

TRAINING UPDATE

Lab Location: SGMC & WOMC
Department: Core Lab

Date Distributed: 2/25/2020
Due Date: 3/25/2020
Implementation: 3/10/2020

DESCRIPTION OF PROCEDURE REVISION

| | |
|---|---|
| Name of procedure: | |
| Malaria SGAH.M0 v6 | |
| Description of change(s): | |
| | |
| Section | Reason |
| Header | Change WAH to WOMC |
| 9 | Added minimum parasitemia level of <0.01% |
| This revised SOP will be implemented on March 10, 2020 | |

Document your compliance with this training update by taking the quiz in the MTS system.

Technical SOP

| | | |
|--------------------|----------------|-----------------|
| Title | Malaria | |
| Prepared by | Ron Master | Date: 5/11/2009 |
| Owner | Ron Master | Date: 5/11/2009 |

| | | |
|--|------------------|------------------------------|
| Laboratory Approval | | Local Effective Date: |
| Print Name and Title | Signature | Date |
| <i>Refer to the electronic signature page for approval and approval dates.</i> | | |

TABLE OF CONTENTS

| | | |
|-----|--|----|
| 1. | Test Information..... | 2 |
| 2. | Analytical Principle..... | 2 |
| 3. | Specimen Requirements..... | 2 |
| 4. | Reagents..... | 4 |
| 5. | Calibrators/Standards..... | 4 |
| 6. | Quality Control..... | 4 |
| 7. | Equipment And Supplies..... | 5 |
| 8. | Procedure..... | 6 |
| 9. | Calculations..... | 7 |
| 10. | Reporting Results And Repeat Criteria..... | 7 |
| 11. | Expected Values..... | 21 |
| 12. | Clinical Significance..... | 21 |
| 13. | Procedure Notes..... | 22 |
| 14. | Limitations Of Method..... | 23 |
| 15. | Safety..... | 23 |
| 16. | Related Documents..... | 23 |
| 17. | References..... | 23 |
| 18. | Revision History..... | 24 |
| 19. | Addenda..... | 25 |

1. TEST INFORMATION

| Assay | Method/Instrument | Test Code |
|--------------------------------|-------------------|-----------|
| Malaria, thick and thin smears | Manual | MAL |

| |
|---|
| Synonyms/Abbreviations |
| Malaria smear, Malaria ID, Malaria Parasites, <i>Plasmodium</i> species |

| |
|-------------------|
| Department |
| Microbiology |

2. ANALYTICAL PRINCIPLE

Examination of stained peripheral blood smears is used for screening and identifying malarial parasites, *Babesia*, trypanosomes, and microfilaria. Malarial and *Babesia* parasites infect circulating red cells and undergo various stages of development within the red cell. The Wright Giemsa stain highlights morphologic features of these stages.

3. SPECIMEN REQUIREMENTS

3.1 Patient Preparation

| Component | Special Notations |
|--|--|
| Fasting/Special Diets | None |
| Specimen Collection and/or Timing | <p>Slides are to be prepared when the patient presents with symptoms of malaria, and every 6 hours for 36 hours. Specimens obtained during the febrile state yield the greatest number of parasites in circulating blood.</p> <p>Prepare fresh finger stick thin smears and thick smears.</p> <p>Thin smears: Collect a small drop of blood near one end of a slide, and then spread the blood over the surface with a second slide. The thin, feathered end should be at least 2 cm long, and the film should occupy the central area of the slide, with free margins on each side.</p> <p>Thick smears: Prepare by touching the slide to the drop of blood (which should be rounded up on the finger). Rotate the slide to form a circular film about the size of a dime that is</p> |

| Component | Special Notations |
|-------------------------------|---|
| | <p>made up barely visible thorough wet smear.</p> <p>Allow for complete air-drying of smears.</p> <p>Label the frosted end of the slides using a pencil. Include the patient name, medical record number, date and accession number.</p> <p>The phlebotomist must hand the slides directly to a technologist.</p> <p>Refer to Phlebotomy procedure Malaria Smear Collection.</p> |
| Special Collection Procedures | <p>Collection procedure for the Germantown Emergency Department ONLY:</p> <p>Because of limitations at Germantown for fingerstick collection, Malaria specimens may be collected in an EDTA lavender tube. Smears must be made at GEC within 30 minutes of collection in order to reduce distortion of the parasites and RBCs.</p> <p>The thin and thick smears (at least 2 of each) will be prepared at the Germantown ED and all of the smears and the EDTA tube will be sent to Shady Grove via STAT courier. The smears will be stained and examined at Shady Grove.</p> |
| Other | A Malaria History Form is to be completed for each patient. |

3.2 Specimen Type & Handling

| Criteria | |
|--|--|
| Type -Preferred -Other Acceptable | <p>Two thin and two thick smears</p> <p>Note: ONLY Germantown Emergency Department may accept an EDTA tube less than 30 minutes old. Thick and thin smears and the EDTA tube should be sent to Shady Grove.</p> |
| Collection Container | See section 3.1 |
| Volume - Optimum - Minimum | N/A N/A |
| Transport Container and Temperature | Slide holder at room temperature |
| Stability & Storage Requirements | <p>Room Temperature: 1 month slides 30 minutes EDTA tube</p> <p>Refrigerated: Unacceptable</p> <p>Frozen: Unacceptable</p> |
| Timing Considerations | N/A |
| Unacceptable Specimens & Actions to Take | If specimen is too old test must not be performed. Improperly prepared or improperly labeled slides. Reject specimen and request recollection. |

| Criteria | |
|---------------------------------------|---|
| Compromising Physical Characteristics | N/A |
| Other Considerations | Treatment with anti-malarial or other antiparasitic drugs may reduce the sensitivity of the test. |

NOTE: Labeling requirements for all reagents, calibrators and controls include: (1) Open date, (2) Substance name, (3) Lot number, (4) Date of preparation, (5) Expiration date, (6) Initials of tech, and (7) Any special storage instructions. Check all for visible signs of degradation.

4. REAGENTS

The package insert for a new lot of kits must be reviewed for any changes before the kit is used. A current Package Insert is included as a Related Document.

4.1 Reagent Summary

| Reagents | Supplier and Catalog Number |
|--------------|---|
| Giemsa Stain | Harleco – 620G-75 |
| Buffer | Alphatec Giemsa (Malaria) Stain Buffer – 033-25 |

4.2 Reagent Preparations and Storage

| Reagent | Giemsa Stain |
|-------------|------------------------------|
| Container | 1 L bottle |
| Storage | 15-30°C |
| Stability | Stable until expiration date |
| Preparation | None |

| Reagent | Alphatec Giemsa (Malaria) Stain Buffer |
|-------------|--|
| Container | 125 mL bottle |
| Storage | 15-30°C |
| Stability | Stable until expiration date |
| Preparation | None |

5. CALIBRATORS/STANDARDS

N/A

6. QUALITY CONTROL

6.1 Controls

Appearance of blood cells is noted every time a patient's smear for malaria is performed.

Romanowsky Color Range

| | |
|--|-------------------------|
| Chromatin of white blood cells | purple |
| Nuclei of parasitic protozoa | red |
| Basophilic cytoplasm of lymphocytes, monocytes, and parasitic protozoa | blue |
| Eosinophilic granules | pink |
| Neutrophilic granules | purple |
| Red blood cells | salmon pink (to bluish) |
| Bacteria | deep blue |

Record QC results on Malaria Stain QC Form.

If controls are unacceptable do not report patient results, notify supervisor.

6.2 Control Preparations and Storage

N/A

6.3 Frequency

Each batch of patient smears is evaluated for proper staining characteristics.

6.4 Tolerance Limits and Criteria for Acceptable QC

A run is rejected if the WBCs, RBCs, and platelets on the thin smear are not stained adequately.

Rejected runs must be effectively addressed by corrective action. Steps taken in response to QC failures must be documented. Patient samples in failed analytical runs must be reanalyzed.

6.5 Documentation

Steps taken in response to QC failures must be documented.

6.6 Quality Assurance Program

N/A

7. EQUIPMENT and SUPPLIES

7.1 Assay Platform

None

7.2 Equipment

Microscope

Wescor Hematology Slide Stainer

7.3 Supplies

- Immersion oil
- Glass slides
- Geimsa Stain
- Buffer

8. PROCEDURE

NOTE: For all procedures involving specimens, buttoned lab coats, gloves, and face protection are required minimum personal protective equipment. Report all accidents to your supervisor.

| 8.1 | Thin Smears |
|-----|---|
| 1. | Allow smear to dry thoroughly before staining. |
| 2. | Place patient thin smears on Wescor Hematology Slide Stainer. (see Hematology procedure for stainer instructions) |
| 3. | Examine thin smears under 10X and 100X (oil immersion) to screen for the presence of malarial parasites, <i>Babesia</i> , microfilaria, and trypanosomes. |
| 4. | At least 300 fields must be viewed with a 100X oil immersion lens for adequate assessment. |

| 8.2 | Thick Smears |
|-----|---|
| 1. | Allow smear to dry thoroughly before staining (at least 2 hours). |
| 2. | Do not fix with alcohol or heat or dry in an incubator. Heat will prevent RBC lysis. |
| 3. | Obtain working Giemsa solution. Into a Coplin jar add 49ml of the phosphate buffer, 1 ml of the Giemsa Blood Stain. Mix well before use. The working stain solution is stable for 24 hours. |
| 4. | Place the thick smears directly into the working solution for 45-60 minutes. The water-based Giemsa stain disrupts the red cell membrane (laking) during the staining procedure exposing the parasites. |
| 5. | Wash the smears by rinsing them with buffer (pH 7.0 to 7.2) for 3-5 minutes. |
| 6. | Record pH of buffer on QC sheet. |
| 7. | Air-dry in a vertical position. |

| 8.3 | Reading |
|-----|--|
| 1. | Scan the smear under low power first to detect presence of microfilaria or trypanosomes. |
| 2. | Next read under oil immersion (100X objective). |
| 3. | At least 300 fields under oil immersion must be examined. |
| 4. | All shifts will stain thin and thick smears and screen thin smears for malaria, <i>Babesia</i> , microfilaria, and trypanosomes. |

| 8.3 | Reading |
|-----|--|
| 5. | If slides must be sent to another site for interpretation, keep 1 set of slides at the originating site. |
| 6. | If positive and the species cannot be determined after review by a microbiology lead tech, supervisor or director, thin and thick smears may be sent to Washington Adventist Hospital to Dr. Beltaifa for pathologist review if she is available. If Dr. Beltaifa is not available, refer the slides to Chantilly. |

9. CALCULATIONS

Parasitemia: In areas of the slide where the RBCs are evenly spread out over the entire field and not overlapping count the number of infected cells per field of 200 cells. Do this on 10 different areas on each thin smear. Calculate the average and divide by 2. The resulting number is the percentage of RBC's infected.

For parasitemia lower than 0.01%, report the parasitemia as <0.01%.

10. REPORTING RESULTS AND REPEAT CRITERIA

10.1 Interpretation of Smears

Smears are examined utilizing 10X and oil immersion lens (100x).

Read a minimum of 300 fields under oil immersion before determining that the thin smears are negative.

Thick smears are to be read before finalizing the report as negative. Thick smears are also a guide to the intensity of the infection. Thick smears allow a large amount of blood to be examined, increasing the detection of parasites in light infections. If parasites are detected on the thick smears, species determination must be made using the thin smear examination. This is determined by the recovery and identification of life cycle stages observed on the thin smear.

10.2 Reporting

10.2.1 General Information

Call both positive preliminary and final results to the nursing unit or physician. The call back information must be documented in the LIS.

Preliminary Reports:

If thin smears are negative, report: "Thin smear presumptive negative, thick smear and final report to follow". (NMLP1)

If thin smears are positive, report: "Presumptive positive, confirmation and identification to follow." (PMAL1)

If *Plasmodium falciparum* can be ruled out, report *Plasmodium* species, not *P. falciparum*. Send all of the smears (thin and thick) to Washington Adventist to Dr. Beltaifa for pathologist review.

If microfilaria or trypanosomes are seen, report their presence and send to Chantilly for confirmation.

*** Do not finalize the thin smear preliminary report in the LIS.**

Final Reports:

If negative, report: "No parasites seen. One set of blood films can not exclude the diagnosis of malaria." (NMAL1)

- If positive report:
1. report genus and species for malaria or "Babesia species" if Babesia is seen.
 2. report the level of parasitemia

Parasitemia:

Report the percentage of cells infected on all positive *Plasmodium* species or *Babesia* species. See section 9 for instructions on performing the calculation.

Report: "x.xx % Parasitemia"

Enter the number and % sign, then enter the code INF2 or enter using the Sunquest keyboard as in 10.2.2 below.

The call back information must be documented in the LIS.

*** More than one technologist must review all initial positive malaria smears. Repeated positive smears on the same patient do not require review by a second technologist.**

Document both tech codes on the LIS workcard.

Reporting to Maryland DHMH:

Smears positive for malaria or *Babesia* species must be reported by the technologist who reported the result to the Maryland Department of Health and Mental Hygiene by completing DHMH form 1281. Reports must be submitted within 1 working day (fax or mail).

Fax the form to the Montgomery County Health Department (240-777-4680).

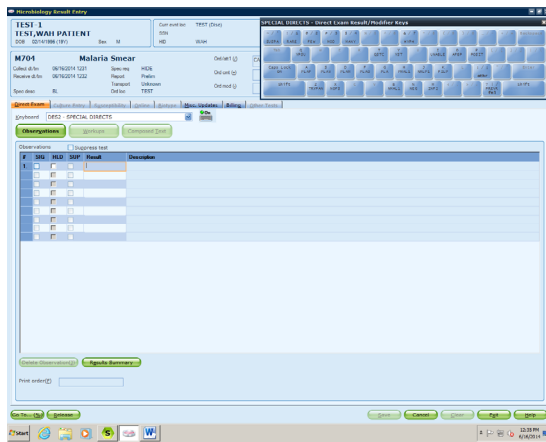
If mailing, mark the sealed envelope "confidential" and send to
Montgomery County Health Department
2000 Dennis Ave
Suite 238
Silver Spring, MD 20992

One set of thick and thin smears must be sent to the Maryland DHMH.

- Place slides in a plastic slide holder and package in a padded shipping envelope or box.
- Form DHMH 4676 must be completed and must accompany the slides.
- Address the package to Maryland Department of Health and Mental Hygiene, 201 W Preston St., Baltimore, MD 21201.
- Place package with the newborn screen samples for courier pickup

10.2.2 LIS resulting

1. Use GUI function **Micro Result Entry**.
2. Key in the accession # and click on **SELECT**.
3. Press on F8 to display the resulting keyboard. Note: to turn off the keyboard press F8 again.



4. Click on the Micro keyboard to enter in your results
5. If you have a positive malaria smear then result as follows:
 - a. **Observation #: Organism** - Click on the organism from the Micro keyboard, then press the tab key until you are at the next observation line in the **result** field.
 - b. **Observation #: Infectivity rate** - Press ; twice (the first ; will not display on the screen but the second one will) and then free text the infectivity rate (example 2.0 %), then press the tab key. From the Micro keyboard, click on the **M** key. This will add **-Parasitemia** to your infectivity rate.

c. **Observation #:** ;CBACK <cr> (expands to 'Called to and read back by:'); ; add free text call documentation <cr> <cr>

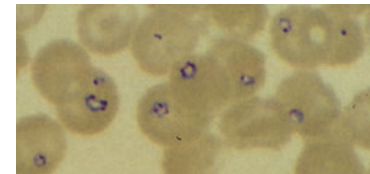
6. Press the tab key twice so that your cursor is in the next result field.
7. If this is a prelim, click on **SAVE**
8. If this is a final, click on the / (**final**) key from the Micro Keyboard and then click on **SAVE**.

* Each call must be documented. Do not delete previous call back information.

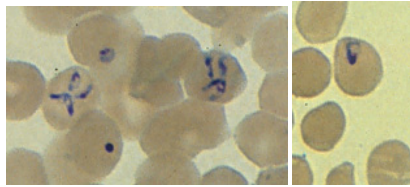
10.3 Interpretation of data

Morphological Characteristics of *Plasmodium falciparum* and *Babesia* species:

| Appearance of parasite | <i>Plasmodium falciparum</i> | <i>Babesia</i> species |
|--------------------------|--|---|
| Size | Small (1/4 to 1/3 RBC diameter, 3-5 µm) | Tiny to small (1/8 to 1/4 RBC diameter, 1-5 µm) |
| Shape | Consistent oval to round ring | Pleomorphic: pear-shaped to round ring |
| Appliqué Forms | Common, either marginal or bulging forms | Common, either marginal or bulging forms |
| Number of Chromatin dots | 1 to 2 | 1 to many ("string of pearls") |
| Multiple rings/RBC | Common | Common; two adjacent parasites may appear to be split into mirror images |
| Tetrads | No | Rarely seen |
| Appearance of RBCs | Normal size and shape | Normal size and shape |
| Parasite stages present | Ring: trophozoite with pigment (in heavy infections); banana-shaped gametocytes (rarely found) | Ring: late ring or trophozoite with no pigment, may contain a white central vacuole not seen in <i>Plasmodium</i> |



A: *Babesia microti* infection, Giemsa-stained thin smear. The organisms resemble *Plasmodium falciparum*; however *Babesia* parasites present several distinguishing features: they vary more in shape and in size; and they do not produce pigment.



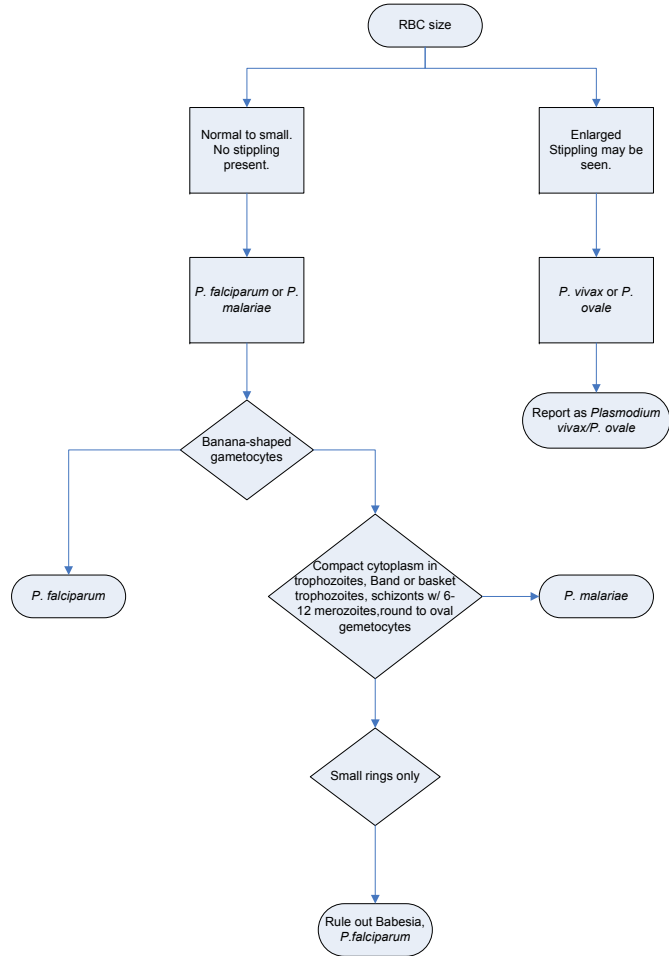
B. and C: Infection with *Babesia*. Giemsa-stained thin smears. Note in **B** the tetrad (left side of the image), a dividing form pathognomonic for *Babesia*. Note also the variation in size and shape of the ring stage parasites (compare **B** and **C**), and the absence of pigment.

Morphology of *Plasmodium* species in Wright-Giemsa stained smears:

| Characteristics | <i>P. falciparum</i> | <i>P. vivax</i> | <i>P. ovale</i> | <i>P. malariae</i> |
|---|---|--|---|--|
| Size and shape of infected erythrocytes | Normal size and shape | Enlarged up to twofold, may be oval | Normal to enlarged, frequently oval, may be fimbriated | Small to normal size, normal shape |
| Stippling | Occasional Mauer's dots, less numerous than Schuffner's | Schuffner's dots (stippling) usually present, except in rings | James' stippling, darker than Schuffner's, present in all stages, including rings | Zeiman's dots rarely seen; requires deliberate over staining |
| Stages seen in peripheral blood | Rings and gametocytes | All | All | All |
| Multiply infected erythrocytes | Common | Occasional | Occasional | Rare |
| Early trophozoites | Delicate ring, frequently with two small chromatin dots, often at the edge of the erythrocyte | Ring up to 1/3 diameter of the erythrocyte; larger chromatin dot than <i>P. falciparum</i> | Similar to <i>P. vivax</i> | Smaller than <i>P. vivax</i> ; otherwise similar |
| Mature trophozoites | Not seen in peripheral blood | Amoeboid shape, fine golden brown pigment | Similar to <i>P. vivax</i> except less amoeboid, pigment darker brown | Compact cytoplasm, oval, round, or band-shaped, dark brown pigment |

| Characteristics | <i>P. falciparum</i> | <i>P. vivax</i> | <i>P. ovale</i> | <i>P. malariae</i> |
|------------------------------|---|--|---|---|
| Schizonts | Not seen in peripheral blood | 12-24 merozoites | 8-12 merozoites | 6-12 merozoites often radically arranged around central pigment in a rosette form |
| Gametocytes | Crescent of banana-shaped | Round to slightly oval | Round to slightly oval | Round to slightly oval |
| Most characteristic findings | Absence of mature trophozoites and schizonts; normal size of infected erythrocytes; multiply infected RBCs; appliqué forms; banana-shaped gametocytes | Enlarged infected erythrocytes; Schuffner's dots frequently present; amoeboid trophozoite; 12-14 merozoites in each schizont | Normal to enlarged, oval or fimbriated infected RBCs, James' stippling may be seen in rings; schizonts with 8-12 merozoites | Normal size of infected erythrocytes; no stippling; "band" trophozoite; rosette schizont with 6-12 merozoites |

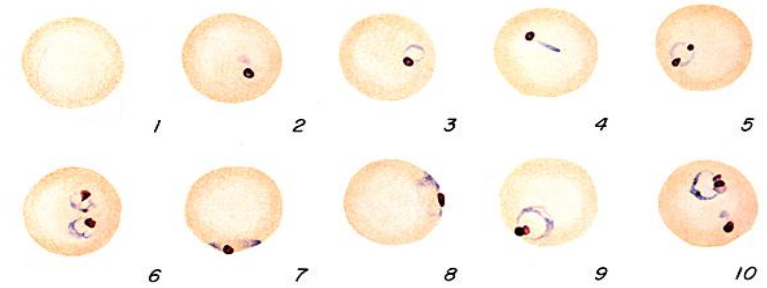
Malaria Identification



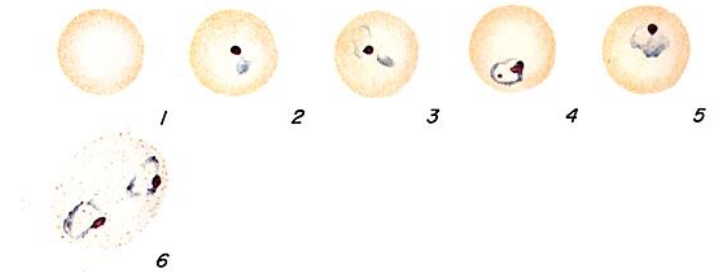
Always calculate and report % parasitemia

Ring Stage Parasites

Plasmodium falciparum: Rings



Plasmodium vivax: Rings



Plasmodium ovale: Rings

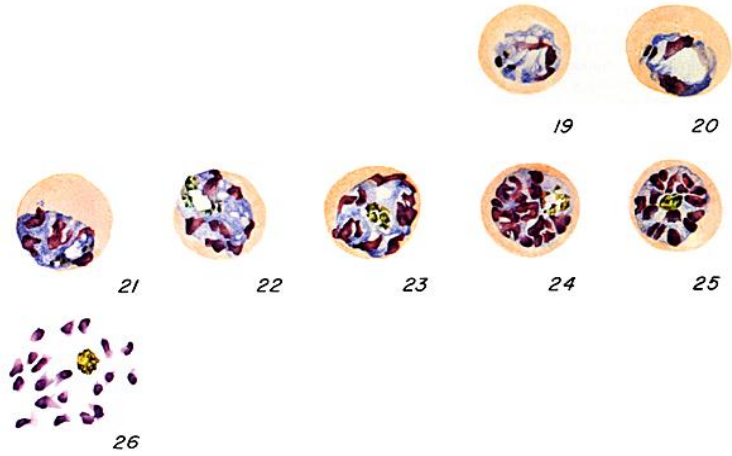


Plasmodium malariae: Rings

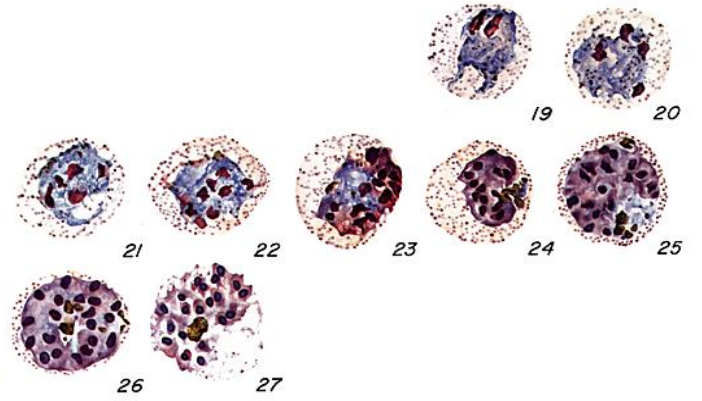


Schizonts

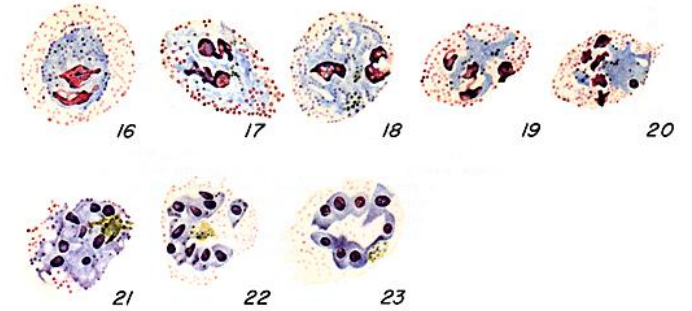
Plasmodium falciparum: Schizonts (usually not seen in blood)



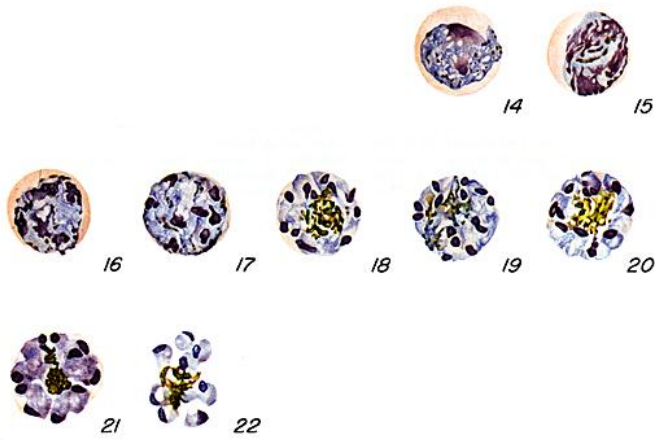
Plasmodium vivax: Schizonts



Plasmodium ovale: Schizonts

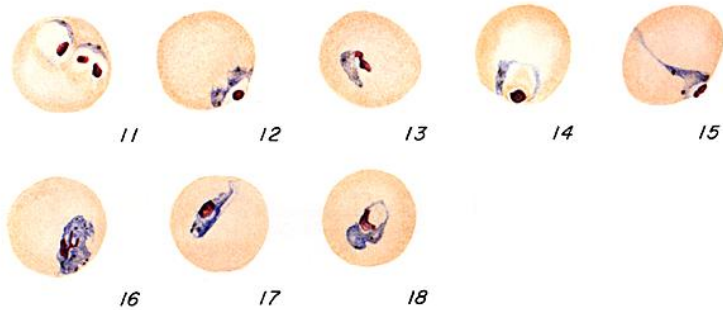


Plasmodium malariae: Schizonts

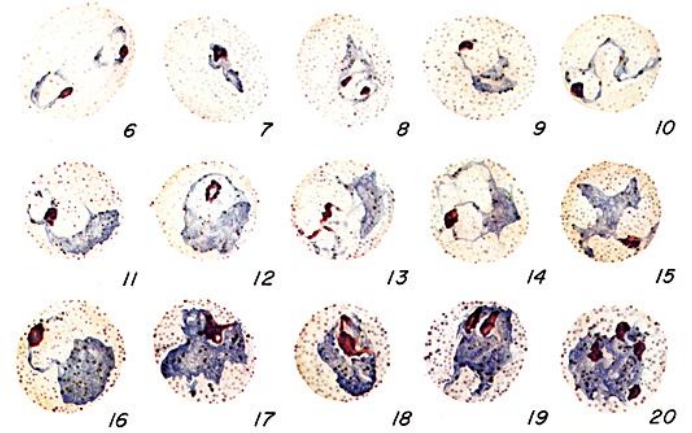


Trophozoites

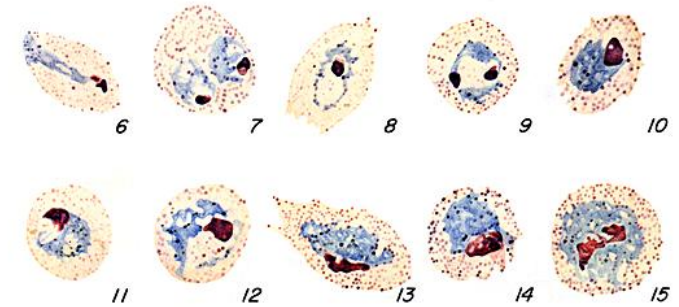
Plasmodium falciparum: Trophozoites (early forms may be seen but later forms usually not seen in blood)



Plasmodium vivax: Trophozoites



Plasmodium ovale: Trophozoites

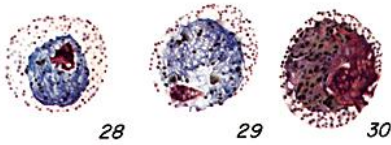


Gametocytes

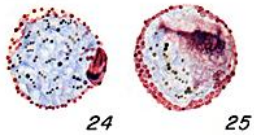
Plasmodium falciparum: Gametocytes



Plasmodium vivax: Gametocytes



Plasmodium ovale: Gametocytes

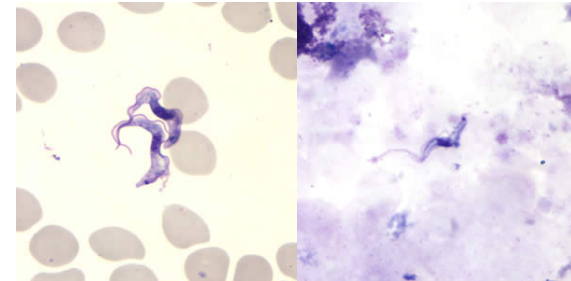
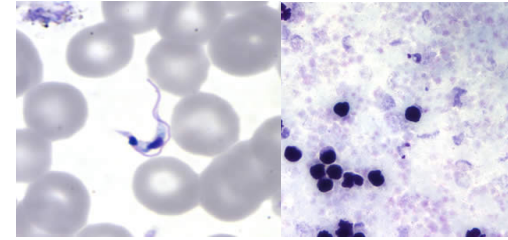


Plasmodium malariae: Gametocytes

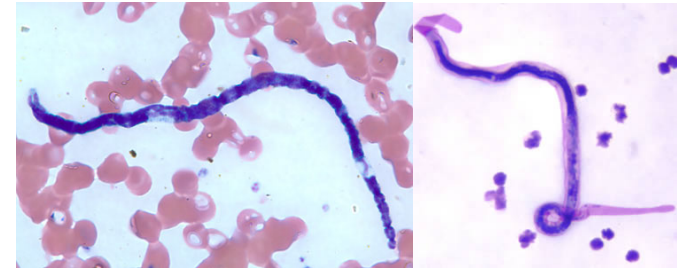


Morphology of trypanosomes and microfilaria.

Trypanosomes



Microfilaria



10.5 Rounding / Units of Measure / Clinically Reportable Range (CRR)

N/A

10.6 Review Patient Data

Review patient results for unusual patterns, trends or distributions in patient results such as an unusually high percentage of abnormal results.

10.7 Repeat Criteria and Resulting

N/A

11. EXPECTED VALUES

11.1 Reference Ranges

No parasites seen.

11.2 Critical Values

Any positive smear

11.3 Standard Required Messages

None

12. CLINICAL SIGNIFICANCE

Malaria is a disease of worldwide importance characterized by fever, anemia and splenomegaly. Although four species of the genus *Plasmodium* (*P. falciparum*, *P. vivax*, *P. malariae*, and *P. ovale*) infect humans, malaria is clinically two diseases; the benign type due to *P. vivax*, *P. malariae*, and *P. ovale*, and the malignant type due to *P. falciparum*.

Determination of parasitemia becomes important when therapy is initiated. The patient's parasitemia is monitored so that possible cases involving drug-resistant strains of *P. falciparum* may be detected. In those cases where the patient is hospitalized, monitoring of the parasitemia should be performed at 24, 48 and 72 hours after initiating therapy. Generally, if the malarial strain is susceptible to the therapeutic regime, the parasitemia will drop significantly within the first 24 hours (often by 50% or more).

Babesia is a malaria-like disease characterized by fever, chills, headache, lethargy and myalgia. Hemolytic anemia and hemoglobinuria are typical and may be severe. The disease is transmitted by the bite of hard ticks of the family Ixodidae. This disease is suspected when individuals have traveled through tick-infested areas and present with a malaria-like illness. The disease becomes apparent 1-3 weeks after the bite of an infectious tick. In splenectomized and immunocompromised patients this disease may be fatal. Determination of % parasitemia helps direct therapy. In severe parasitemia (>10%), exchange transfusion may be considered.

Parasitemia and Clinical Correlation

| Parasitemia | Parasites /uL | Clinical Correlation |
|----------------|-------------------|---|
| 0.0001-0.0004% | 5 – 20 | Number of organism that are required for a positive thick film (sensitivity) |
| 0.002% | 100 | Patients may be symptomatic below this level |
| 0.2% | 10,000 | Level above which immune patients will exhibit systems |
| 2% | 100,000 | Maximum parasitemia of <i>P. vivax</i> and <i>P. ovale</i> (infect young RBSs only) |
| 2-5% | 100,000 – 250,000 | Hyperparasitemia, severe malaria, increased mortality |
| 10% | 500,000 | Exchange transfusion may be considered, high mortality |

13. PROCEDURE NOTES

- **FDA Status:** LDT without message
- **Validated Test Modifications:** None

Any alcohol left on the skin prior to collection may fix the red cells and then they will not clear in the staining procedure.

Do not dry smears using heat, as this will fix the red cells.

Slides prepared from EDTA blood are not optimal as they may cause distortion in the parasites, making identification difficult. However, the Emergency Center at Germantown is the ONLY location where an EDTA specimen is acceptable rather than fingertip smears.

Organisms are most likely to be detected if the smears are obtained immediately upon the onset of fever, or immediately before the fever is anticipated. In patients with a strong clinical history, but repeatedly negative results, multiple sampling throughout the fever may prove successful.

Platelets sitting on top of red blood cells may have the appearance of a ring form of malaria.

Precipitated stain may obscure malarial forms on the smear.

Identification to species should not be based solely on the examination of the thick smear preparation. Both thick and thin smears are required for a comprehensive blood parasite examination.

The patient's travel history may provide helpful information in the identification of malaria, *Babesia* species, and other blood parasites. Blood parasites are endemic to certain regions of the world; knowing what countries the patient has visited will aid in diagnosis.

The chart below can be used as a guide for diagnosis. It is not to be used as the primary diagnostic factor:

| Blood Parasite | Endemic Area(s) |
|------------------------------|---|
| <i>Plasmodium falciparum</i> | Africa, Asia, Indian subcontinent, South America (Tropical areas worldwide) |
| <i>Plasmodium vivax</i> | Tropical and Temperate areas worldwide |
| <i>Plasmodium malariae</i> | Africa, Asia, Indian subcontinent, South America (Tropical areas worldwide) |
| <i>Plasmodium ovale</i> | West Africa, India, South America, some South Pacific Islands |

14. LIMITATIONS OF METHOD

14.1 Analytical Measurement Range (AMR)

N/A

14.2 Precision

N/A

14.3 Interfering Substances

N/A

14.4 Clinical Sensitivity/Specificity/Predictive Values

N/A

15. SAFETY

Refer to your local and corporate safety manuals and Safety Data Sheet (SDS) for detailed information on safety practices and procedures and a complete description of hazards.

16. RELATED DOCUMENTS

Hematology Slide Stainer Cyto centrifuge, Wescor Aerospray; Hematology procedure
 Resulting Microbiology Direct Exams, Microbiology procedure
 Malaria Smear Collection, Phlebotomy procedure
 Malaria Smear Collection – GEC Only, GEC Microbiology procedure
 Malaria History Form (AG.F289)
 Reportable Results to State and Outside Agencies, Laboratory policy

17. REFERENCES

- Jacobs DS, et al, Laboratory Test Handbook, 4th edition, Hudson, OH: Lexi-Comp, Inc., 1999, pp. 332-333.
- Kjeldsberg C, et al, Practical Diagnosis of Hematologic Disorders, 2nd edition, Chicago, IL: ASCP Press, 1995, pp. 172-173.
- Atlas of Human Parasitology, 3rd edition, Chicago, IL: ASCP Press.

- Hansheid, T. 1999. Diagnosis of malaria: A review of alternatives to conventional microscopy. Clin. Lab. Haematol. 21:235-245.
- Wilkinson, R.J., J.L. Brown, G. Pasvol, P.L. Chiodini, and R.N. Davidson. 1994. Severe falciparum malaria: predicting the effect of exchange transfusion. Q.J. Med. 87:553-557.

18. REVISION HISTORY

| Version | Date | Section | Reason | Reviser | Approval |
|---------|-----------|-----------------------|--|---------------|-----------|
| | | | Supersedes SOP M028.005 | | |
| 000 | 10/12/09 | 10.2.2 | LIS upgrade to GUI system | A. Sears | R. Master |
| 000 | 10/12/09 | 16 | Added procedure for resulting | L. Barrett | R. Master |
| 001 | 9/19/2011 | 3.1, 13 | Added use of EDTA specimen at GEC | C. Reidenauer | R. Master |
| 001 | 9/19/2011 | 4.2 | Changed storage temperature for buffer | R. Master | R. Master |
| 001 | 9/19/2011 | 8 | Remove statement regarding pkg insert | L. Barrett | R. Master |
| 001 | 9/19/2011 | 8.3 | Added trypanosomes and microfilaria | R. Master | R. Master |
| 001 | 9/19/2011 | 11.2 | Update title to local terminology | L. Barrett | R. Master |
| 002 | 11/19/12 | 9 10.2.1 10.2.2 | Change report to "Parasitemia followed by the % infectivity". Changed steps on how to report the % infectivity (English Text code first then free text the rate %) | M. Sabonis | R. Master |
| 003 | 7/17/14 | 3.1, 3.2 | Changed EDTA time to 30 min. Removed sending all thick smears to WAH. | R. Master | R. Master |
| 003 | 7/17/14 | 8.2 | Add stability of working solution. Removed comment to send all think smears to WAH. | R. Master | R. Master |
| 003 | 7/17/14 | 8.3, 10.2 | Add to send to Dr. Beltaifa if species could not be determined. | R. Master | R. Master |
| 003 | 7/17/14 | 9 | Clarified calculation | R. Master | R. Master |
| 003 | 7/17/14 | 10.2 | Add preliminary report of <i>Plasmodium</i> species, not <i>P. falciparum</i> . | R. Master | R. Master |
| 003 | 7/17/14 | 10.2.1 | Change order of reporting parasitemia, deleted redundant calculation | R. Master | R. Master |
| 003 | 7/17/14 | 10.2.2 | Add instructions for entering results in GUI version of LIS | R. Master | R. Master |
| 003 | 7/17/14 | 10.3 | Added flow chart | R. Master | R. Master |
| 003 | 7/17/14 | 16 | Update titles, add form number | L. Barrett | R. Master |
| 003 | 7/17/14 | Footer | Version # leading zero's dropped due to new EDCS in use as of 10/7/13 | L. Barrett | R. Master |
| 4 | 9/20/16 | Header | Add WAH | L. Barrett | R. Master |
| 4 | 9/20/16 | 3.1 | Add date and MR# to slide label | R. Master | R. Master |

| | | | | | |
|---|---------|--------|--|-----------|-----------|
| 4 | 9/20/16 | 4 | Update to new standard labeling instruction | L Barrett | R. Master |
| 4 | 9/20/16 | 8.3 | Add stain thin and thick smears. Add instructions to keep 1 set of slides at originating site. Clarified options if species cannot be determined | R. Master | R. Master |
| 4 | 9/20/16 | 10.2 | Add reporting of identification and parasitemia of Babesia | R. Master | R. Master |
| 4 | 9/20/16 | 10.2 | Clarified reporting of patients with repeat positive results. Added reporting and sending slides to Maryland DHMH | R. Master | R. Master |
| 4 | 9/20/16 | 10.6 | Move patient review from section 6 | R. Master | R. Master |
| 4 | 9/20/16 | 15 | Update to new standard wording | L Barrett | R. Master |
| 4 | 9/20/16 | 16 | Added SOP Reportable Results to State and Outside Agencies | R. Master | R. Master |
| 4 | 9/20/16 | 19 | Added Maryland DHMH forms | R. Master | R. Master |
| 5 | 2/18/20 | Header | Changed WAH to WOMC | L Barrett | R. Master |
| 5 | 2/18/20 | 9 | Added minimum parasitemia level of <0.01% | R. Master | R. Master |

19. ADDENDA

Maryland form DHMH 1281
 Maryland form DHMH 4676

**CONFIDENTIAL REPORT: LABORATORY EVIDENCE OF CERTAIN COMMUNICABLE DISEASES
 USE FOR REPORTING TO: MARYLAND STATE DEPARTMENT OF HEALTH AND MENTAL HYGIENE**

USE FOR ALL COMMUNICABLE CONDITIONS EXCEPT HIV and CD4. (Use form DHMH 4492 for HIV and CD4.)

(PLEASE TYPE OR PRINT USING BLACK INK.)

| | | | | | | | |
|---|--------|--------------------------------|----------------|---|-----|--|-------------------|
| PATIENT LAST NAME | | FIRST | MIDDLE INITIAL | HOSPITAL NUMBER | | PREGNANT? (FEMALE) | |
| | | | | | | YES <input type="checkbox"/> NO <input type="checkbox"/> | |
| DATE OF BIRTH | | AGE | SEX | ETHNICITY | | RACE | |
| | | | | HISPANIC <input type="checkbox"/> NON-HISPANIC <input type="checkbox"/> | | | |
| NUMBER | STREET | APT | CITY | STATE | ZIP | COUNTY | (AREA CODE) PHONE |
| | | | | | | | |
| ORDERING PROVIDER | | NAME | | | | | |
| | | | | | | | |
| NUMBER | STREET | SUITE | CITY | STATE | ZIP | COUNTY | (AREA CODE) PHONE |
| | | | | | | | (AREA CODE) FAX |
| ORDERING FACILITY NAME | | | | | | | |
| | | | | | | | |
| NUMBER | STREET | SUITE | CITY | STATE | ZIP | COUNTY | (AREA CODE) PHONE |
| | | | | | | | |
| DATE SPECIMEN COLLECTED | | DATE SPECIMEN RECEIVED | | DATE RESULTED | | LAB ACCESSION NUMBER | |
| | | | | | | | |
| TYPE OF SPECIMEN | | | | | | | |
| Sputum <input type="checkbox"/> | | Stool <input type="checkbox"/> | | Pharyngeal Swab <input type="checkbox"/> | | Discharge <input type="checkbox"/> | |
| Blood <input type="checkbox"/> | | CSF <input type="checkbox"/> | | Washing <input type="checkbox"/> | | Other (Specify) _____ | |
| SITE OF SPECIMEN (CERVIX, EYE, ETC.) | | | | | | | |
| | | | | | | | |
| NAME OF TEST | | | | TEST NUMBER OR CODE | | | |
| | | | | | | | |
| RESULT WITH REFERENCE RANGE & INTERPRETATION | | | | | | | |
| | | | | | | | |
| (IF AN ORGANISM RESULT: INCLUDE SPECIES, SEROGRUOPING, OR OTHER SUBTYPING IF KNOWN) | | | | | | | |
| IF A HEPATITIS C RESULT: | | | | | | | |
| Signal to Cut-Off Ratio (SCO) | | Critical Value for SCO | | Hepatitis A IgM Result | | Hepatitis B Core IgM Result | |
| | | | | | | | |
| LAB NAME (LAB PERFORMING THE TEST) | | | | | | LAB CLIA NUMBER | |
| | | | | | | | |
| LAB ADDRESS | | | | | | | |
| | | | | | | | |
| LAB DIRECTOR | | | | LAB (AREA CODE) PHONE | | DATE OF REPORT | |
| | | | | | | | |

DHMH 1281 **SEND TO YOUR LOCAL HEALTH DEPARTMENT**
 Revised JAN 26, 2012 For more forms or information, go to <http://idsha.dhmh.maryland.gov/SitePages/what-to-report.aspx>



Laboratories Administration MD DHMH
 201 W. Preston St. • Baltimore, MD 21201
 P.O. Box 2355 • Baltimore, MD 21203-2355
 410-767-6100 www.dhmh.state.md.us/labs
 Robert A. Myers, Ph.D., Director

STATE LAB
Use Only

INFECTIOUS AGENTS: CULTURE/DETECTION

| | | |
|---|--|--|
| TYPE OR PRINT REQUIRED INFORMATION OR PLACE LABELS ON ALL FOUR COPIES | <input type="checkbox"/> HPI <input type="checkbox"/> PP <input type="checkbox"/> IMTY:PN <input type="checkbox"/> ONOD <input type="checkbox"/> DSTD <input type="checkbox"/> TB <input type="checkbox"/> CDC <input type="checkbox"/> UDOR | Patient SS# (last 4 digits): |
| | Health Care Provider | Last Name |
| | Address | First Name M.I. Maiden |
| | City County | Date of Birth (mm/dd/yyyy) / / |
| | State Zip Code | Address |
| | Contact Name: | City County |
| | Phone# Fax# | State Zip Code |
| | Test Request Authorized by: | |
| | Sex: <input type="checkbox"/> Male <input type="checkbox"/> Female <input type="checkbox"/> Transgender M to F <input type="checkbox"/> Transgender F to M | Ethnicity: Hispanic or Latino Origin? <input type="checkbox"/> yes <input type="checkbox"/> no |
| | Race: <input type="checkbox"/> American Indian/Alaska Native <input type="checkbox"/> Asian <input type="checkbox"/> Black/African American <input type="checkbox"/> Native Hawaiian/Other Pacific Islander <input type="checkbox"/> White | |
| Case # DOC# Outbreak # | Submitter Label# | |
| Collect Date: Collect Time: <input type="checkbox"/> AM <input type="checkbox"/> PM Onset Date: | | |
| Reason for Test: <input type="checkbox"/> Screening <input type="checkbox"/> Diagnosis <input type="checkbox"/> Contact <input type="checkbox"/> Test of Cure <input type="checkbox"/> 2-3 Months Post Rx <input type="checkbox"/> Suspected Carrier <input type="checkbox"/> Isolate for ID <input type="checkbox"/> Release | | |
| Therapy/Drug Treatment: <input type="checkbox"/> No <input type="checkbox"/> Yes | Therapy/Drug Type: Therapy/Drug Date: | |

| ↓ SPECIMEN CODE | ↓ SPECIMEN CODE | ↓ SPECIMEN CODE |
|--|--|---|
| BACTERIOLOGY | SPECIAL BACTERIOLOGY | RESTRICTED TESTS |
| Bacterial Culture - Routine | Legionella Culture | Pre-approved submitters only |
| Additional specimen codes: | Leptospira | Chlamydia trachomatis GC NAAT |
| Bordetella pertussis | Mycoplasma | Chlamydia trachomatis only/NAAT |
| Group A Strep | MYCOBACTERIOLOGY/AFB/TB | Norovirus ** (see comment on back) |
| Group B Strep Screen | A-F:TB Culture and Smear | OTHER TESTS FOR INFECTIOUS AGENTS |
| C. difficile Toxin | AFB:TB Referred Culture for ID | Test name: |
| Diphtheria | M. tuberculosis Referred Culture for Genotyping | |
| Foodborne Pathogens (E. cereus, C. perfringens, S. aureus) | Nucleic Acid Amplification Test for M. tuberculosis Complex (MTD) | Prior arrangements have been made with the following DHMH Laboratories Administration employee: |
| Gonorrhea Culture: Incubated? <input type="checkbox"/> Yes <input type="checkbox"/> No | PARASITOLOGY | |
| Hrs. incubated: Add'l specimen codes: | Blood Parasites: | SPECIMEN CODE: |
| MRSA (rule out) | Country visited outside US: | PLACE CODE IN BOX NEXT TO TEST |
| VRE (rule out) | Cvs & Parasites: Immigrant? <input type="checkbox"/> Yes <input type="checkbox"/> No | B Blood |
| ENTERIC INFECTIONS | Cryptosporidium | BW Bronchial Washing |
| Campylobacter | Cyclospora/Isospora | CSF Cerebrospinal Fluid |
| E. coli O157 typing | Micospordium | CX Cervix/Endocervix |
| Enteric Culture - Routine (Salmonella, Shigella, E. coli O157, Campylobacter) | Pinworm | E Eye |
| Salmonella typing | VIRUS ISOLATION/CHLAMYDIA | F Feces |
| Shigella typing | Adenovirus* | N Nasopharynx/Nasal |
| V. parahaemolyticus | Arbovirus Panel (WNV, EEEV, SLEV) | P Penis |
| Yersinia | Chlamydia trachomatis | R Rectum |
| REFERENCE MICROBIOLOGY | Cytomegalovirus (CMV) | SP Sputum |
| ABC'S (BIDS) # | Enterovirus (Inc. Echo & Coxsackie) | T Throat |
| Organism: | Herpes Simplex Virus (Types 1 & 2) | URE Urethra |
| Bacteria Referred Culture for ID | Influenza (Types A & B)* | UVU Urine (First Void) |
| Specify: | Parainfluenza (Types 1, 2 & 3)* | UCC Urine (Clean Catch) |
| | Respiratory Syncytial Virus (RSV)* | V Vagina |
| | Varicella (VZV) | W Wound |
| | *MAY INCLUDE RESPIRATORY SCREENING PANEL. | O Other: |
| | Comments: | |

DHMH 46/6 Revised 04/2

ORIGINAL