-609-6900	301-932-2222	301-934-4632
Dorchester	3 Cedar Street Cambridge MD 21613	410-228-3223	410-228-3223	410-228-9319
Frederick	350 Montevue Lane Frederick MD 21702	301-600-1029	301-600-0311	301-600-3111
Garrett	1025 Memorial Drive Oakland MD 21550	301-334-7777	301-334-1930	301-334-7771
Harford	120 South Hays Street P.O. Box 797 Bel Air MD 21014-0797	410-838-1500	Comm. Disease 443-243-5726 Environ. Health 410-638-3400	410-638-4952
Howard	8930 Stanford Boulevard Columbia, MD 21045	410-313-1412	410-313-2929	410-313-6108
Kent	125 S. Lynchburg Street Chestertown MD 21620	410-778-1350	Comm. Disease 410-708-5611 Environ. Health 410-778-1371	410-778-7913
Montgomery	401 Hungerford Drive, 5th Floor Rockville MD 20850	240-777-1741	240-777-4000	301-279-1692
Prince George's	1701 McCormick Drive Largo MD 20774	301-883-7834 301-883-7879	301-883-4748 301-883-7879	301-883-7896

HEALTH DEPARTMENT	ADDRESS	PHONE#	EMERGENCY PHONE#	FAX NO.
Queen Anne's	206 N. Commerce Street Centreville MD 21617	410-758-0720	410-758-3476 410-778-5173	410-758-2838
Somerset	7920 Crisfield Highway Westover MD 21871	443-523-1700	443-523-1750	410-651-5680
St. Mary's	21580 Peabody Street, P.O. Box 316 Leonardtown MD 20650	301-475-4330	301-475-8016	301-475-4350
Talbot	100 S. Hanson Street Easton MD 21601	410-819-5600	410-822-0095	410-819-5690
Washington	1302 Pennsylvania Avenue Hagerstown MD 21742	240-313-3260	301-573-6375	240-313-3201
Wicomico	108 East Main Street Salisbury MD 21801	410-543-6930	410-543-6996	410-543-6975
Worcester	P.O. Box 249 6040 Public Landing RD. Snow Hill MD 21863	410-632-1100	410-632-1311	410-632-0906

# I. ACRONYMS

AFB	acid fast bacillus
AFP	alpha fetoprotein
Ag	Antigen
ВСК	branch chain ketoacids
САН	congenital adrenal hyperplasia
CF	complement fixation
СНЅ	Childhood Screening
CMV	Cytomegalovirus
CSF	cerebrospinal fluid
DF	dark field
DFA	direct fluorescent antibody
DHMH	Department of Health and Mental Hygiene
EBNA	Epstein Barr virus nuclear antigen
EBV	Epstein Barr virus
EEE	Eastern Equine Encephalitis
EIA	enzyme linked immunosorbent assay
ELISA	enzyme linked immunosorbent assay
GALT	Galactose 1-phosphate uridyl transferase
HAVAb	Hepatitis A virus antibody
Hb	Hemoglobin

HIV	Human Immunodeficiency virus
HSV	Herpes Simplex virus
HTLV I/II	Human T Lymphocytic virus
, IFA	indirect fluorescent antibody
IFA	Immunofluorescent antibody
lgG	Immunoglobulin G
lgM	Immunoglobulin M
IHA	indirect hemagglutination
ІМ	infectious mononucleosis
LCM	lymphocytic choriomeningitis
LIMS	Lab Information Management System
LT	lavender top tube
MAC	Mycobacterium avium complex
MCAD	medium chain acyl-dehydrogenase deficiency
МСНС	mean corpuscular hemoglobin concentration
mg/dL	milligram per deciliter
NBS	Newborn Screening
NP	nasopharyngeal
PCR	polymerase chain reaction
PFGE	pulsed-field gel electrophoresis
PKU	phenylketonuria
RFFIT	rapid fluorescent focus inhibition technique
RPR	rapid plasma reagin
RSV	Respiratory Syncytial virus
RT	red top tube
RT-PCR	Reverse-transcribed polymerase chain reaction
SPS	yellow blood collection tubes containing sodium polyanethol sulfonate
TIBC	total iron binding capacity
VIR-IMM	Virology Immunology Division
VCA	viral capsid antigen
VTM	viral transport media
VZV	Varicella-Zoster virus
WB	Western Blot
WEE	Western Equine encephalitis